Textbook of Tinnitus
REFLECTIONS ON A 1,000-DAY ADVENTURES IN A RESEARCH PROJECT.

October is a very nice month in the Egyptian desert. It is also when the “Rally of the Pharaons” takes place; an intensive ride in the sand where the main objective is not to get stuck or lost and to arrive at the right place before most of the others.

In 2004, like other times, I was participating and enjoying the concentration, the scenery, and the short nights in a camp, preparing the mind and the equipment for the next day. The next day, half an hour before the end of the stage, I passed the wheel to an impatient navigator who wanted his moment of piloting glory.

A few minutes later, the car went to the wrong side of a mountain, “rolled over” several times, and landed upside down at the bottom of the hill.

Whiplash, stressful emotion, and lack of oxygen to the ear (dissection of the carotid artery); I had just landed at the perfect scenario for developing something that was totally unknown to me until then: TINNITUS!!

After 6 months of panic and useless wondering to find a cure, I was left with two choices: live with it or try to do something about it. Although accepting to live with it was probably the best cure at that moment, I chose to try to do something about it. Not out of generosity or because I thought I was called upon the task by higher duties, because:

1. Unlike other pathologies, time was on my side: I was not going to die or get worse over time
2. I had experience in organizing research
3. I had the motivation to walk in other people’s lives and invite them into a project I believed in
4. I had the time, having sold my main business believing I could not lead as well anymore
5. I had the money, and
6. I did not want to regret that I had not tried

The “program” turned out to be a venture in frustration and hope, a balancing act between logic and instinct, and maybe a little, but important milestone for successful therapies in the future. Also, and not surprisingly, it was a human adventure about people and their beliefs, their weaknesses, and their strengths. Here is how I remember it and what I would consider if it started again.

As an independent entrepreneur, I wanted to give some structure to my program, but without losing flexibility and making sure I would not “play doctor.” The main immediate points were:
1. How to finance it and through what entity
2. How to choose the people
3. How to choose and coordinate the research program, and my role in it, and
4. How and when to end it, the businessman’s “exit strategy”

How to Finance It and Through What Entity

(a) An existing pharmaceutical company would seem the most immediate choice. However, their managers are guided by long-term survival of their companies and, consequently by considerations such as short-term cash flow, risk, time to market a product, and reimbursement by health care, and are often not open to innovation if it overlaps existing businesses (like in the case of new hearing aids).

(b) Co-investing with government funding was not really an option. Tinnitus, not being a life-threatening disease, would not get a lot of attention. Moreover, government projects have a long bureaucratic approval process and once funded, they lack the flexibility to change directions during the research if the interim results so suggest.

(c) An existing association was another obvious choice. Scott Mitchell, member of the board of ATA, has written many interesting articles and believes that public non-profit organizations appear to be the best vehicle for funding tinnitus research. Although I agree with him to some extent, it is normal that every time you are managing other people’s money, you are somewhat restricted by present logic and paradigms, and have to allocate a lot of time and resources for explanations and accounting to “shareholders,” in addition to public awareness, prevention, support to patients, etc.

(d) Direct funding to individuals by an individual

As more individuals live longer and achieve financial success, they reach a point where they feel they can use their money and their experience to make a difference in a field other than their own – and make it their “legacy.”

Teaming up with one of these individuals would be risky because they are, in all likelihood strong personalities who bring into a program their style, their objectives, and their people, and since it is their “legacy” after all, often want a lot of exposure.

In addition, I wanted to try to bring together cross-border and interdisciplinary knowledge into a field where not enough was yet known to make it interesting to future participants (industry, governments, and associations) and had my own ideas on what was important – and what was going to make this possible.

Chances of improving were higher because we started from zero.

My program would be based on the idea that tinnitus research was still in a phase where to get to the next step it was better to stay away from too many “models,” and that some of it had to be done by somebody who was willing to fail, make mistakes, change his mind, not understand, and ultimately not base his decisions on risk/reward, but on people who were willing to work on a project for the right reasons and with the right attitude.

“Life is like a game of chess; the first moves are very important, but until the game is over you still have some good moves to play.”

Anne Frank
How to Choose the People

I have always been involved in science – and yet know very little. My father was a brilliant scientist, with many researchers around him. I never tried to compete directly, but learned a lot from “back stage” and over the years. He had a sign in his office that said: “if you want to lose money spend it on boats, women and research.” Even if we had not spent a lot of time together, I must have taken that part from him!

The process of choosing the scientists whom I would have liked to meet each other and work together was very intuitive, but I can try to list a few characteristics that I think are common to successful scientists – they:

- Are optimistic, but realistic
- Do not promise more than what they can deliver
- Are capable of giving bad news
- Take pleasure and attention in the growth of people around them
- Simplify and explain complicated things in a simple way
- See a problem and turn it into an opportunity
- Do not have what is called the “not invented here syndrome”: they listen with an open mind to other people’s ideas
- Recognize today’s assumptions and question them
- Look beyond the obvious
- Find a way to look at something new without rejecting the current concept
- Don’t look at an idea only to see what is wrong with it and how they can reject it
- Think and work a lot – genius ideas are a result of it
- Have a high sense of responsibility
- Always want to do things better and
- Try to do the best they can.

Some of these characteristics usually surface even in a short interview and I always saw some of them in the people who have at some stage participated in the TRI research program. I am naturally honored that they have accepted to work with TRI as I never took it for granted.

“The scientific mind does not so much provide the right answers as ask the right questions.”

Claude Levi Strauss

How to Choose and Coordinate the Research Program, and My Role in it

A traditional program would have three main components.

Leadership, to clearly identify the objectives so as to produce the results.
Organization, to identify the different functions and to allocate them to the best people.
Administration, to allocate the resources where and when necessary.

One difference in this case was that none of the participants was directly employed and that the relationship was based more on attitude and trust than otherwise. Each had their own existing activity.
The main objective was not to organize an effective research program, but to encourage multidisciplinary, interdisciplinary exchange in the belief that the right people would seize the opportunity.

Personal interaction coupled with the exposure to different therapeutic areas would combine the knowledge without setting boundaries of research, and ultimately, individuals would choose their partners in the program.

Their partners would possibly be from different areas, different levels, and different countries and cultures, and that combination would increase understanding, innovation, and the feeling that the “mission” was doable.

Over time strategic groups and their performance obligations would form. Diversification would increase the effort of coordinating their work, but would naturally identify specific areas of research.

Workgroups in pharmacology, neurostimulation, auditory stimulation, somatosensory modulation, and eventually tinnitus clinics (when the need for integrating research and clinical medicine became more evident) were formed, but these were based more on the individuals who chose to work together than on an imposed structure or organization.

Somehow the dynamics were quite different than those of a company.

Later, I would have worked more closely to improve the connection between innovation and actual therapy. I knew that existing commercial compounds generated less problems. I also had learned that successful players design the most incisive clinical trials and were not necessarily hung up on publishing a lot.

The dynamics were a strange mix of what I had lived in the past, and my role was going to shape accordingly.

Rod Davis, coach of the Team New Zealand sailing team, wrote an interesting article to explain coaching and support: The Invisible Hand. He says coaching is a weird combination of teaching, mentoring, being the hatchet man (at times), and being a “nanny,” throw it all in a blender and make something good out of it. Coaching, Rod writes, is not rocket science. In fact, it is not a science at all, it is art. Coaches provide the environment for driven talent to become champions. The ones with talent who take full advantage of the opportunities presented became champions.

Environment means unloading distractions. It means create a belief in the ability to perform in tasks that are the most important to them. He adds that a big part of self-confidence is self-responsibility: if someone knows that it is up to him to be in control of his own destiny and knows he has done all that is needed to be ready, how can he not be self-confident?

This improves the chances of success, but there are no guarantees. There are thousands of pieces to the puzzle, but if the environment is right, the end result is certainly more likely to be positive.

Interestingly enough, two successive research coordinators failed in their mission, probably because they did not see the program the same way.

I was going to try and follow Rod’s “art,” keeping in mind that it was also my role – at least at the beginning, to add strong leadership and sense of the mission, just like Grant Dalton does with the very successful Team New Zealand.

“I came in understanding that the magnitude of the issues facing the country required that I put together a team that I could delegate a whole range of different tasks to and who would be able to work well together. Over the last 6 months I have relearned that lesson – that my most important job is to get the right people in the right place, give them the freedom to innovate and to think creatively about problems,
hold them accountable for results, and make sure they are cooperating with each other and communicating on an ongoing basis.”
President Barack Obama, August 2009

How and When to End it

Basic research delivers the technology platform, the ideas, and concepts, but they are often not at first accepted by industry or peers. This is the innovation gap and it needs to be bridged by the public hand. At a certain point, there needs to be an investment of the government to share the risk: political will is not only the weakest link in the chain, but also the hardest to fix.¹

Governments, whose biggest expense is becoming health care, have a difficult task in choosing priorities. As an example, a very small percentage of cancer research spending would make a huge difference in other areas, including tinnitus.

Maybe a better way to look at it would be to present the issue in a more global way. Now that the majority of researchers agree that tinnitus is a malfunction or reorganization that takes place with the neurons in the brain, its research implications go together with the understanding of other pathologies such as Alzheimer’s or Parkinson’s that are more easily understood as terribly detrimental.

Public nonprofit organizations should help bridge the gap to government involvement in addition to encouraging awareness and prevention.

Contrary to many, I believe that it is important that at a certain point the individual sponsor disappears. A more structured and long-term mechanism has to take place. People and programs should not depend solely on the sponsor.

In this specific case, the objective was to install new energy toward an “undervalued” problem and contribute to make it a stand-alone research area for medicine. Only time will tell how much has been achieved toward that end.

“You can have a dialogue about solving future problems all you like, but if you do not behave any differently when you go out of here, it won’t make any difference.”

Dennis Meadows
“Limits to growth”

Conclusions

Strategy is about the future and then making decisions based on that. The worst thing you can do is not to have an opinion, and not make decisions.²

More than ever, success depends on our ability to learn and to create value from what we learn.

In these times of uncertainty, scientists and physicians have to be agents of change in the right direction, accelerate science, advance medicine, and also direct it in a more integrated and patient-driven experience that is comprehensive to all.

¹Peter Gruss, President Max-Planck-Society
²Alan Mulally, President Ford Motor Company
Individuals still play an important role in sponsoring and discovery. It is everybody’s task to create the environment and attitude for positive change.

Whether we made a change, and if the change was meaningful we will not know for years and maybe never. But I believe it would be a mistake to loose the momentum and coordination that TRI has created.

On a personal note, I have met some extraordinary people and scientists, although my tinnitus is still there, I believe that we have cured people who otherwise would still be suffering. I believe I will be cured in the next 3–5 years and that I will have that cure available before it enters the global market.

Is that enough?

It is one of the best things I ever did!

Matteo de Nora
Preface

Tinnitus (ringing in the ears) has many forms, and the severity of tinnitus ranges widely from being a slight nuisance to affecting a person’s daily life. How loud the tinnitus is perceived does not directly relate to how much it distresses the patient. Thus, even tinnitus very close to the hearing threshold can be a disabling symptom that amounts to a major burden, it can reduce the quality of life by generating anxiety and concentration problems impairing the ability to do intellectual work, making it difficult to sleep; causing depression and tinnitus can ultimately lead to suicide. Tinnitus can occur at young age, but its prevalence steadily increases with the degree of age-related hearing loss and can reach 12–15% for people aged 65 and over. Moreover, tinnitus incidence is increasing dramatically with increased leisure noise, more work-related noise trauma, and longer lifespan.

The different forms of tinnitus have similarities with different kinds of pain; many forms of pain and tinnitus are phantom sensations. Another important commonality is that pain and tinnitus lack detectable signs; imaging tests (structural MRI, CT, etc.) and common electrophysiological test results are the same whether or not a person has tinnitus.

For a long time, it was believed that the anatomical location of the physiological abnormalities that caused the tinnitus was the ear. However, it was later understood that most forms of tinnitus are caused by abnormalities in the central nervous system and that these abnormalities are often caused by expression of neural plasticity.

Many structures of the body, such as the ear, the auditory nervous system, the somatosensory system, other parts of the brain, and muscles of the head and the neck are directly or indirectly involved in different forms of tinnitus. To treat and understand the pathology of tinnitus, therefore, requires the involvement of many specialties of medicine, surgery, psychology, and neuroscience.

Tinnitus may occur after noise exposure and administration of pharmacological agents, but the cause of subjective tinnitus is often unknown. Severe tinnitus is often accompanied by symptoms, such as hyperacusis (lowered tolerance to sound) and distortion of sounds. Affective disorders, such as phonophobia (fear of sound) and depression, often occur in individuals with severe tinnitus. With such differences in attributes, it is not reasonable to expect that a single cause can be responsible for severe tinnitus, again a factor that makes managing the tinnitus patient a challenge for health care professionals.

Realizing the complexity of tinnitus has highlighted the importance of interdisciplinary research, and the fact that most forms of tinnitus are disorders of the nervous system has put emphasis on neuroscience, both in studies and in the treatment of tinnitus.
However, few clinicians are specifically trained in tinnitus treatment, and there is a lack of suitable books that describe how to diagnose and treat each of these many forms of tinnitus most effectively.

Each of the authors contributing to the “Textbook of Tinnitus” were, therefore chosen from many specialties of medicine, surgery, psychology, and neuroscience, and came from diverse areas of expertise, such as Neurology, Neurosurgery, Audiology, Otolaryngology, Psychiatry, Clinical- and Experimental Psychology, Pharmacology, Dentistry, and Neuroscience.

Unlike pain, which has considerable literature, including a book with the title “Textbook of Pain” now in its fifth edition, there is no comprehensive book that covers the many aspects of tinnitus. This book, therefore, fills a void by providing relevant information about tinnitus as a disease and how to treat it effectively. The “Textbook of Tinnitus” is directed toward the clinician and gives detailed information about the diagnosis of many different forms of tinnitus and their treatment. The book also provides an overview of what is known about the pathophysiology of different kinds of tinnitus.

It has become more and more evident that neural plasticity plays an important role, not only in adapting the nervous system to changes in demand and after injuries, but also as a cause of symptoms and signs of disease. Such diseases have been called “plasticity disorders.” The role of neural plasticity in creating symptoms of disease, such as many forms of tinnitus, has only been described in a few books directed to neurologists and researchers in neuroscience. This means the medical community in general is often unaware that functional changes in the nervous system can be the cause of a patient’s complaints, and that hampers the diagnosis of disorders, such as tinnitus. Therefore, the effective treatment of tinnitus also requires knowledge about neural plasticity as a cause of diseases. This is one of the aspects of tinnitus that is covered in the “Textbook of Tinnitus.”

The fact that tinnitus is not a single disease, but a group of diseases means tinnitus cannot be effectively treated by a single approach, and several disciplines of health care must be involved in managing the patient with tinnitus. Treatment of the patient with severe tinnitus requires collaborations between clinicians in many different fields of medicine, audiology, and psychology. Accordingly, tinnitus research and treatment have been performed by a variety of disciplines, viewing the problem from various perspectives, focusing on different targets, and using diverse approaches. New developments regarding the treatment have prompted the involvement of neurosurgeons, neurologists, psychiatrists, and dentists. Therefore, an important challenge for the future consists in improving cooperation between different disciplines involved in tinnitus research and treatment.

It is a challenge to translate the results from basic research into clinical practice. The “Textbook of Tinnitus” provides the basis for multidisciplinary management of the tinnitus patient using the most modern methods of treatment. The book represents a new and broad interdisciplinary approach to tinnitus by bringing together in a single book, contributions from many different areas of basic science and clinical research and health care to guide the management of the tinnitus patient. This is the first time that such broad efforts have been made regarding the treatment of tinnitus.

The 95 chapters in this book express the independent views of the authors, some of which may diverge and some may complement one and another. The editors have made no attempts to modify individual authors’ views, only attempts have been made to achieve a similar style of writing in the different chapters.
The book describes both the theoretical background of the different forms of tinnitus and detailed knowledge of state-of-the-art treatment of tinnitus written for clinicians by clinicians and researchers in tinnitus. It provides up-to-date information in forms that are suitable for those who diagnose and treat patients with tinnitus in their clinical praxis as otolaryngologists, neurologists, psychiatrists, neurosurgeons, clinical audiologists, dentists, and psychologists. The book can also serve as a reference for clinicians who do not treat tinnitus patients routinely because of its organization and extensive subject index.

The book has five sections, I Basics about tinnitus, II Causes of tinnitus, III Differential diagnosis of tinnitus, IV Clinical characteristics of different forms of tinnitus, and V Management of tinnitus.

The first section describes the basic aspects of tinnitus and the symptoms that often accompany the disorder, such as hyperacusis and misophonia. This section includes chapters on the epidemiology of tinnitus in children as well as adults and discusses the role of genetics in tinnitus. The anatomy and physiology of the normal auditory system and the pathologic system are the topics of other chapters; chapters on pain and similarities between tinnitus and pain are also included, as are chapters that discuss the use of special forms of neuroimaging for studies of tinnitus. Modeling of the pathologies of tinnitus is the topic of two chapters, and one chapter discusses how clinical trials are performed. The last part of the section concerns how tinnitus is perceived and approached by members of different specialties in the research and treatment of tinnitus, including a chapter about how tinnitus is viewed by the patients themselves.

Section II has chapters about different causes of tinnitus, such as the role of disorders of the ear, age, and exposure to noise and ototoxic substances. Diseases associated with tinnitus, such as vestibular schwannoma and Ménière’s disease, are the topics of other chapters in this section. Yet another chapter covers the cause of somatosensory tinnitus. Other chapters concern the role of different disorders of the central nervous system. The role of disorders of the masticatory system, including that of the temporomandibular joint, is the topic of the last chapter in the section.

Section III discusses the diagnosis of tinnitus and a chapter presents a diagnostic algorithm for tinnitus, followed by chapters on how the different diagnostic methods are performed. Chapters covering otologic, audiologic, and neuro-otologic assessment and examination follow a chapter about history and questionnaires. A chapter describes the diagnosis of somatosensory tinnitus, and another the assessment of temporomandibular disorders. The last chapter in the section covers psychological and psychiatric assessments.

The chapters of Section IV cover the clinical characteristics of the different forms of tinnitus. In order to better meet the need of clinicians, the section is organized according to symptoms and syndromes as presented by the patients. The chapters describe the management of tinnitus with sudden hearing loss, hyperacusis and phonophobia, intermittent tinnitus, and pulsatile tinnitus. Tinnitus that occurs together with other symptoms, such as, Ménière’s disease, headache, and psychiatric disorders (depression, anxiety, and insomnia), are also covered in separate chapters. Finally, posttraumatic tinnitus and tinnitus caused by blast injuries that occur in wars are described.

The chapters of Section V concern management of the various forms of tinnitus. The chapters provide an extensive coverage of the available treatments. The chapters review treatments, such as counseling, cognitive behavioral treatment, and auditory
training, which include various forms of sound stimulation. Specific treatment pro-
grams, such as the Tinnitus Retraining Therapy (TRT) and the Neuromonics program
are described. The chapters also discuss different kinds of pharmacologic treatment.
Treatment using botulinum toxin and different forms of surgical treatment are cov-
ered in separate chapters. Other chapters describe different forms of neuromodula-
tion, and one chapter discusses complementary treatments. The two final chapters
include the treatment of tinnitus and pain and strategies for TMJ disorders as their
topics.

Many of the contributors to “Textbook of Tinnitus” are involved in research spon-
sored by the international research organization, “The Tinnitus Research Initiative”
(TRI). The goal of the TRI is to improve the treatment for tinnitus through advances
in the understanding of the pathophysiology of tinnitus. This organization has pro-
moted collaborative interdisciplinary research on tinnitus during the past 5 years. It
has now been converted into an international research foundation, the TRI
Foundation.

TRI’s goal is to provide a basis for collaborations between researchers and clini-
cians from different fields to achieve an integrated approach to studies of the
pathophysiology of tinnitus and develop and test treatments of different forms of
tinnitus.

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Dallas, February 2010

Aage R. Møller
Berthold Langguth
Dirk De Ridder
Tobias Kleinjung
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Contributors

Umberto Ambrosetti, MD
Department of Specialist Surgical Sciences, University of Milan, Fondazione IRRCCS Ca Granda Ospedale Maggiore Policlinico, Via Pace 9, 20122, Milano, Italy
umberto.ambrosetti@unimi.it

George E. Anthou Esq
132 Greens Ave, Cannonsburg PA, 15317, USA
ganthou@hotmail.com

Moisés A. Arriaga, MD, MBA, FACS
Department of Otolaryngology, Louisiana State University Health Sciences Center, New Orleans, LA, USA
Our Lady of the Lake Hearing and Balance Center, 7777 Hennessy Blvd, Suite 709 Baton Rouge LA, 70808, USA
maa@neurotologic.com

David M. Baguley, BSc MSc MBA PhD
Cambridge University Hospitals, Hills Road, Cambridge, CB2 2QQ, UK
dmb29@cam.ac.uk

Carey Balaban, PhD
Department of Otolaryngology, Eye and Ear Insitute, University of Pittsburgh, 203 Lothrop St, Pittsburgh PA, 15213, USA
cbalaban@pitt.edu

Giovanna Baracca
Fondazione Ascolta e Vivi, via Foppa 15, 20144, Milano, Italy
baracca.giovanna@libero.it

Michael Behr, Dr. med. dent
Department of Prosthodontics, Regensburg University Medical Center, Franz-Josef-Strauss-Allee 11, 93053, Regensburg, Germany
michael.behr@klinik.uni-regensburg.de
Eric C. Bielefeld, PhD, CCC-A
The Ohio State University, 110 Pressey Hall, 1070 Carmack Road, Columbus, OH, 43210, USA
bielefeld.6@osu.edu

Eberhard Biesinger, Dr.med. (PhD)
Department of Klinikum Traunstein, Maxplatz 5, 83278, Traunstein, Germany
Dr.Eberhard.Biesinger@t-online.de

Luca Del Bo
Fondazione Ascolta e Vivi, via Foppa 15, 20144 Milano, Italy
delbo@sordita.it

Daniel J. Bosnyak
Department of Psychology, Neuroscience, and Behavior, McMaster University, Hamilton ON, Canada, L8S4K1
bosnyak@mcmaster.ca

Ralf Bürgers, PhD, DMD
Department of Prosthodontics, University Medical Center Regensburg, Franz-Josef-Strauss-Allee 11, 93053, Regensburg, Germany
ralf.buergers@klinik.uni-regensburg.de

Anthony T. Cacace, PhD
Department of Communication Sciences & Disorders, Wayne State University, 207 Rackham, 60 Farnsworth Detroit MI, 48202, USA
cacacea@wayne.edu

Claudia Barros Coelho, MD, PhD
Rua Mostardeiro, 32/32 Porto Alegre- RS -Brazil, 90430-000
claudiabarroscoelho@gmail.com

Tatjana Crönlein, Dr. phil
Dept of Psychiatry and Psychotherapy, University Hospital of Regensburg, Universtitaetsstr. 84, 93053, Regensburg, Germany	
tatjana.croenlein@medbo.de

Paul B. Davis, PhD MAudSA (CC)
Audiology, Health Professions Division, Nova Southeastern University, 3600 South University Drive, Fort Lauderdale, FL, 33328, USA
pauldavi@nova.edu

Dirk De Ridder, MD, PhD
BRAF1N TRI Tinnitus Clinic & Dept of Neurosurgery, University Hospital Antwerp, Wilrijkstraat 10, 2650, Edegem, Belgium
dirk.de.ridder@uza.be

Isabel Diges, PhD
Department of Otorhinolaryngology, Tinnitus and Hiperacusis Clinic, Hospital Universitario Fundacion Alcorcon, c/ Budapest, 1, 28922
Alcorcon Madrid, Spain
idiges8@gmail.com
Ana Belén Elgoyhen, PhD
University of Buenos Aires, School of Medicine,
National Research Council (CONICET), Institute for Research in Genetic
Engineering and Molecular Biology, Vuelta de Obligado 24, 901428,
Buenos Aires, Argentina
elgoyhen@dna.uba.ar

Paolo Enrico, PhD
Department of Biomedical Sciences, University of Sassari,
V.le S. Pietro 43/B07100, Sassari, Italy
enrico@uniss.it

Stella Forti
Audiology Unit, Fondazione IRRCCS Ca’ Granda Ospedale Maggiore Policlinico,
Via Pace 920122, Milan, Italy
aut_est@yahoo.it

Peter Geisler, MD
Dept of Psychiatry, Psychosomatics and Psychotherapy, University Hospital of
Regensburg, Universitaetsstr. 8493053, Regensburg, Germany
peter.geisler@medbo.de

Vénéra Ghulyan-Bédikian, PhD
106, Bd de Hambourg, 13008, Marseille, France
V_Ghulyan@hotmail.com

Ron Goodey, MD Otolaryngologist
3 Wootton RoadRemuera, Auckland1050, New Zealand
rongoodey@xtra.co.nz

Martin Gosau, MD, DMD
Department of Cranio-Maxillo-Facial Surgery,
University Medical Center Regensburg, Franz-Josef-Strauss-Allee 11, 93053,
Regensburg, Germany
martin.gosau@klinik.uni-regensburg.de

Karoline V. Greimel, PhD
Salzburg University Hospital, Muellner Hauptstasse 48, 5020, Salzburg, Austria
k.greimel@salk.at

Sebastian Hahnel, DMD
University Medical Center Regensburg, Department of Prosthodontics,
Franz-Josef-Strauss-Allee 11, 93053, Regensburg, Germany
sebastian.hahnel@klinik.uni-regensburg.de

Göran Hajak, Dr. med.
Dept of Psychiatry and Psychotherapy, University Hospital of Regensburg,
Universtitaetsstr. 84, 93053, Regensburg, Germany
goeran.hajak@medbo.de

Thomas Hartmann, Dipl.-Psych
Department of Psychology, University of Konstanz, P.O. Box 25, 78457,
Konstanz, Germany
thomas.hartmann@uni-konstanz.de
Carlos Herraiz, MD, PhD
Tinnitus and Hiperacusis Clinic, Department of Otorhinolaryngology, Hospital Universitario Fundacion Alcorcon, c/ Budapest, 128922, Alcorcon, Madrid, Spain cherraizp@seorl.net

Michael E. Hoffer, MD
Department of Otolaryngology, Naval Medical Center San Diego, 34800 Bob Wilson Drive San Diego CA, 92134, USA Michael.hoffer@med.navy.mil

Pawel Jastreboff, PhD, DSc
Department of Otolaryngology, Tinnitus and Hyperacusis Center, Emory University School of Medicine, Atlanta GA, USA pjastre@emory.edu

James A. Kaltenbach, PhD
Department of Neurosciences, NE-63, Lerner Research Institute, Cleveland Clinic, 9500 Euclid Avenue, Cleveland, OH, 44195, USA kaltenj@ccf.org

Eman Khedr, MD
Department of Neurology, Assiut University Hospital, Assiut 71511, Egypt emankhedr99@yahoo.com

Andrea Kleine-Punte, MSci
University Department of Otorhinolaryngology, Head and Neck Surgery, Antwerp University Hospital, University of Antwerp, Wilrijkstr 10, 2650, Edegem-Antwerp, Belgium Andrea.kleine.punte@uza.be

Tobias Kleinjung, MD
Department of Otorhinolaryngology, University Hospital of Regensburg, Regensburg, Germany tobias.kleinjung@klinik.uni-regensburg.de

Michael Koller, PhD
Center of Clinical Studies, University Hospital of Regensburg, Regensburg, Germany michael.koller@klinik.uni-regensburg.de

Orianna Kong MAud (Hons)
The University of Auckland, 92019, Auckland, New Zealand audiology@auckland.ac.nz

Birgit Kröner-Herwig, PhD
Department of Clinical Psychology & Psychotherapy Georg-Elias-Müller-Institute of Psychology, University of Goettingen, Gosslerstr. 14, 37073, Göttingen, Germany bkoene@uni-goettingen.de

Benoit Lafont, MD
Hôpitaux Hopital Nord, 13915, Marseille Cedex 20, France Beloit.Lafont@ap-hm.fr
Michael Landgrebe, MD  
Department of Psychiatry and Psychotherapy, University Hospital of Regensburg, Regensburg, Germany  
michael.landgrebe@medbo.de

Berthold Langguth, MD  
Department of Psychiatry and Psychotherapy, University of Regensburg, Universitätsstraße 84, 93053, Regensburg, Germany  
Berthold.Langguth@medbo.de

Miguel J A Láinez, MD, PhD  
Department of Neurology, University Clinic Hospital, University of Valencia, Avda Blasco Ibáñez 17, 46010, Valencia, Spain  
jlaineza@meditex.es

Edward Lobarinas, PhD, CCC-A  
Department of Communicative Disorders and Sciences, Center for Hearing and Deafness, State University of New York at Buffalo, 137 Cary Hall, 3435 Main StreetBuffalo NY, 14214, USA  
el24@buffalo.edu

Isabel Lorenz, Dipl.-Psych  
Department of Psychology, University of Konstanz, D2578457, Konstanz, Germany  
isabel.lorenz@uni-konstanz.de

Jacques Magnan, MD  
University Aix-Marseille II, Hopital Nord, 13915 Marseille Cedex 20, France  
jmagnan@ap-hm.fr

Jane E Magnusson  
Department of Sport and Exercise Science, The University of Auckland, 92019, Auckland, New Zealand  
j.magnusson@auckland.ac.nz

William Hal Martin, PhD  
Department of Otolaryngology, Oregon Health and Science University, Portland, OR, USA  
martinw@ohsu.edu

Jason G. May, MD  
Department of Otolaryngology – Head and Neck Surgery, School of Medicine, Wayne State University, 4201St Antoine #5E, Detroit, MI, 48201, USA  
jmay@med.wayne.edu

Manuela Mazzoli, MD  
ORL-Otochirurgia, Az. Ospedaliera-Università di Padova, via Giustiniani 2, Padova 35128, Italy  
manuela.mazzoli@gmail.com

Don J. McFerran, MA, FRCS  
Department of Otolaryngology, Colchester Hospital University, NHS Foundation Trust, Lexden Rd., Colchester CO33NB, UK  
donmcferran@aol.com
Olivier Meeus, MD  
Department of Otorhinolaryngology, Head and Neck Surgery, Antwerp University Hospital, University of Antwerp, Wilrijkstr 10, 2650, Edegem-Antwerp, Belgium  
Olivier.meeus@uza.be

Nadia Müller, Dipl. Psych  
University of Konstanz, P.O. Box 25, 78457, Konstanz, Germany  
nadia.mueller@uni-konstanz.de

Aage R. Møller, PhD (DMedSci)  
The University of Texas at Dallas School of Behavioral and Brain Sciences, GR 41, 800 W Campbell Rd, Richardson, TX, 75080, USA  
amoller@utdallas.edu

Matteo De Nora  
Tinnitus Research Initative Foundation, Bezirksklinikum Regensburg, Universitätsstr. 84, 93053, Regensburg, Germany  
Foundation@tinnitusresearch.org

Arnaud Norena, PhD  
Université de Provence, Centre St Charles, Pôle 3C - Case B, 3, Place Victor Hugo F 13331, Marseille Cedex 03, France  
arraud.norena@univ-provence.fr

Michel Paolino, MD  
Centre Médical Clairval, 317,Bd du Redon13009, Marseille, France  
michel.paolino@wanadoo.fr

Anna Piera, MD  
Department of Neurology, University Clinic Hospital, University of Valencia, Avda Blasco Ibáñez 17, 46010, Valencia, Spain  
jlaineza@meditex.es

Alejandro Ponz, MD, PhD  
Department of Neurology, University Clinic Hospital, University of Valencia, Avda Blasco Ibáñez 17, 46010, Valencia, Spain  
jlaineza@meditex.es

Benjamin Questier  
1 Place de l’Eglise, 69270, Saint Romain au Mont d’Or, France  
bquestier@gmail.com

Virginia Ramachandran, AuD  
Division of Audiology, Department of Otolaryngology - Head and Neck Surgery, Henry Ford Hospital, 2799W. Grand Blvd, Detroit MI, 48202, USA  
vramach1@hfhs.org

Charbel Rameh, MD, PhD  
Hopital Nord, 13915, Marseille Cedex 20, France  
charbelramer@hotmail.com
Larry E. Roberts, PhD
Department of Psychology, Neuroscience, and Behavior, McMaster University,
1280 Main Street West Hamilton ON, Canada, L8S4K1
roberts@mcmaster.ca

Carina Andrea Bezerra Rocha,
Rua São Vincente de Paulo, 650/82, São Paulo-SP-Brazil,
01229-010
carinabr.fisio@gmail.com

Carla Vanina Rothlin, PhD
School of Medicine, Yale University,
300 Cedar St TAC S625A, New Haven, CT, 06520, USA
carla.rothlin@yale.edu

Richard Salvi, PhD
Center for Hearing & Deafness, 137 Cary Hall, University of Buffalo,
3435 Main Street Buffalo NY, 14214, USA
salvi@buffalo.edu

Philipp G. Sand, MD
Department of Psychiatry, University of Regensburg, Franz-Josef-Strauss-Allee 11,
Regensburg, Germany
philipp.sand@klinik.uni-regensburg.de

Tanit Ganz Sanchez, MD, PhD
Discipline of Otolaryngology, University of São Paulo School of Medicine, Instituto
Ganz Sanchez, Av Padre Pereira de Andrade, 545/174F, São Paulo-SP-Brazil,
05469-000
tanitsanchez@gmail.com

Winfried Schlee, PhD
University of Konstanz, P.O. Box 25, 78457, Konstanz, Germany
winfried.schlee@uni-konstanz.de

Hannah Schulz, Dipl. Psych
University of Konstanz, P.O. Box 25, 78457, Konstanz, Germany
Hannah.schulz@uni-konstanz.de

Grant D Searchfield, BSc MAud (Hons) PhD (Audiology) MNZAS
Section of Audiology School of Population Health,
The University of Auckland, Auckland, New Zealand
g.searchfield@auckland.ac.nz

Georgina Shakes, BSc (Hons), DClinPsychol, CPsychol
Mt Eden Road, Symonds Street, P.O. Box 8050, Auckland1150, New Zealand
prac92@ihug.co.nz

Susan E Shore, PhD
Departments of Otolaryngology and Molecular and Integrative Physiology,
Kresge Hearing Research Inst, University of Michigan, 1150 West Medical Center Drive, Room 5434A Ann Arbor MI, 48109-5616, USA
sushore@umich.edu
Paul F. Smith, PhD  
Dept. of Pharmacology and Toxicology, School of Medical Sciences, University of Otago Medical School, Dunedin, New Zealand  
paul.smith@stonebow.otago.ac.nz

Wei Sun, PhD  
Center for Hearing and Deafness, 137 Cary Hall,  
University of Buffalo, Buffalo NY, 14214, USA  
weisun@buffalo.edu

Chiemi Tanaka, MA, CCC-A PhD  
Department of Communicative Disorders and Sciences, Center for Hearing and Deafness, State University of New York at Buffalo, 137 Cary Hall, 3435 Main StreetBuffaloNY, 14214, USA  
ctanaka@buffalo.edu

Dayse Távora-Vieira, BSc (Sp Path & Aud) MAudSA (CC)  
University of Western Australia, Perth, Medical Audiologist Services, 51, Colin St, West Perth, WA6005 Australia  
dayse.tavora@gmail.com

Ambrosetti Umberto  
Audiology Unit, Department of Specialist Surgical Sciences, University of Milan, Fondazione IRCCS Ca’ Granda Ospedale Maggiore Policlinico, Via Pace 9, 20122, Milan, Italy  
umberto.ambrosetti@unimi.it

Paul Van de Heyning, MD, PhD  
Department of Otorhinolaryngology Head and Neck Surgery, Antwerp University Hospital, University of Antwerp, Wilrijkstr 10, 2650, Edegem-Antwerp, Belgium  
paul.van.de.heyning@uza.be

Sven Vanneste, MA, MSc  
BRAFiN TRI Tinnitus Clinic and Department of Neurosurgery, University Hospital Antwerp, Wilrijkstraat 10, 2650, Edegem, Belgium  
sven.vanneste@ua.ac.be

Nathan Weisz, Dr. rer. nat  
Department of Psychology, University of Konstanz, P.O. Box 25, 78457, Konstanz, Germany  
nathan.weisz@uni-konstanz.de

Yu-Lan Mary Ying, MD  
Department of Otolaryngology, Baylor College of Medicine, One Baylor Plaza, NA-102 Houston TX, 77030, USA  
ylmying@yahoo.com

Florian Zeman, MA  
Center of Clinical Studies, University Hospital of Regensburg, Regensburg, Germany  
florian.zeman@klinik.uni-regensburg.de
Part I
Basics About Tinnitus
Chapter 1
Introduction

Aage R. Møller

Keywords  Tinnitus • Objective tinnitus • Subjective
           tinnitus • Impact of tinnitus • Treatment • Neural
           plasticity • Hyperacusis • Phonophobia

Abbreviations

CNS   Central nervous system
EEG   Electroencephalography
PAG   Periaqueductal gray

Introduction

Tinnitus can affect the entire life of an individual, can
prevent intellectual work, and impair the quality of life in
general; in some instances, tinnitus can cause suicide.
Severe tinnitus is often accompanied by hyperacusis and
affective disorders such as phonophobia and depression.

Tinnitus and auditory hallucinations are perceptions
of sounds in the absence of external noise. Subjective
 tinnitus and hallucinations are phantom sounds. Tinnitus
is different from hallucinations and objective tinnitus
that is caused by sounds generated in the body and
conducted to the ear. Tinnitus is hearing of meaning-
less sounds. Hallucinations consist of meaningful
sounds such as music or speech and occur in schizo-
phrenia, after intake of certain drugs, and it may occur
(rarely) in temporal lobe disorders. This book will not
cover hallucinations.

There are two main kinds of tinnitus, namely, objective
and subjective tinnitus. Objective tinnitus is caused
by sounds generated in the body and conducted to the
ear. It may be caused by turbulence of blood flow or
muscle contractions. Individuals with subjective tinni-
itus have no visible signs of disease, and the disease has
few detectable physical correlates. Objective tinnitus
may be detected by an observer using auscultation,
whereas subjective tinnitus can only be observed by
the person who has the tinnitus.

Subjective tinnitus can have many forms: it can be
high frequency sounds similar to the sounds of crickets,
like a high- or low-frequency tone, and constant or pul-
satile. Tinnitus can be present at all times or can appear
only sometimes. However, it is usually not possible to
relate a specific event to the appearance of tinnitus.

Patients’ description of their symptoms is the only
cue, and this may be misleading because they point to
the ear, which is rarely the site of the pathology. It is
abnormal neural activity in the brain that causes sub-
jective tinnitus. This abnormal neural activity may
originate in the ear but it is more likely generated
somewhere in the brain.

There are two ways in which abnormal neural
activity that may be interpreted as a sound can occur in
the brain. One is through neural activity in the periphery
of the auditory system that emulates the activity elic-
ted by sound, which reaches the ear. The other way is
through abnormal neural activity generated somewhere
in the ascending auditory pathways. The way the neu-
ral activity that causes tinnitus is generated is not
known in detail, but recent studies indicate that the
activity is different from that elicited by sound stimu-
lation, which means that the different forms of tinnitus
may be generated in different ways.

There is evidence that tinnitus, after some time
(chronic tinnitus), becomes fundamentally different
from acute tinnitus. This change over time is important for treatment of tinnitus, and there is evidence that treatments are less effective after tinnitus has persisted for more than 5 years [1].

Tinnitus is not perceived in the same way as normal physical sounds, and there are indications that the way tinnitus is perceived has to do with perception of “self” (see Chap. 73) [2].

It is not known where in the nervous system sensory activation reaches conscious awareness, and neural activity in other parts of the CNS than that of normal sounds may give rise to the tinnitus sensation. It is not known what features of neural activity are important for eliciting awareness of a sensory signal, and even less is known about which kind of neural activity causes awareness of tinnitus (see Chap. 10) [3].

Contemporary understanding of which qualities of neural activity gives awareness of sensory stimulation includes neural synchrony, coherence of activity in many neurons in cortical or other structures, and neural connectivity. There is considerable evidence that activation of neural plasticity plays an important role in many forms of tinnitus (see Chap. 12). These characteristics of tinnitus have similarities with equally variant forms of pain. In particular, central neuropathic pain has many similarities with severe tinnitus, as will be discussed in this book (Chap. 14). Tinnitus and neuropathic pain are typical examples of “plasticity disorders” [4], where the symptoms are caused by plastic changes that are not beneficial to an individual person.

Sensory awareness and affective reactions (distress) are probably caused by different kinds of neural activity and probably occur in different parts of the CNS. Such separation of perception is known for pain, where the lateral tract of the spinothalamic system produces awareness while the medial system produces the affective and emotional reaction to pain and activates distress networks.

More recently, some abnormal physiological signs have been found to be abnormal in individuals with some forms of tinnitus. One abnormality is with regard to the high-frequency component of electroencephalographic (EEG) recordings, known as gamma activity (see Chap. 21). The amplitude of the gamma activity is increased while the amplitude of another common component of the EEG, the alpha activity, is decreased (see Chap. 17). Animal experiments have shown that some forms of evoked potentials are altered (often increased) after exposure to sounds of an intensity that in humans causes tinnitus, and which has shown signs of hyperactivity in recordings from specific nuclei [5, 6].

The signs of tinnitus at a local anatomical level are often different from those of a global brain level, and there are indications that non-auditory regions of the brain are activated abnormally in some forms of subjective tinnitus (see Chap. 17) [7]. Many different parts of the CNS are involved with tinnitus and there is evidence that parts that normally are not activated by sounds may also be involved in generating the sensation of tinnitus (see Chap. 73).

Also, animal experiments have shown evidence of non-auditory structures, for example, the hippocampus, being involved [8, 9]. Studies in humans have shown evidence of involvement of limbic structures [10]. Other studies have indicated that nonclassical pathways are abnormally involved in some forms of tinnitus [11, 12].

The degree and the impact of tinnitus on an individual person vary widely for the different kinds of tinnitus and also from person to person. It often fluctuates over time and with differing circumstances. Tinnitus is common, but only in a relatively few individuals does it cause distress or other problems. Many people who do not have tinnitus under normal environmental circumstances will experience tinnitus when placed in a room that is silent, such as the test rooms used for audiological testing.

Tinnitus is a phantom sensation of different kinds of sounds, but rarely are these sounds comparable with natural sounds or even with sounds that can be synthesized electronically.

Different methods have been used to estimate the intensity (loudness) of tinnitus. Visual analog scales have been used to estimate the strength of tinnitus, but methods such as loudness balance often give results that are unrealistically low [13]. The results of loudness matching show that most forms of tinnitus have loudness in the range of 20 dB even in situations where the tinnitus is regarded to be unbearable.

The effect of tinnitus on an individual person varies, and the degree of annoyance is not directly related to the perception of tinnitus. Like the impact of severe pain depends on whether it is regarded to be escapable or inescapable, also the impact of tinnitus on a person’s quality of life largely varies. Studies have indicated that inescapable and escapable pain involved different lamina of the PAG [14] and the hypothalamic–midbrain neural circuits [15].
While tinnitus is described as a sound, similar sensations cannot be evoked by sound stimulation and it is assumed that the neural activity that causes tinnitus is different from that evoked by sound stimulation. The abnormal neural activity that causes tinnitus cannot be detected by imaging methods that are available. Some physiological methods can provide some insight in abnormal neural activity, but most of these methods are restricted to use in animals.

Tinnitus, especially severe tinnitus, is often accompanied by abnormal perception of (physical) sounds such as hyperacusis (lowered tolerance for all kinds of sounds) (see Chap. 3) and phonophobia (fear of sound). Hyperacusis also occurs in connection with other diseases such as autism.

In some individuals, tinnitus is associated with distress of affective (emotional) symptoms. These two qualities, perception and distress, are caused by activation of different parts of the nervous system. This is similar to pain where the lateral spinothalamic system is engaged in the perception of pain, whereas the medial spinothalamic system mediates the distress or affective component of pain. Animal experiments have indicated that pain that is perceived as escapable involves anatomically different parts of the periaquaductal gray (PAG) than pain that is perceived as inescapable. It is not known if there are similarities regarding tinnitus.

It is particularly true that when limbic structures (the emotional brain) become activated, tinnitus becomes a problem [2] (see Chaps. 10 and 73).

Treatment of Tinnitus

Subjective tinnitus is the most challenging of common disorders of hearing. So far, the available forms of treatment have had little to moderate success. Many different treatments are in use and even more have been tried and discarded. Often the goal of treatment of severe tinnitus has been to eliminate the symptoms, but this is rarely achieved. However, it is often possible to reduce some of the effects of the tinnitus, so that a patient gains quality of life and would perhaps be able to work in spite of the remaining effects of the disorder. This means that it is often possible to gain quality of life for the patient by such management of the tinnitus. Setting the goal to eliminate tinnitus will often make the patient disappointed when this goal is not met, and the patient may try to find another treatment option, which most likely will be equally disappointing.

There are no known objective tests that can determine the severity of tinnitus and even detect whether tinnitus is present or not. Treatment must therefore rely on the patient’s own assessment of his/her tinnitus. Some functional abnormalities have been detected in some individuals with tinnitus using functional imaging methods that can relate the abnormalities to specific brain regions. However, these methods are still in development and are not yet available for general clinical diagnosis of tinnitus.

Research on tinnitus has lagged behind similar disorders such as pain. There are two kinds of sound perception that are not caused by sounds reaching the ear from outside the body: tinnitus and auditory hallucinations.

Tinnitus Can Occur Together with Other Diseases

Tinnitus may occur together as one of the symptoms of a specific disease, such as Ménière’s disease (see Chaps. 38 and 60), where tinnitus is one of the three (or four) symptoms that define the disease (the others are paroxysmal vertigo and fluctuating low-frequency hearing loss). Vestibular schwannoma are almost always accompanied by tinnitus (see Chap. 39). Individuals with Wilson’s disease often have tinnitus. Tinnitus is often one of the symptoms of intracranial hypotension [16]. Traumatic injuries to the auditory nerve often result in tinnitus. Down’s syndrome may also be associated with a higher incidence of tinnitus than non-Down’s syndrome individuals. It has been reported that autistic individuals have an abnormal perception of loudness [17], but little is known about tinnitus.

Many conditions have tinnitus as part of their symptoms; most noticeable are Ménière’s disease and vestibular schwannoma.

Tinnitus is often associated with hearing loss of various kinds, but hearing loss also occurs without tinnitus. Individuals with tinnitus often have hearing loss, but tinnitus may also occur, although rarely, in individuals with normal or near-normal hearing. In a study by Friedland and co-authors [18], a correlation was found between low-frequency hearing loss and
risk of cardiovascular diseases. These investigators found that the shape of a person’s audiogram correlated strongly with cardiovascular changes and peripheral arterial disease. Hypertension has been found to be associated with a lower incidence of tinnitus, as compared to normotension and hypotension [19].

Tinnitus often occurs after head injuries. Injury to the auditory nerve, which may occur from surgical manipulation or head trauma, often results in tinnitus. Blast injuries, such as those occurring in recent wars, result in a high incidence of tinnitus in connection with closed head injuries.

Tinnitus is more prevalent at old age, but results of epidemiologic studies vary widely, mainly because the criteria for tinnitus chosen in the different studies have been different. Most studies have concerned people who have sought professional help for their tinnitus.

Tinnitus may occur after exposure to loud noise and as complication in treatment with certain drugs such as some antibiotics (ototoxic antibiotics), aspirin, idometacin, and diuretic (furosemide) quinine (see Chap. 42).

Tinnitus often occurs together with depression [20], and it is often said that depression is a co-morbidity to tinnitus. However, it could also be possible that the physiological abnormalities that cause tinnitus are similar or that tinnitus and depression have the same risk factors. Misophonia (dislike of specific sound) may occur together with tinnitus or alone. The “exploding head syndrome” may also occur with tinnitus or alone (see Chap. 4).

**Plastic Changes in the Brain Can Cause Tinnitus**

Tinnitus is regarded to be a complex hyperactive disease, or rather tinnitus is a symptom with complex causes that indicate hyperactive neural activity. There is evidence that the neural activity that causes at least some forms of tinnitus is different from that evoked by sound. Earlier it was assumed that tinnitus was caused by increased firing rate of neurons occurring without sensory input. Recent studies indicate that other forms of abnormal activity somewhere in the nervous system, in particular how neural activity in populations of nerve cells are inter-related, may be the cause of some forms of tinnitus. Evidence has been presented that abnormal synchrony and temporal coherence of the activity in populations of neurons may be the important factors for causing tinnitus [21, 22]. Activation of the nervous systems with temporal (periodic or non-periodic) signals, such as those occurring from sensory stimulation with sounds, creates coherence in the neural activity in a population of neurons because many neurons are activated by the same source. There are reasons to believe abnormal communications between nerve fibers or nerve cells (ephaptic transmission) may be involved in creating an abnormally high degree of temporal coherence of neural activity without any physical sensory input (see Chaps. 10 and 13).

There is considerable evidence that activation of neural plasticity plays an important role in many forms of tinnitus (see Chap. 12). Activation of neural plasticity can alter the connectivity in the brain by unmasking dormant synapses. This is another factor that may be involved in some forms of tinnitus. There is also some evidence that the anatomically located regions activated in tinnitus are different from those that are activated by sound. There are indications that the neural activity that causes the awareness (conscious perception) of tinnitus is different from that which causes the affective (distress) reactions. Such separation in processing of sounds that represent different kinds of information may be similar to the separation of different kinds of sensory signals described as stream segregation. The separation processing that leads to conscious perception and the processing that causes distress may indicate that these occur in different parts of the thalamus: the ventral part for processing of awareness and the medial and dorsal parts for the activity that causes affective symptoms. The dorsal and medial thalamus has subcortical connections to the amygdala. All these forms of changes in the function of the nervous system have few or no detectable morphological correlates.

Many aspects of tinnitus that have lasted a long time (e.g., more than 5 years) are different from tinnitus that has only lasted a short time (less than 5 years). Perhaps most important, tinnitus that has lasted a long time is more difficult to treat than tinnitus that has only lasted a short time [1].

**Impact of Tinnitus on an Individual Person**

The degree and the impact on an individual person from tinnitus vary widely from person to person and often vary over time. Only rarely has it been possible to relate the character and the severity to events or specific diseases.
Introduction

There are no objective tests that can determine the existence of tinnitus nor is it possible to evaluate the severity of tinnitus by any known test. The lack of objective tests may sometimes set the patients’ description into question. The cause (meaning what caused the tinnitus to start) is often elusive. Only rarely has it been possible to relate the character and the severity to events or specific diseases.

The lack of objective signs to classify tinnitus according to severity has affected attempts to study the epidemiology of tinnitus. This is probably the most important reason why different studies typically show different incidence and prevalence values.

References

Chapter 2
Different Forms of Tinnitus

Aage R. Møller

Keypoints

1. Subjective tinnitus has many forms and may be regarded as a group of disorders rather than a single disorder.
2. There are a few objective ways to distinguish between the different forms of tinnitus.
3. Tinnitus has been classified subjectively according to:
   (a) Intensity: Often using a visual analog scale or loudness matching.
   (b) Character: High frequency (like crickets), low frequency (rumbling), tonal, pulsatile, constant, or intermittent.
   (c) Other features such as the ability to modulate the tinnitus by manipulating their jaw, moving their eyes, or applying pressure on neck regions.
   (d) Whether referred to one ear, both ears, or perceived as being inside the head.
4. Some diseases, such as Ménière’s disease, are accompanied with tinnitus; such tinnitus may be different from other forms of tinnitus.
5. Some forms of tinnitus are associated with affective disorders such as depression or phonophobia.
6. Subjective tinnitus is often accompanied by abnormal perception of sounds, known as hyperacusis (lowered tolerance for sounds) or hypersensitivity to sounds.

Keywords
Objective tinnitus • Subjective tinnitus
• Somatosensory tinnitus • Modulation of tinnitus
• Abnormal perception of sounds

Abbreviations
AVM Arterio-venous malformations
EEG Electroencephalography
MEG Magnetoencephalography
TMJ Temporomandibular joint

Introduction

Subjective tinnitus is a broad group of sensations that are caused by abnormal neural activity in the nervous system that is not elicited by sound activation of sensory cells in the cochlea. Subjective tinnitus is by far the most common kind of tinnitus. Subjective tinnitus is phantom sounds that have similarities with the phantom limb symptoms and central neuropathic pain (see Chap. 14) [1, 2].

It is a general problem that the same name (tinnitus) is used for so many different forms of subjective tinnitus with different characteristics, different severities, and different causes. Having the same name used for fundamentally different disorders, such as the different forms of tinnitus, is an obstacle in treatment as well as research. The fact that tinnitus is not a single disorder but many makes epidemiological studies difficult to interpret. Different epidemiological studies have come up with very different numbers for the prevalence of tinnitus to some extent, because different definitions of tinnitus and its severity were employed in different studies.

It is agreed that the incidence of tinnitus increases with age and is more common in people who have had exposure to loud noise. Studies of the prevalence of
tinnitus in individuals above the age of 50 years have shown values from 7.6% to 20.1% (see Chap. 5).

In general, subjective tinnitus has no physical signs, and there are no objective clinical diagnostic tests that can distinguish between the different forms of subjective tinnitus. Only the patient’s own description can serve as a basis for a clinical evaluation. Only recently have laboratory research methods been developed that might provide some insight into the different anatomical locations of the abnormalities associated with different forms of tinnitus. Neuroimaging methods are now beginning to provide some information on the functional changes in the brain of individuals with tinnitus (see Chap. 18). Electrophysiologic tests (electroencephalography, EEG, and magnetoencephalography, also known as MEG) can provide some information about plastic changes in the brain associated with tinnitus (see Chap. 20). These methods may become the basis for future clinical tests that can make a differential diagnosis of the different kinds of tinnitus possible and then relate it to pathology.

**Subjective for Objective Measures of Tinnitus**

Loudness matching and the use of a visual analog scale have been used for estimations of the loudness of an individual’s tinnitus. However, loudness matching results in unrealistically low values [3–5]. The use of a visual analog scale seems to give more realistic values.

In the absence of objective tests, tinnitus has been classified according to its perceived severity. Reed classifies tinnitus into three broad groups: mild tinnitus, moderate tinnitus, and severe chronic tinnitus [3]. Mild tinnitus is defined as tinnitus that does not interfere noticeably with everyday life, moderate tinnitus may cause some annoyance and may be perceived as unpleasant, and severe chronic tinnitus affects a person’s entire life. These classifications rely on the individual person’s own description of their tinnitus. Similar classifications have been used for pain (see [6]).

**The Anatomical Location of the Physiological Abnormality**

Like other phantom sensations, such as phantom limb syndrome, tinnitus is often referred to a different anatomical location than that of the pathology. Since tinnitus has the character of sound, it is often referred to one or both ears. Naturally, tinnitus has been regarded as a pathology located in the ear. Therefore, individuals with tinnitus often seek medical assistance from an ear specialist. However, examination of the ear in most cases finds nothing to be wrong. Also much of the research conducted early had been directed to the ear for studies of the pathology of tinnitus.

The anatomical location of the physiological anomaly of subjective tinnitus is often unknown and is likely to be different from where the tinnitus is referred (one ear, both ears, or in the middle of the head). Instead, the anatomical location of the abnormality that causes tinnitus is the brain. However, it is not obvious which region of the brain the pathology is located, and the abnormal function is not necessarily restricted to regions that are normally activated by sound stimulation.

Many forms of tinnitus are caused by activation of neural plasticity, which makes it difficult to identify the cause and the location of the primary pathology.

Activation of neural plasticity may change many neural processes, re-route information, alter the relation between inhibition and excitation, and change temporal coherence of activity in the population of neurons that may be involved in different forms of tinnitus.

It is possible that different characteristics of tinnitus distinguish the different kinds of tinnitus. There is recent evidence that the pathology of tinnitus that is pulsating is different from tinnitus that is not pulsating (see Chap. 59).

The pathology of tinnitus that is caused by external factors may be different from tinnitus that occurs without external factors being involved.

Deprivation of sensory input may constitute such external factors. It is known to be powerful in turning on neural plasticity, and there are many examples of how restoring input to the auditory nervous system can alleviate tinnitus [7] (see Chaps. 74 and 77). The fact that these methods provide relief from tinnitus supports the hypothesis that neural plasticity has been activated by the absence of signals to the nervous system.

Tinnitus occurs together with age-related hearing loss (see Chap. 36) and noise-induced hearing loss (see Chap. 37), as well as after administration of ototoxic antibiotics, some diuretics (furosemide), and quinine [8].
Different Forms of Tinnitus

Tinnitus caused by noise exposure may normally abate after ending the exposure, but the tinnitus may sometimes remain present after ending exposure and may last indefinitely, which indicates that generation of tinnitus is caused by a stable pathologic state of neural circuits. These neural networks, which generate that kind of tinnitus, have bistable properties: one normal and another pathologic.

Exposure to loud sounds can cause tinnitus (see Chap. 37), and so can administration of ototoxic drugs. It is not known if the cause is the reduction in input to the auditory nervous system that turns neural plasticity on, or if it is overstimulation or possibly the morphological damage from overstimulation that activates neural plasticity.

There is evidence that the pathology of subjective tinnitus that occurs in Ménière’s disease (see Chaps. 38 and 60) is different from other forms of tinnitus because it can be reduced or eliminated by sympathectomy [9], which has not been shown effective in other kinds of tinnitus. Tinnitus in Ménière’s disease may therefore be a specific form of tinnitus that is different from other forms.

Tinnitus almost always occurs together with vestibular schwannoma (earlier known as acoustic tumors) (see Chap. 39). There are reasons to believe that the pathology of these forms of tinnitus is also different, although studies have not been published that could support this hypothesis. It has also been shown that there are other specific differences between the tinnitus that accompanies vestibular schwannoma and other forms of tinnitus. Thus, Cacace (1994) found some specific signs that occurred regarding tinnitus after operations for vestibular schwannoma [10], consisting of gaze-evoked or gaze-modulated tinnitus (see Chap. 39). He ascribed it to a phenomenon of deafferentation-induced plasticity. Acoustic schwannoma is one of the few risk factors for tinnitus that is almost 100%. The tinnitus does not normally disappear after removal of the tumor [11]. Injury of the auditory nerve from trauma, surgical operation, or viral infection (neuritis) is also associated with a high risk of tinnitus.

Traumatic head injuries are often associated with tinnitus.

Subjective tinnitus is often accompanied by an abnormal perception of sounds, such as hyperacusis (decreased tolerance for sounds in general, see Chap. 3), phonophobia (fear of sound), and misophonia (dislike of specific sounds) (see Chap. 4). Some individuals with tinnitus hear sounds as being distorted, spoiling the enjoyment of music. This distortion may also make it difficult to understand speech.

Many individuals who have tinnitus (about two-thirds) can modulate their tinnitus by signals from the somatosensory system, such as from eye movements [16], manipulations of their jaw, and applying various pressure on specific neck regions [17–19]. These forms of tinnitus can be managed by somatosensory-oriented treatment [20], and such individuals may be a subgroup with a different pathology.

Affective symptoms accompany some forms of tinnitus [21]. It seems likely that such forms of tinnitus are different from other forms and that their pathology may differ as well (see Chap. 62).

Conclusion

Tinnitus is not a single disorder and the symptoms vary substantially. The causes of different individual’s tinnitus also have wide variants. The fact that a disorder with such differences has the same name is an obstacle in studies of tinnitus and patient management.
References

Chapter 3
Hyperacusis and Disorders of Loudness Perception

David M. Baguley and Don J. McFerran

Keypoints

1. There are several forms of loudness perception disorder.
2. The terminology of such disorders is often confused.
3. The most commonly used terms in an audiological context are hyperacusis, denoting a generalized reduced tolerance for sound, as well as phonophobia, denoting a fear of sounds.
4. The majority of people with a loudness perception disorder also have tinnitus. Just under one half of individuals with tinnitus also describe some degree of loudness perception disorder.
5. There are few rigorous studies regarding the epidemiology of loudness perception disorders; the true prevalence of hyperacusis and phonophobia remains a matter of conjecture.
6. Some loudness perception disorders are associated with disorders of facial nerve function with consequent loss of the acoustic reflex. Most cases have no such association and the underlying pathological mechanism is unclear.
7. Various management strategies have been suggested, including the use of tinnitus therapies, with or without the use of sound therapy, and psychological therapies.

Keywords Tinnitus • Hyperacusis • Hypersensitivity • Loudness discomfort • Migraine

Introduction

Most people dislike certain sounds, irrespective of their intensity; chalk screeching on a blackboard or the sound of skin catching on a child’s balloon are common examples of this. Many people recognize that their sound tolerance varies with their mood, so that someone who is tired, stressed, or anxious may find sounds within their normal tolerance zone unpleasantly loud. Similarly, one person’s unbearably loud concert may be another’s ideal outing. Because of this interpersonal and temporal variation, clinical disorders of sound tolerance were not recognized until relatively recently, and even when recognized, were thought to be exceptional. In 1987, Vernon [1] stated “In our Tinnitus Clinic, where more than 4,000 patients have been seen, hyperacusis has been seen only four times.” As knowledge of tinnitus has improved, recognition of disorders of loudness tolerance has also improved. However, this is still a confused and under researched area.

The fact that clinical recognition of disorders of sound tolerance is relatively recent is not to say that these issues have arisen in modern times. For example, Wilkie Collins uses hyperacusis as an essential element of the plot in his gripping novel “the Woman in White” (1860). Mr Fairlie is the uncle and guardian of Laura, and is derelict in his duty (leading to his niece’s downfall) as he is unable to tolerate spoken conversation and thus advise her. For example, Mr Fairlie states:

“Pray excuse me, but could you contrive to speak in a lower key? In the wretched state of my nerves, loud sound of any kind is indescribable torture to me You will pardon an invalid?” (see also Chap. 57).
Definitions

There is still no unified standard of nomenclature for disorders of sound tolerance. Some of the commonly used words are shown in Table 3.1.

Part of the reason for this wide range of terminology is that disorders of sound tolerance are treated by several disciplines. As well as Audiology and Otolaryngology, Neurologists and Psychiatrists encounter patients with symptoms of altered sound tolerance and, hence, have developed their own terminology. Sound tolerance is also an important consideration for those involved in the public health issues of environmental and occupational noise.

Hyperacusis

The word “hyperacusis” first appeared in the medical literature in 1938 [2]. A later modification to “hyperacusis dolorosa” [3] captured the emotional impact but was not widely adopted. The dictionary definition given in Table 3.1 implies the ability to detect sound at abnormally low intensities, or, in other words, better than average hearing: this is not how the term is used in the clinical literature. Subsequent attempts to define hyperacusis have included “unusual tolerance to ordinary environmental sounds” [1], “consistently exaggerated or inappropriate responses that are neither threatening nor uncomfortably loud to a normal person” [4], and “abnormal lowered tolerance to sound” [5]. A more recent definition [6] describes hyperacusis as “abnormal increased sound-induced activity within the auditory pathways”. As a result, sounds that are nonintrusive, or unnoticed by the general population, are uncomfortable to people with hyperacusis. The common thread to all these definitions is that sounds in general, rather than specific sounds, are unpleasant to individuals with hyperacusis.

Some workers have applied a different meaning to hyperacusis. Gordon [7] defined it as increased sensitivity to quiet sounds or, in other words, unusually acute hearing and coined the term “audiosensitivity” for what audiologists would regard as hyperacusis. The word “audiosensitivity” is not commonly used. To further complicate matters, a new term has recently appeared, namely, “conductive hyperacusis” [8, 9].

Table 3.1  Some of the words and phrases used to describe disorders of loudness tolerance

<table>
<thead>
<tr>
<th>Synonyms</th>
<th>Derivation</th>
<th>Definition</th>
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<tbody>
<tr>
<td>Recruitment</td>
<td>Loudness recruitment</td>
<td>Fr recruter</td>
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<tr>
<td>Hyperacusis</td>
<td>Hyperacousia Hyperacusia Hyperacusis Acoustic hyperaesthesia Auditory hyperaesthesia</td>
<td>Gk hyper (above) akousis (hearing)</td>
</tr>
<tr>
<td>Phonophobia</td>
<td></td>
<td>Gk phone (voice or sound) phobia (fear)</td>
</tr>
<tr>
<td>Misophonia</td>
<td></td>
<td>Gk misos (hatred) phone (voice or sound)</td>
</tr>
<tr>
<td>Dysacousis</td>
<td>Auditory dyesthesia dysacousia dysacusis</td>
<td>Gk dys (bad) akousis (hearing)</td>
</tr>
<tr>
<td>Odynacusis</td>
<td></td>
<td>Gk odyne (pain) akousis (hearing)</td>
</tr>
<tr>
<td>Auditory alldynia</td>
<td></td>
<td>L auditorius (pertaining to hearing) Gk allos (other) odyne (pain)</td>
</tr>
<tr>
<td>Collapsed sound tolerance</td>
<td></td>
<td>Term coined by the Hyperacusis Network (<a href="http://www.hyperacusis.net">www.hyperacusis.net</a>) and synonymous with hyperacusis</td>
</tr>
</tbody>
</table>
This is a phenomenon associated with dehiscence of the superior semicircular canal in which the person may have normal air conduction thresholds on pure tone audiometry but better-than-normal bone conduction. This results in an air-bone gap, and the person often complains of hyper-awareness of somatosounds.

**Phonophobia**

Phonophobia, literally meaning fear of sound, is a widely used term in neurology, particularly in association with migraine. Woodhouse and Drummond [10] reported that at least 50% of migraine attacks are accompanied by increased sensitivity to sound, and uncomfortable loudness levels are reduced during attacks. From an audiological point of view, however, phonophobia implies reaction to certain sounds that have specific emotional associations for that person. Thus, the reduced sound tolerance seen in migraine might be better described as hyperacusis. True phonophobia in isolation is unusual; Hazell et al. [11] reported that only 56% of individuals with reduced loudness tolerance had pure phonophobia.

**Misophonia**

The condition misophonia (see also Chap. 4) was proposed in 2003 [12] to convey many of the same sentiments as phonophobia but removing the phobic connotation as an automatic accompaniment. This is potentially useful as in some health economies it is not lawful to treat a phobic condition unless one is a licensed psychologist or psychiatrist. In 2004, Jastreboff and Hazell [6] describe misophonia as “a negative reaction to sound results from an enhanced limbic and autonomic response, without abnormal enhancement of the auditory system.” They suggest that phonophobia is a subsection of misophonia where fear is the chief component. The word “misophonia” has yet to enter widespread usage and is not a recognized term in many healthcare databases such as Medline. It does, however, add a useful definition to the terminology of reduced loudness tolerance, and its usage should probably be encouraged.

**Recruitment**

Recruitment or, to use the full title, loudness recruitment [13, 14], is a common finding in individuals with cochlear hearing loss associated with outer hair cell dysfunction. It is characterized by an abnormally large increase in the perceived loudness of a sound caused by a slight increase in its intensity. This is not modulated by mood or levels of anxiety.

The boundaries between these definitions can occasionally seem blurred, and it is also quite possible for a person to have more than one form of reduced sound tolerance. For example, a person with a cochlear hearing loss may display recruitment but also have phonophobia. In the Audiology/Otology literature, hyperacusis is quite frequently used as an all-embracing term for all forms of reduced sound tolerance, adding to the confusion. Additionally, present terminology does not describe some clinical presentations, such as the hyper-vigilance to novel auditory events seen in individuals with autism spectrum disorder [15, 16] or the marked auditory startle seen in some with cerebral palsy [17].

**Acoustic Shock**

Recently, considerable interest has been developed about auditory symptoms arising in response to sudden, unexpected sounds [5, 18] (see also Chap. 4). The causative signal does not have to be particularly loud and does not reach a level that causes noise-induced hearing loss. The phenomenon has developed particular relevance among people wearing headsets or using telephone handsets in working environments such as call centers. Acoustic shock undoubtedly predates call centers, and wearing a headset is probably not essential to the diagnosis. Almost all affected individuals describe pain in or around their ears. Other symptoms include tinnitus, vestibular disturbance, hyperacusis, hyper-vigilance, anxiety, headache, numbness, burning, tingling, blockage, pressure, fullness, echoing, or hollow feelings in the ear. Much remains to be discovered about the character of the sounds that trigger this condition, the characteristics of the individuals who develop the symptoms, and the correct methods of managing the disorder. The pathogenesis of the condition has included theories of overactivity of the tensor tympani muscle, cochlear damage, central auditory mechanisms,
or a post-traumatic stress disorder. A UK working group has been set up, “The Acoustic Shock Programme” and has proposed the following definitions for those individuals who develop acoustic shock in the workplace while using communications equipment:

- An acoustic incident is a sudden, unexpected, noise event which is perceived as loud, transmitted through a telephone or headset.
- Acoustic shock is an adverse response to an acoustic incident resulting in alteration of auditory function.

**Epidemiology of Reduced Loudness Tolerance**

There is still a dearth of published work on the demographics of reduced sound tolerance, but it certainly seems more common than Vernon’s original observation [1]. A Polish study into the prevalence of tinnitus [19] included a question on hyperacusis; 10,349 people responded, of whom 15.2% reported hyperacusis. The symptom was more common in men, more common in those of higher socio-economic class, and more common in urban dwellers. Among individuals who had tinnitus, the prevalence of hyperacusis was 40%. A well-designed study by Andersson et al. [20] examined responses to a questionnaire administered partly over the internet and partly via the conventional postal system. This showed a point prevalence of hyperacusis of 9% in the web respondents and 8% in the postal group. The prevalence of requiring ear protection for everyday sounds was notably lower at 2 and 3% for the web and postal respondents, respectively. Interestingly, a proportion of respondents also reported sensitivity to other sensory modalities, particularly light and odours, and this increased sensitivity was higher in the respondents who also reported sound sensitivity. In addition, those who reported hyperacusis were also more likely to report dizziness, hearing loss, and headaches. These estimates of the prevalence of hyperacusis do not make a distinction between people who have a mild dislike of extremely loud sounds and those whose sound tolerance has a significant impact on their ability to live a normal life. Consequently, they almost certainly over-estimate the number of people who have clinically important hyperacusis (see also Chaps. 5 and 6).

One way to obtain an approximate prevalence figure for significant hyperacusis is to use a process of extrapolation from other data sources. The prevalence of hyperacusis in individuals who have tinnitus and vice versa has been well documented. Patients attending a tinnitus clinic have a hyperacusis prevalence of approximately 40% [21, 22]. Among those whose chief complaint is hyperacusis, the prevalence of tinnitus has been reported as 86% [23]. If 5% of the adult population have troublesome tinnitus [24], and 40% of those have troublesome hyperacusis, then a prevalence of significant hyperacusis of 2% can be derived [5].

Altered loudness tolerance is seen in conjunction with several other common conditions, most notably migraine [25] and post-traumatic stress disorder [26]. It is also thought to be more common in conditions such as depression, though there is little robust scientific support for this assertion.

**Pathophysiology of Reduced Loudness Tolerance**

A review by Katzenell and Segal [27] separated disorders of loudness tolerance into those associated with conditions of the peripheral auditory system, diseases of the central nervous system, hormonal diseases, and infectious diseases. However, they also concluded that in many cases, there was no identifiable cause, and that in these cases, the central auditory system was the likely culprit. The peripheral causes discussed by Katzenell and Segal [27] included Bell’s palsy, Ramsay Hunt syndrome, and individuals who have undergone a stapedectomy. However, in all these cases, the stapedial reflex might have been affected, either due to direct damage to the stapedius muscle (stapedectomy) or due to damage to the facial nerve that innervates stapedius (Bell’s palsy, Ramsay Hunt syndrome). Without a functioning stapedius, part of the ear’s protective reflex is lost and more sound energy can reach the cochlea. As the auditory system is then responding correctly to the amount of energy reaching the cochlea, it is a moot point as to whether this constitutes a true abnormality of loudness tolerance.

Peripheral causes of hyperacusis, however, are relatively uncommon. The majority have no obvious cause, but a number of cases of hyperacusis are associated with specific conditions; these examples of so-called
“syndromic hyperacusis” are shown in Table 3.2. Because the underlying pathology for some of these conditions is at least partially understood, it is useful to examine their pathological mechanisms to try and obtain clues about hyperacusis in general.

Some cases of familial migraine have been shown to be associated with mutations in a central nervous system calcium gene. It has been speculated that if this faulty gene is present, calcium channels within the cochlea or central auditory pathways could be involved, resulting in the episodic hypersensitivity to sound [28].

Lyme disease is a tick borne infection caused by a bacterium, *Borrelia burgdorferi*. The infection affects many organs including the nervous system, and hyper-sensitivity to sound is a well-recognized symptom [29]. In some cases, the facial nerve is affected. Hence, the stapedial reflex may be deficient, resulting in the same mechanism described above for Bell’s palsy. However, there are also cases where the facial nerve function and stapedial reflex are normal and, in these cases at least, the problem is likely to be in the central auditory system.

Several of the other conditions associated with central hyperacusis, such as depression, migraine, posttraumatic stress disorder, and posthead injury syndrome, are thought to be related to disturbances of 5 hydroxytryptamine (5 HT, serotonin) function [18, 27]. 5 HT is known to be involved in central auditory pathways. It has been suggested that the hyperacusis is a manifestation of this disturbed 5 HT function [30].

Williams syndrome is a rare chromosomal abnormality caused by deletion of part of chromosome 7, which includes the Elastin Gene. Affected individuals have characteristic elfin features, developmental delay, cardiac problems, and hypercalcaemia. Diagnosis is accomplished by detecting the abnormal gene sequence using fluorescence in situ hybridization (FISH test). Traditionally, it has been thought that at least 90% of individuals with William’s syndrome experience hyperacusis; and for all intents and purposes, that symptom has been regarded as a defining characteristic of the condition (Table 3.3). It is interesting, however, to note that when a validated questionnaire is used, as in the Blomberg et al. study [31], the prevalence of hyperacusis falls, and it may be that what is experienced in Williams syndrome is an aversion to all sound rather than an abnormality of loudness tolerance. Marriage and Barnes [30] suggested that the hyperacusis of Williams syndrome is another example of hyperacusis, secondary to problems of 5 HT function. This theory is yet to be proved.

There are several theories as to the cause of nonsyndromic hyperacusis. The medial efferent part of the central auditory system sends neurons to the outer hair cells; it is thought that these modulate the cochlea’s response to sound [32]. Thus, a defect of the medial efferent system might lead to reduced damping of the cochlea. Sahley and Nodar [33] suggested that stress causes the release of endogenous opiates or dynorphins under the inner hair cells. This could potentiate the cochlear neurotransmitter glutamate, which might lead to enhanced auditory nerve activity.

The neurophysiological model supplies possible mechanisms for both hyperacusis and misophonia (including phonophobia) [6]. In hyperacusis, the incoming auditory signal undergoes a process of abnormal enhancement or amplification in subconscious auditory pathways. This then causes secondary activation of the limbic system and autonomic nervous system. The mechanism by which the incoming signal is enhanced is obscure. In misophonia, the auditory

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**Table 3.2** Conditions associated with reduced loudness tolerance

<table>
<thead>
<tr>
<th>Condition</th>
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<tbody>
<tr>
<td>Migraine</td>
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<tr>
<td>Depression</td>
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<tr>
<td>Posttraumatic stress disorder</td>
</tr>
<tr>
<td>Head injury</td>
</tr>
<tr>
<td>Lyme disease</td>
</tr>
<tr>
<td>Williams syndrome</td>
</tr>
<tr>
<td>Multiple sclerosis</td>
</tr>
<tr>
<td>Addison’s disease</td>
</tr>
<tr>
<td>Fibromyalgia/pain</td>
</tr>
<tr>
<td>Multiple sclerosis (case report)</td>
</tr>
<tr>
<td>Middle cerebral artery aneurysm (case report)</td>
</tr>
</tbody>
</table>

**Table 3.3** Hyperacusis in Williams syndrome

<table>
<thead>
<tr>
<th>References</th>
<th>Percentage with hyperacusis</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Klein et al. (1990) [4]</td>
<td>65</td>
<td>95</td>
</tr>
<tr>
<td>Van Borsel et al. (1997) [52]</td>
<td>82</td>
<td>95</td>
</tr>
<tr>
<td>Levitin et al. (2005) [51]</td>
<td>118</td>
<td>91</td>
</tr>
<tr>
<td>Blomberg et al. [31]</td>
<td>38</td>
<td>13</td>
</tr>
</tbody>
</table>

References

Klein et al. (1990) [4]
Van Borsel et al. (1997) [52]
Levitin et al. (2005) [51]
Blomberg et al. [31]
pathways behave normally but the limbic and autonomic nervous systems are in a heightened state of excitation and therefore react abnormally to normal auditory input.

An elegant piece of work by Formby et al. [34] examined auditory plasticity by allocating normal hearing volunteers to wear either sound attenuating earplugs or sound generators for a 2-week period. Wearing ear plugs resulted in the participants reporting increased loudness perception, whereas wearing sound generators resulted in increased sound tolerance. This experiment supports the hypothesis that loudness perception is directly related to central auditory gain and not only demonstrates the plasticity of the auditory system but also provides support for the use of sound therapy in the management of loudness perception disorders (see also Chap. 13).

Baguley and Andersson [5] attempted to incorporate beliefs and thoughts about the effects of noise and listening situations, in which discomfort is experienced with a model explored in the literature on pain, called the “fear avoidance model” [35, 36], originally developed by Lethem et al. [37]. The central concept of the model is fear of pain, with varying degrees of severity from just plain pain to exacerbation of pain following exposure. The two end-points in the process are either “confrontation” or “avoidance,” with the latter leading to maintained avoidance and possibly even a phobic state. The observation from the pain literature that fear of pain can serve a causal role in leading to disability (fear of injury leads to inactivity, and that inactivity in itself leads to even more pain and disability) is relevant for hyperacusis as well. In light of the experimental evidence recently provided by Formby et al. [34] that ear protection leads to increased noise sensitivity, it was postulated that avoidance of auditory stimulation is likely to sensitize the auditory system, which in turn can exacerbate the hyperacusis. In a recent book on tinnitus [24], a three-component understanding of hyperacusis was proposed that involved consideration of sensitivity, annoyance, and fear of injury (Fig. 3.1). While the first two of these factors have been extensively researched in the literature on noise sensitivity [38], fear of the pain experience in itself, the risk of becoming hearing impaired, getting worse tinnitus, and so on, might be a further factor that plays a significant role in explaining the avoidance of sounds in hyperacusis. It is too narrow to just focus on fear of injury, however; and in a slightly revised version of the three-component model, “fear” is considered a factor which is a more broad construct including fear of the actual pain experience when noise is confronted. The noise sensitivity refers to the actual sensation of pain that is an aversive reaction not necessarily involving cognitive appraisal. The annoyance/irritation dimension is similar to the construct proposed more recently by Jastreboff and Hazell [6] – misophonia – and is more closely linked to cognitive appraisal. What is left out in the figure and in the discussions on hyperacusis overall is the possible effect of noise sensitivity on cognitive capacity, such as attention and memory performance. Moreover, the link between noise exposure and stress responses, including cardiovascular responses, sleep, etc., has largely been ignored in the literature on hyperacusis.

Engel [39] proposed that disease should be considered within a biopsychosocial framework, and this idea has come to underpin much of health psychology. The biological element is the pathophysiology of the condition, the psychological element, the emotional and behavioural impact, and the social element – initially society’s view of a symptom/condition – but more literally taken to mean the social consequences (e.g., work or respect) of that state. Baguley and Andersson [5] considered hyperacusis within a biopsychosocial framework, arguing that an exclusive focus upon only one of these elements (such as the pathophysiology of hyperacusis) is not helpful.

A novel perspective of hyperacusis has been provided by Dubal and Viaud-Delmon [40], who sought to determine if an association exists between hyperacusis and magical ideation (the latter concept describing “nonrational” beliefs about the world), it may be a model of the distorted cognitions in psychosis. Using the Khalfa et al. [41] questionnaire (see below) to assess hyperacusis, Dubal and Viaud-Delaman demonstrated an association, proposing that magical ideation might give a predisposition for heightened auditory sensitivity. More work in this area is awaited with interest.
Assessment

Clinical History

The diagnosis of hyperacusis is essentially made by taking a detailed clinical history, which will often take considerable time to elucidate. A structured framework for a hyperacusis history has been proposed by Baguley and Andersson [5] (Table 3.4).

Examination

Clinical examination of patients with hyperacusis is frequently normal and does not contribute to the diagnosis or direct subsequent treatment. Nevertheless, there are occasions when the examination may supply useful information and it is important that all patients have the reassurance that a thorough assessment provides. In addition to examination of the ears, particular attention should be given to cranial nerve function as a small but significant number of hyperacusis patients have disorders of cranial nerve function, in particular, disorders of the facial nerve.

Audiometry

All audiometric testing in hyperacusis patients must be undertaken with the utmost care. Exposing people to the sensory stimulus that distresses them runs the risk of increasing that distress and exacerbating the situation rather than helping. Many patients require nothing more than a pure tone audiogram. Even this can prove upsetting for some patients, and the initial presentation of the test tone may need to be at a much lower level than normally used. The use of loudness discomfort levels (LDLs) has been advocated for this patient population [6], both as an aid to diagnosis and as an outcome measure. This is not without issue, as there is marked inter- and intra-participant variability in results [42, 43], evident in both tonal and speech-based protocols [5]. Further, subjecting the patient to sound, either at or above their threshold of discomfort, may undermine clinical rapport and therapeutic trust. Arguments for undertaking the test are the need to determine the extent of the problem for individuals, perhaps most pertinently in a medico-legal context, and to determine the efficacy of therapy. There is no consensus on this issue at present. Other tests that use sound stimuli that are likely to exceed the threshold for discomfort should be avoided. These include stapedial reflex estimation and evoked response audiometry.

Table 3.4 Structured diagnostic hyperacusis interview [5]

<table>
<thead>
<tr>
<th>Background questions</th>
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<tbody>
<tr>
<td>1. Family situation</td>
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<tr>
<td>2. Work situation</td>
</tr>
<tr>
<td>– Current or previous work history?</td>
</tr>
<tr>
<td>3. Sick leave</td>
</tr>
<tr>
<td>– On extended sick-leave? (for the last six months for example)</td>
</tr>
<tr>
<td>– Part-time or full time?</td>
</tr>
<tr>
<td>Noise sensitivity questions</td>
</tr>
<tr>
<td>4. Onset of noise sensitivity?</td>
</tr>
<tr>
<td>5. Gradual or sudden?</td>
</tr>
<tr>
<td>6. Development over time (worse–better)?</td>
</tr>
<tr>
<td>7. Laterality?</td>
</tr>
<tr>
<td>8. Type of sounds? (e.g., clatter, talk, paper noises etc)</td>
</tr>
<tr>
<td>9. Perception of sounds being unclear/distorted? If so, what kind of sounds?</td>
</tr>
<tr>
<td>11. Hearing impairment and related compensations (e.g., hearing aids)?</td>
</tr>
<tr>
<td>12. Tinnitus and related distress?</td>
</tr>
<tr>
<td>13. What is most bothersome? Hearing loss, tinnitus, or the hyperacusis?</td>
</tr>
<tr>
<td>14. Does exposure to loud sounds increase the sensitivity?</td>
</tr>
<tr>
<td>Other diagnoses and medical history of relevance</td>
</tr>
<tr>
<td>15. Episodes of depression? If yes, how many episodes in life?</td>
</tr>
<tr>
<td>16. Any contact with psychiatry?</td>
</tr>
<tr>
<td>17. Migraine?</td>
</tr>
<tr>
<td>18. Tension headache?</td>
</tr>
<tr>
<td>19. Other sensitivities and medical problems?</td>
</tr>
<tr>
<td>(a) Light</td>
</tr>
<tr>
<td>(b) Touch</td>
</tr>
<tr>
<td>(c) Pain</td>
</tr>
<tr>
<td>(d) Smell</td>
</tr>
<tr>
<td>(e) Allergy</td>
</tr>
<tr>
<td>(f) Balance disturbance</td>
</tr>
<tr>
<td>20. Whiplash?</td>
</tr>
<tr>
<td>21. Temporomandibular joint dysfunction or problems with teeth?</td>
</tr>
<tr>
<td>22. Hypertension or other cardiovascular issues?</td>
</tr>
<tr>
<td>23. Medications?</td>
</tr>
<tr>
<td>24. Avoidance of places and activities because of hyperacusis?</td>
</tr>
<tr>
<td>(a) Things not done/stopped because of hyperacusis?</td>
</tr>
<tr>
<td>(b) Thing not done yet in life and now very unlikely/impossible because of hyperacusis?</td>
</tr>
<tr>
<td>(c) Use of ear protection? What kind, when and where?</td>
</tr>
</tbody>
</table>
Imaging

Patients with hyperacusis may require imaging, with similar indications as for tinnitus. These indications include asymmetric symptoms, asymmetric audiometric findings, or associated neurological symptoms or signs. Care should be taken when choosing the imaging modality. Magnetic resonance imaging (MRI) is the usual modality of choice for investigating the cerebellopontine angles, but it produces considerable sound levels. Although ear defenders are routinely employed when performing MRI scans, these may not attenuate the sound sufficiently to make it a comfortable experience for a hyperacusis patient. Consideration may need to be given to using a quieter, less-sensitive modality such as computed tomography (CT).

Self-Report

Until recently, there was no method of recording the impact that reduced sound tolerance had on individuals. However, this has been addressed and there are now two self-report questionnaires. Khalfa et al. [41] devised a questionnaire based on 14 items and normalized on 201 volunteers from the general population. A three-factor solution of attentional, emotional, and social factors was derived. In a clinical setting, this questionnaire has low negative impact on patients, but the fact that of the sample population only 4 or 5 might have been expected (from the data above) to have clinically significant hyperacusis. Nelting et al. [44] developed a similar tool using 27 items on 226 individuals with a complaint of hyperacusis. A three-factor solution was derived, with cognitive reactions, actional or somatic behaviour, and emotional factors identified. This latter questionnaire is currently only available in German, but hopefully the arrival of these questionnaires marks a step forward in the study of reduced sound tolerance.

An alternative approach to the scaling of the severity of hyperacusis was proposed by Dauman and Bouscau-Faure [45], who formulated a multiple activity scale for hyperacusis (MASH) which measures the impact of the symptom upon everyday activities. Whilst this undoubtedly captures an aspect of the disability associated with hyperacusis, in that patients may describe a major impact upon family activities such as supermarket shopping or cinema attendance, it should be noted that what is a commonplace activity for an individual in one culture may be exceptional in another context.

Management

The first response of many patients and clinicians is to try and escape from sound, whether by moving to a naturally quiet environment or by using sound attenuating devices such as ear plugs or earmuffs. Unfortunately, observation would suggest that this generally makes the situation worse. It is thought that by reducing the expected sound input, the central auditory gain is increased which exacerbates the loudness hypersensitivity. It is important to explain this carefully to patients, as to many it seems counter-intuitive to expose someone who is distressed by noise to the very thing that is causing the distress. If their job or recreation involves noise exposure, they may need appropriate sound protection devices such as general earmuffs and plugs or musician’s, shooter’s, or motorcyclist’s plugs. It must be carefully stressed that these are only to be worn when the noise levels are genuinely high and must not be worn at other times.

Any associated condition such as Lyme disease, depression, or post-traumatic stress disorder should be treated by the appropriate clinical teams at the same time as the loudness perception is addressed.

Sound therapy is widely used in the treatment of hyperacusis, either as a stand-alone treatment modality or as part of Tinnitus Retraining Therapy (TRT). Using therapy on its own can be undertaken using techniques of either desensitization or recalibration. With decalibration, sound generators are set to levels just below the patient’s threshold of discomfort and slowly increased as tolerance improves. There is some support for this method [46], though in practice relatively few clinicians use this approach. For recalibration, the devices are set to a comfortable, consistent level with the intention of resetting the central auditory gain in much the same fashion that Formby et al. [34] demonstrated with normal volunteers. Norena and Chery-Croze [47] used what they described as an “enriched auditory environment” in patients with hyperacusis. The stimuli were used for several hours a day (for 15 weeks) and consisted...
of pure tone stimuli (100-ms duration and 100-ms intervals) within the audiometric area affected by hearing loss. This meant that the sound stimulation was given at the same frequency range as the hearing loss: i.e., if the hearing loss was in the region 3–6 kHz, sound stimulation of 3–6 kHz was administered.

The sound stimulation was achieved by listening to a CD, and significant improvements in measures of loudness scaling were reported. Due to the small number of patients and some of their characteristics (the presence of hearing loss being an example), this work should be regarded as preliminary but of major interest. For sound therapy for hyperacusis in general, there is a paucity of robust evidence of efficacy to date.

TRT was introduced by Jastreboff and Hazell (1993) [6, 53] as a novel method of dealing with tinnitus. It was recognized that this could be adapted slightly and applied to patients with reduced sound tolerance. In TRT, the first therapeutic step is to allocate each new patient to a category. Category 0 patients have mild or recent onset symptoms. Category 1 and 2 patients have significant tinnitus with normal hearing or hearing loss, respectively. Category 3 patients have hyperacusis without prolonged enhancement from sound exposure. Category 4 patients have tinnitus and/or hyperacusis with prolonged worsening of the symptoms following sound exposure. Those patients with significant hyperacusis, namely category 3 and 4 patients, receive counselling and treatment with wearable binaural sound generators. The protocol for wearing the sound generators varies according to the category. In category 3, patients are advised to slowly increase the sound to the highest level that does not cause annoyance or discomfort or interfere with the hearing. The generators should be worn continuously and may need frequent adjustment. In category 4 patients, the generator output should be set close to threshold or even put in the ear but not initially switched on. The output is then slowly increased, changing the level every 6–8 weeks. If the patients have tinnitus as well as decreased loudness tolerance, the loudness tolerance should be addressed first even if this runs the risk of temporarily worsening the tinnitus. There are a limited number of trials showing the outcome of TRT in the treatment of hyperacusis and all have methodological flaws. However, the studies that are available by Gold et al. [48], McKinney et al. [49], and Hazell et al. [11] have all shown positive outcomes. In the latest of these trials [11], 60.4% of treated patients had normal LDLs by 25 months [11].

Psychological treatments, particularly cognitive behavioural therapy (CBT) (Andersson et al. 1999 [54]), probably have a role to play, in particular for those patients who have significant associated anxiety and distress. There is a paucity of evidence for this promising approach to date, but what evidence does exist is reviewed by Baguley and Andersson [5].

There is no evidence about how to treat patients with acoustic shock. Many clinicians treat it as a form of acute phonophobia, but this is based on intuition rather than science. Much effort is being spent investigating the nature of sounds that triggers acoustic shock in the hope that telecommunications equipment can be fitted with suitable filtering circuitry.

As with tinnitus, there is considerable value in a self-help approach to hyperacusis. The Hyperacusis Network (www.hyperacusis.net) is an important resource in this regard, with well-informed and well-moderated forums and much positive advice.

Summary

Disorders of loudness tolerance have received much less attention than tinnitus and still remain in the shadows. Far from being a rare and obscure condition, research has shown hyperacusis to be a common symptom, especially among patients with tinnitus. Various mechanisms have been suggested, and it seems likely that different mechanisms can apply to different patients. Although there is meagre published information on the management of sound hypersensitivity, the research that is available suggests that a programme of careful and gradual desensitization is effective for the majority.

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References

32. Sahley, TL and Nodar, RH. A biochemical model of peripheral tinnitus. Hear Res 2001; 182: 43–54
41. Valente, M, Potts, LG, and Valente, M. Differences and intersubject variability of loudness discomfort levels measured in sound pressure level and hearing level for TDH-50P and ER-3A earphones. J Am Acad Audiol 1997; 8: 59–67
42. Nielting, M, Rienhoff, NK, Hesse, G, and Lamparter, U. The assessment of subjective distress related to hyperacusis with
a self-rating questionnaire on hypersensitivity to sound. Laryngorhinootologie 2002; 81: 32–34
50. Baguley, DM. Current perspectives on hyperacusis J Royal Society of Medicine, 2003, 96, 1–4
Chapter 4
Misophonia, Phonophobia, and “Exploding Head” Syndrome

Aage R. Møller

Keypoints

1. Misophonia, phonophobia, and “exploding head” syndrome have symptoms that may occur together with some forms of tinnitus or they can occur alone.
2. These sensations are different from hyperacusis which is a lowered tolerance to most kinds of sounds.
3. Misophonia is a dislike of specific kinds of sounds.
4. Attempts have been made to treat misophonia using the same methods as used for treating tinnitus.
5. Phonophobia is a fear of specific sounds related to the implication of the sounds.
6. The non-classical auditory pathways providing a subcortical route to the amygdala may be involved in phonophobia.
7. The “exploding head” syndrome is the experience of a very loud and sudden noise that seems to originate from within the head. It often occurs during sleep and wakes up the individual.
8. The “exploding head” syndrome may have similarities with REM sleep behavior disorder (RBD).

Keywords  Tinnitus  •  Misophonia  •  Phonophobia  •  Exploding head syndrome

Introduction

There are several forms of decreased sound tolerance (DST); probably, the most common one is hyperacusis, which is a decreased tolerance level of (nonspecific) sounds, independent on their significance or importance (see Chap. 3). Misophonia is a decreased tolerance to specific sounds and phonophobia is fear of sounds: both disorders are based on the perceived implications or meanings of those sounds, whereas hyperacusis is not related to the comfort of the sound. The “exploding head” syndrome is hearing loud unexpected sounds, mostly during sleep or drowsiness. The prevalence of misophonia and phonophobia is unknown, and there are no known effective treatments. These abnormal reactions to sound are different from hyperacusis because the reactions are related to the significance of the sounds and are different from the common reaction to loud sounds that occur unexpectedly and which can cause a general body reaction known as a startle response. The “exploding head” syndrome is not a reaction to sound but occurs spontaneously. Even less is known about these symptoms than misophonia and phonophobia, and the available treatments are unsatisfactory.

The fact that these three different syndromes are not generally known opens the possibility of many kinds of maltreatment and misinformation from health care professionals, who often administer large batteries of tests, which are ineffective. Patients often go to one physician after another searching in vain for help.

Misophonia

Misophonia is defined as “dislike of certain specific sounds,” thus comparable with the term “phonophobia.”
Misophonia has been regarded a phantom sensation similar to tinnitus [1]. It has been discussed in connection with tinnitus and tolerance to sounds [2, 3]. Misophonia is different from hyperacusis in that it is only experienced in response to specific sounds, unlike hyperacusis, which is a lowered tolerance to all sounds (above a certain intensity) (see Chap. 3). A better word than misophonia may be “unpleasant” or “annoying.” These sounds that are unpleasant may also elicit autonomic reactions of various kinds.

Misophonia can occur together with tinnitus and hyperacusis, but may also occur alone. Treatment of misophonia has been discussed by Jastreboff, who suggested similar treatment to that used for tinnitus, namely the tinnitus retraining therapy (TRT) [4]. Beneficial effects of treatment of misophonia using TRT have been reported [2, 5, 6].

One can only guess about the anatomical location of the physiological abnormalities that cause misophonia. Since misophonia is related to specific sounds (its difference from hyperacusis), the anatomical location of the physiological abnormality that causes misophonia must be structures that are activated by highly processed sounds, thus located after auditory information has been subjected to considerable processing and selection. It seems likely that the location would be that of object (and frequency) processing sounds. It was mentioned in Chap. 8 that different kinds of sound are processed in different parts of the brain (stream segregation). This means the anatomical location of the abnormality that causes misophonia is located more central than that of hyperacusis because more neural processing of the sounds has occurred to cause misophonia compared with hyperacusis.

The anatomical location of the physiological abnormality may be in the inferior part of the temporal lobe where processing of object (“what”) information in humans occurs (see Chap. 8).

**Phonophobia**

Phonophobia means fear of sound and it is related to the content (or significance) of the sound. Some kinds of sounds can invoke fear in most people. Sounds that are understood not to be signaling an eminent danger usually do not invoke fear. This can be explained by considering the normal route of sensory signals to the amygdala through the “high route” [7, 8] (see Fig 9. in Chap. 8). The input to the high route comes through the classical auditory pathways where sounds use the ventral part of the thalamus from where connections lead to the primary auditory cortex, secondary cortex, association cortices, and from there to the lateral nucleus of the amygdala. This allows control by higher CNS regions of the flow of information in the high route and can therefore control the information that reaches the amygdala.

The situation is different if the non-classical pathways are active because there is a subcortical route to the amygdala from the dorsal and medial thalamus that is not controlled by higher CNS centers. Some individuals with severe tinnitus have signs that they use the non-classical auditory pathways [9], (see Chaps 8 and 10).

Functional imaging studies have supported the results of the reports that indicate an increased activity of structures of the limbic system [10].

**“Exploding Head” Syndrome**

“Exploding head” syndrome is a condition that causes the sufferer to occasionally experience a tremendously loud noise as if originating from within his or her own head. The “exploding head” symptoms usually occur during sleep or drowsiness [11]. Individuals with these symptoms explain it as explosions in the head. This syndrome can also cause the sufferer to feel an extreme rush of adrenaline kick going through his or her head, sometimes multiple times.

The “exploding head” syndrome and the abnormal perceptions that some people with tinnitus may experience is unpleasant and even described as a terrifying sensation of flashing lights, the sound of an explosion, gunshot, door slamming, roar, waves crashing against rocks, loud voices, a ringing noise, or the sound of an electrical short circuit. In some cases, an instant flash of what is perceived as video “static” is reported [12]. The “exploding head” syndrome may have similarities with audiogenic seizures, which has been studied in animals where it was found that the inferior colliculus was involved [13].

The exploding head phenomenon may be a failure to prepare the nervous system for sleep. It may be an exaggeration of the events that normally occur in the transition between being awake and being at sleep.
The normal transition between wakefulness and sleep requires that the reticular system changes the excitability of not only the motor system but also other CNS systems. Many people experience sounds that are perceived to be louder moments before falling asleep. This may have to do with the different steps needed in the process of changing the excitability (or gain) in sensory systems to preparation for sleep that are not fully synchronized.

The “exploding head” syndrome may be a result of failure of the automatic gain control that normally compresses the range of amplitudes of sounds. The auditory nervous system would not be able to process sounds in the enormous range of intensities of normal sounds without extensive gain control. Different stages of the auditory system have automatic gain control. The first structure that performs gain control is the cochlea, where amplification in the cochlea by the action of the outer hair cells decreases with the intensity of sounds. The amplification of this “cochlear amplifier” is to some extent controlled by the central nervous system through the olivocochlear bundle that is a part of the descending auditory pathway (see [14] and Chap. 8).

The “exploding head” phenomenon may have similarities with what is known as REM sleep behavior disorder (RBD) [15]. In some individuals, the system that normally keeps skeletal muscles paralyzed during REM sleep malfunctions causing violent behavior during REM sleep [16]. RBD is assumed to be caused by failure of the reticular system to maintain paralysis of skeletal muscles. Many people experience hyperacusis just before falling asleep, thus a sign that the reticular formation has affected the processing of auditory information.

Other forms of little known malfunctions of the reticular formation may be responsible for similar phenomena that may occur immediately after waking up. Some individuals can occasionally experience total paralysis for a few moments. This seems to be caused by a failure of the reticular formation to release the paralysis that occurs normally during REM sleep.

The symptoms of the “exploding head” can be reduced by reassurance of the harmlessness of the condition and the symptoms often ameliorate spontaneously with time. In a study, clomipramine, a tricyclic agent with both antidepressant and antiobsessional properties, has been reported to provide immediate relief of the symptoms [11]. None of the participants in these studies had any neurological disorders [11].

References

Chapter 5
Epidemiology of Tinnitus in Adults

Aage R. Møller

Keypoints

1. Many studies have addressed the prevalence of tinnitus, but the definition of tinnitus has varied.
2. Some studies have reported that as many as 80% of the adult population experience tinnitus at some point.
3. Six large population studies in different countries reported prevalence of prolonged tinnitus, varying between 4.4 and 15.1% for adults and between 7.6 and 20.1% for individuals below the age of 50 years. One of the studies reported that 2.4% of the population responded “yes” to the description of tinnitus as “tinnitus plagues me all day.”
4. A study in four cities in England found that tinnitus, on average, occurred in 17.5% of the participants in the age group of 40–60 years and 22.2% in participants above the age of 60 years.
5. Since tinnitus has many forms and its prevalence varies with age and, to some extent, gender, the prevalence of tinnitus cannot be described by a single number.
6. The prevalence of tinnitus increases monotonically up to the age of approximately 70 years, above which the prevalence either becomes constant or decreases slightly with age.
7. The prevalence of tinnitus is lower in women up to 75 years, above which the gender difference becomes small.
8. There is some evidence that noise exposure increases the risk of tinnitus.
9. The odds of having tinnitus increases with the degree of hearing loss when measured at 4 kHz.
10. While reported “trouble hearing” increases monotonically with age, “bothersome tinnitus” increases with age only up to the age group of 65–74, after which it becomes independent of age or decreases slightly with age.

Keywords Tinnitus • Epidemiology • Prevalence • Adults • Hearing loss • Noise exposure

Introduction

Understanding the incidence and prevalence of a disease in a defined population is important for improvement of health and prevention of diseases. Accurate determination of the prevalence of a condition, such as tinnitus, which does not have objective signs, depends on the ability to define the disease to the members of the population that is studied.

Tinnitus affects different groups of people differently, such as different age groups, and the prevalence of tinnitus in women and men is also different. This means that a single number cannot describe the prevalence of tinnitus. It is therefore important to define the part of the population that is studied.

Tinnitus is often accompanied by hyperacusis (lowered tolerance for sound, see Chaps.3 and 57), misophonia (dislike of certain sounds), and phonophobia (fear of certain sounds); see Chaps. 4 and 57. While the prevalence of tinnitus, in general, is poorly known, the prevalence of these symptoms is even less known. The effect of tinnitus on a person’s quality of life depends more on the distress it causes and less on how a person perceives his or her tinnitus. However, the prevalence of distress from tinnitus is poorly known.
When discussing the prevalence of tinnitus, it is the troubled tinnitus that is of the greatest interest because that is the form of tinnitus that affects the quality of life and which may have severe consequences for the person who has tinnitus. Troubled tinnitus may result in the inability to work and may have such a severe effect on a person that it causes suicide.

This chapter discusses population studies of the prevalence of tinnitus. Few studies have addressed the incidence of tinnitus which will not be discussed, and the natural history of tinnitus is not understood (see Chaps. 63 and 64) [1].

For studies of the prevalence of tinnitus, the greatest challenge lies in defining the tinnitus. As has been discussed in many of the chapters in this book, tinnitus has many forms (see especially Chaps. 2–4 and 17). Tinnitus varies widely among individuals not only in strength but also in character, and many investigators have proposed different classification schemes for tinnitus (for a review see Heller 2003) [2]. An individual’s tinnitus can vary widely from time to time. Many forms of tinnitus change from day to day and even change over the course of one day.

In that way, tinnitus has many similarities with pain. When the task is to obtain accurate information regarding its prevalence of tinnitus and pain, there are many aspects of these two symptoms that must be taken into account as has been discussed in Chaps. 14 and 94. Tinnitus can noticeably decrease the quality of life or it can just be a small annoyance. In fact, most people who have tinnitus do not regard it as anything important. One study reported that 0.5–1% of individuals with tinnitus indicated that the condition severely affected their ability to live a normal life [3]. Other studies have reported different estimates of prevalence of such forms of tinnitus.

The degree of distress tinnitus can cause is not related to the character or the perceived strength of the disorder as it is described by the persons who have tinnitus. The perceived severity of tinnitus depends on many different factors; one being a person’s personality (see Chaps. 27, 63 and 64). The perception of tinnitus is also influenced by external circumstances. These factors all make it difficult to obtain an accurate estimate of the prevalence of tinnitus that affects a person’s life. Different definitions of such forms of tinnitus have been used by individual investigators. This is one of the reasons that the results reported by different epidemiologic studies differ considerably, and different studies report prevalence of tinnitus that varies from study to study. The lack of objective signs of tinnitus is another source of uncertainty in studies of this disorder, and only self-reported evaluation of a person’s tinnitus is available. Most epidemiologic studies have not attempted to distinguish between the different origins of the tinnitus, not even distinguishing between objective and subjective tinnitus.

Another source of variation in the results of different epidemiologic studies of tinnitus is shared with other voluntary studies, namely, that not all persons selected for a study respond. Normally, epidemiologic studies will spend a considerable effort finding out if the group of non-responders is different from the group that responds.

Another reason for varying results in different studies is that questions are formulated differently. Some studies have used written questions distributed to groups of people more or less representative of the general population. Some studies have enrolled individuals seeking professional help for their tinnitus. The participants in some studies must therefore be regarded as being a selected group of individuals that may not be representative of the general population. Tinnitus depends on many factors, which makes it important to obtain a multi-dimensional description of its epidemiology. Thus, it is not meaningful to just describe the prevalence with a single number.

Estimates of the Prevalence of Tinnitus

Data from the National Center for Health Statistics, US Department of Health, Education, and Welfare (1968), indicate that 30% of the general population are affected by tinnitus, and that 6% of them (1.8% of the general population) have incapacitating symptoms [2]. Other studies have presented values of prevalence that vary between 7.6 and 20.1% (see Table 5.1).

Prevalence of Tinnitus as a Function of Age

One of the main variables in the prevalence of tinnitus is age, and studies have therefore expressed the prevalence of tinnitus as a function of age. Table 5.1 compares the
Epidemiology of Tinnitus in Adults

reported prevalence in several studies from the United Kingdom, Sweden, Norway, and the US.

All published studies show values of prevalence of tinnitus that are not the same for different age groups. All published studies seem to agree that the risk of getting tinnitus increases with age up to about 65 years, after which age prevalence is either independent of or decreases slightly with increasing age.

The design of the studies, the results of which are shown in Table 5.1, all had differences, which make it difficult to compare the results. The first study (United Kingdom National Study of Hearing) used a postal questionnaire sent to people in Cardiff, Glasgow, Nottingham, and Southampton in age groups between 17 and more than 80 years. In the questionnaire, tinnitus was defined as “prolonged spontaneous tinnitus” that lasts for more than 5 min and occurs not exclusively after loud sounds [5].

In the study from Gothenburg, Sweden, questionnaires were mailed and had blinded response (no follow-up of non-responders). Tinnitus was defined as an ear noise that occurs “often or always” and sounds like a peep, chirping, roaring, wind blowing in the trees, etc., [6]. In the same study, 2.4% of the population suffered from the worst severity degree defined as “tinnitus plaques me all day.”

The United States National Health Interview Survey (US NHIS) is a household survey with personal interviews of non-institutionalized civilians from randomly chosen areas constituting a nationally representative sample. The participants in this study had tinnitus that was defined as “having been bothered by ringing in the ears or other funny noises in the head in the past 12 months” [7]. The 1994–1995 US NHIS Disability Supplement study, Phase I, used an impairment and disability-screening questionnaire. Chronic tinnitus was defined in the interview as “now having a ringing, roaring, or buzzing in the ears that has lasted for at least 3 months” [8, 9].

The participants in the Beaver Dam, WI Hearing Loss study had significant tinnitus that was defined as “bothered by ringing, noise in the ears in the past year of at least moderate severity and/or tinnitus that caused difficulty in falling asleep” (Nondahl 2002) [10].

The Nord Trondelag, Norway Hearing Loss study used a self-administered questionnaire filled out in study clinics prior to the hearing examination [11]. Tinnitus was defined as “bothered by ringing in the ears.” The participants in this study were thus individuals who had sought professional help for their tinnitus. The results may therefore not be representative of the general population.

While there are large differences between the values of prevalence arrived at by different studies, there is agreement that the prevalence of tinnitus increases with age. Less clear is the relationship to gender, but studies show a tendency for tinnitus to occur more frequently in men than in women. Epidemiological studies of prevalence show a slightly larger prevalence of tinnitus in men, but the results are not consistent.

Most studies show an increase in the prevalence of tinnitus with age in the age groups up to 65–74 and considerably lower prevalence in individuals above 75 years than in the age group of 65–74 years. Since tinnitus of an individual rarely decreases the prevalence arrived at in epidemiologic studies, at least up to

<table>
<thead>
<tr>
<th>Age (year)</th>
<th>I (%)</th>
<th>II (%)</th>
<th>III (%)</th>
<th>IV (%)</th>
<th>V (%)</th>
<th>VI (%)</th>
</tr>
</thead>
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<tr>
<td>20–29</td>
<td>5.7</td>
<td>7.5</td>
<td>5.1</td>
<td>1.4</td>
<td>9.8</td>
<td></td>
</tr>
<tr>
<td>30–39</td>
<td>7.4</td>
<td>5.8</td>
<td>6.0</td>
<td>2.0</td>
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<td>40–49</td>
<td>9.9</td>
<td>8.9</td>
<td>7.2</td>
<td>3.7</td>
<td>11.8</td>
<td></td>
</tr>
<tr>
<td>50–59</td>
<td>12.5</td>
<td>18.6</td>
<td>10.1</td>
<td>5.7</td>
<td>7.3</td>
<td>16.9</td>
</tr>
<tr>
<td>60–69</td>
<td>16.3</td>
<td>20.3</td>
<td>13.0</td>
<td>7.9</td>
<td>10.1</td>
<td>20.2</td>
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<td>14.4</td>
<td>21.3</td>
<td>12.6</td>
<td>9.4</td>
<td>8.7</td>
<td>24.0</td>
</tr>
<tr>
<td>&gt;80</td>
<td>13.6</td>
<td>14.1</td>
<td>8.3</td>
<td>5.5</td>
<td>22.9</td>
<td></td>
</tr>
<tr>
<td>&lt;50</td>
<td>14.2</td>
<td>20.1</td>
<td>12.1</td>
<td>7.6</td>
<td>8.2</td>
<td>20.1</td>
</tr>
<tr>
<td>Adult</td>
<td>10.2</td>
<td>14.2</td>
<td>8.4</td>
<td>4.4</td>
<td>15.1</td>
<td></td>
</tr>
<tr>
<td>Participants</td>
<td>34,050</td>
<td>2,556</td>
<td>59,343</td>
<td>99,435</td>
<td>3,737</td>
<td>47,410</td>
</tr>
</tbody>
</table>


Data from Hoffmann and Reed [4]
the age of 70 can be regarded as being cumulative, above which it does not seem to increase very much.

This pattern of age relations of prevalence of tinnitus is clearly seen in the graphic representation of the results of another study (Fig. 5.1).

This graph shows that the prevalence of tinnitus for both men and women increases with age and that it levels out and even decreases above a certain age. The prevalence of tinnitus is higher for men than for women up to the age of 75, and the prevalence of tinnitus reaches its highest value earlier in life for men than for women. At the age of 75, the prevalence in women catches up with that of men, and above the age of 75 the prevalence of tinnitus is about the same for men and women. This pattern is thus similar to that of the prevalence of cardiovascular disease.

That the prevalence is less in the group of 80 years of age may have to do with tinnitus being associated with diseases, which may have caused early death. Therefore, those who die before the age of 80 may have had a higher prevalence of tinnitus than those who live beyond the age of 80 years. Again, it may suggest that tinnitus has similar risk factors as diseases that can cause early death, such as cardiovascular disorders.

Animal studies have shown that rats genetically predisposed for high blood pressure also acquire more hearing loss from noise exposure than animals not genetically predisposed for hypertension (normotensive rats) [12, 13]. There may also be (unknown) relations between cardiovascular factors and tinnitus in humans.

Those individuals who survive the age of 80 may, thus, have fewer risk factors that shorten lifespan and which are the same risk factors for tinnitus. It may therefore be more representative to compare the values for the higher age groups, such as the 60–69 years, where the prevalence arrived at in different studies varies between 20.3 and 8.7%.

Other studies found similar trends of prevalence of tinnitus increasing with age between 50 and 75 years, above which the prevalence decreased in two different studies: one from the US and one from Australia (Table 5.2).

It has been pointed out that the influence on daily life from tinnitus is different and that the ways in which questions asked of participants in epidemiologic studies

Table 5.2

<table>
<thead>
<tr>
<th>Age (years)</th>
<th>Women</th>
<th>Men</th>
<th>Participants</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 60</td>
<td>23.6</td>
<td>32.3</td>
<td>28.0</td>
</tr>
<tr>
<td>60–69</td>
<td>30.5</td>
<td>35.1</td>
<td>32.7</td>
</tr>
<tr>
<td>70–79</td>
<td>28.7</td>
<td>32.7</td>
<td>30.5</td>
</tr>
<tr>
<td>80+</td>
<td>27.7</td>
<td>21.5</td>
<td>25.4</td>
</tr>
<tr>
<td>All ages</td>
<td>28.6</td>
<td>32.2</td>
<td>30.3</td>
</tr>
</tbody>
</table>

Data from Sindhusake et al. [14]
Epidemiology of Tinnitus in Adults

A study from four cities in the UK (Table 5.3) [15], which reported that tinnitus occurred in an average of 17.5% of individuals, showed that annoyance was classified to be moderate or severe in 5.3% on average for the cities studied. This study also showed a similar pattern of age relationship (14.5% in persons younger than 40 years, 17.5% in the group of 40–60 years, and 22.2% in individuals older than 60 years).

Although the reported prevalence of tinnitus varies between different studies, twenty percent of people who say they have tinnitus reported their condition as “severe tinnitus.” This means that about 80% of patients with tinnitus suffer little and are not seeking treatment.

### Hearing Loss and Tinnitus

A study shows that the risk of having tinnitus (expressed as odds ratio) increases with the degree of hearing loss at 4 kHz [16] (Fig. 5.2).
“Trouble Tinnitus” Compared with “Trouble Hearing”

There is considerable difference between having tinnitus and being troubled by tinnitus (having “trouble tinnitus”). The same is the case for hearing; having “trouble hearing” is a distinction from just having hearing loss. In a study based on the US National Health Interview Survey, participants were asked to report whether they had a “lot of trouble hearing” or “any trouble hearing,” and for tinnitus the participants were asked if they had “bothersome tinnitus.” In this study, the reported hearing trouble only increased slightly after age of 65, when self-reported problems were concerned (Fig. 5.3).

It is evident that while hearing loss increases monotonically with age, the prevalence of bothersome tinnitus levels off and even decreases after the age of 70 years, thus, in agreement with other studies reported above. That bothersome tinnitus reaches a level of about 14% in the 65- to 74-year age group should be noted.

Can Hearing Loss Cause Tinnitus?

The prevalence of tinnitus increases with age, and it has been discussed whether hearing loss can be a contributing cause of this increase in the prevalence of tinnitus with age. Audiometric data show that hearing loss increases with age [17, 18] (see Chap. 36) The results of studies discussed in this chapter show that the prevalence of tinnitus increases with age, but it is not known if it is age-related changes in the ear and the nervous system that cause the tinnitus to increase with age or if it is the age-related hearing loss that causes the increase in prevalence of tinnitus. Population studies have shown that individuals with tinnitus, on average, have hearing loss affecting mostly high frequencies [19] (Fig. 5.4).

Other studies (Table 5.4) have confirmed the difference in prevalence of tinnitus between men and women.

There is considerable evidence that deprivation of input to the auditory nervous system can cause tinnitus (see Chaps. 11–13 and 21). There is also evidence that noise exposure can cause hearing loss (see Chap. 37), but it is not known if it is the noise exposure, as such, or the associated hearing loss that causes the tinnitus.

The hearing loss shown in Fig. 5.4 is slightly greater in males than in females, resembling what is found in population studies of hearing and showing signs of a 4-kHz dip, indicating that some of the hearing loss is likely to have been caused by noise exposure.

As an example of the diversity of tinnitus, it has been shown that the type of hearing (shape of the audiogram) is related to the character of the tinnitus.
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Epidemiology of Tinnitus in Adults

Patients with low-pitched tinnitus (<1,500 Hz) tend to have much more severe hearing losses, especially in the low frequencies, than do patients with higher pitched tinnitus.

Again, it must be emphasized that these data are also from tinnitus clinics, thus, only including people who have sought professional help. While tinnitus does occur in individuals with normal hearing, people with tinnitus usually have hearing loss, and deprivation of sound activation of the nervous system can cause tinnitus by activating neural plasticity, as discussed in Chap. 12. However, the prevalence of troubled tinnitus does not increase above the age of 65, while audiometric hearing loss does continue to increase with age above 65.

The fact that tinnitus cannot be measured objectively as can hearing loss means that comparing tinnitus with audiometrical hearing loss may be regarded to be an invalid comparison. However, as seen in Fig. 5.3, the subjective trouble with hearing also increases with age while the prevalence of tinnitus is not changing above the age of 65. This could be because the debut of tinnitus above that age is rare or that some individuals who had tinnitus before the age of 65 improve and that counteracts an increase in the new cases of tinnitus. This question cannot be answered because the natural history of tinnitus is poorly known. Studies that have concerned the natural history of tinnitus have only reported on the presence of the disorder, not its severity [20, 21, 22] or how the individuals perceive their tinnitus. There are two reasons why the prevalence of tinnitus may be higher in a population of individuals with hearing loss. One reason is that hearing loss implies a certain degree of deprivation of input to the auditory system (see Chap. 11), which is known to be able to activate neural plasticity, and known to be involved in many forms of tinnitus (see Chaps. 12 and 13). Another reason that there may be a relationship between the prevalence of tinnitus and hearing loss is that the same factors that cause hearing loss may cause tinnitus. Such common factors may be age, cardiovascular disorders, and noise exposure.

When the kind of tinnitus individuals with hearing loss have is correlated with the shape of their audiograms, it follows that there are distinct correlations between the pitch of an individual’s tinnitus and the.

Table 5.4 Prevalence in percentages for males and females

<table>
<thead>
<tr>
<th>Age</th>
<th>Male (%)</th>
<th>Female (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>18–24</td>
<td>4.3</td>
<td>5.2</td>
</tr>
<tr>
<td>25–44</td>
<td>5.8</td>
<td>6.2</td>
</tr>
<tr>
<td>45–64</td>
<td>10.6</td>
<td>9.5</td>
</tr>
<tr>
<td>65+</td>
<td>12.3</td>
<td>13.9</td>
</tr>
</tbody>
</table>

Prevalence based on a self report of “bothersome tinnitus” as a function of age, sex, and percentage of each characteristic in the population; based on the 1990 Hearing Supplement to the National Health Interview Survey. Data from Hoffmann and Reed [4]
individual’s audiogram (Fig. 5.5). This is yet another complexity of tinnitus that makes it difficult to establish clear data on its prevalence.

It was shown in a study of patients with tinnitus of different pitch that patients with low-pitched tinnitus (less than 1,500 Hz) tend to have much more severe hearing losses, especially at the low frequencies, than do patients with higher pitched tinnitus.

The studies reported in the two preceding graphs concerned patients who had sought help for their tinnitus. This means that the participants do not represent a random selection of people. There are many reasons why people seek professional help and equally many reasons why people do not seek professional help.

**Other Risk Factors for Tinnitus**

Noise exposure and noise that induced hearing loss is another factor anecdotally reported to cause tinnitus in individuals.

Risk factors for tinnitus other than age are hearing loss, diseases such as middle-ear disorders, Ménière’s disease, cerebrovascular diseases, and, in particular, hearing loss of various causes and environmental factors such as exposure to noise and administration of certain medications such as ototoxic antibiotics and acetylsalicylate.

Noise exposure increases the risk of tinnitus, and, at the same time, it causes hearing loss. The question is, therefore, if the tinnitus from noise exposure is caused by the hearing loss associated with noise exposure. The increased prevalence of tinnitus in males may have to do with the increased noise exposure in males and subsequent higher frequency of hearing loss in men [17].

**Tinnitus and Suffering**

Tinnitus is a sensation, and suffering may be related to and possibly a consequence of having tinnitus. The prevalence of tinnitus, counting all forms, is of little interest from a health care perspective because most individuals with tinnitus are not bothered to an extent that it affects their daily life, and few will seek medical attention except for those who want to be sure that their tinnitus is not a sign of a serious disease. In that way, tinnitus has similarities with pain. Most people have experienced pain in one form or another, but only a few have severe pain that causes suffering.

**References**


Chapter 6
Epidemiology of Tinnitus in Children

Claudia Barros Coelho

Keypoints

1. Children experience tinnitus and might present similar suffering as observed in adults but they rarely mention the symptom unless directly asked about it.
2. Difficulty on concentration, sleeping, hearing, leisure activities, sports practice, and hyperacusis are the most frequent complaints associated to tinnitus in children.
3. Only a few population studies have been performed and have disclosed prevalence rates from 6% to 59%. Many factors might be implicated in the large inter-study variability of tinnitus prevalence in children.
4. Age, gender, hearing loss, motion sickness, hyperacusis, and noise exposure have been suggested as risk factors to development of tinnitus in children.
5. A proper model to investigate children should be developed for the purpose of obtaining accurate information about the prevalence of tinnitus in children.
6. Preventive measures should aim at hearing education about the risk of hearing loss and tinnitus. Prevention of noise exposure should be promoted as early as possible.

Keywords Tinnitus • Epidemiology • Children • Hyperacusis • Preventive measures • Sleep

Abbreviations

OR OR
TTS Temporary threshold shift
HL Hearing Level

Introduction

Children rarely mention tinnitus unless they are asked specifically about it. The frequency with which they mention the symptom spontaneously ranges from 1.6% to 6.5% [1–4]. Therefore, the observed proportion of children who seek professional help does not represent all children with tinnitus. Also, investigating tinnitus is seldom a part of routine pediatric otolaryngological practice. For these reasons, the prevalence of the symptom is generally underestimated in childhood [5].

Children who experience tinnitus may suffer in a similar way as adults with tinnitus. Difficulty in concentration, sleeping, hearing, and hyperacusis are the most frequent complaints associated with tinnitus in children [1, 6–8]. The symptoms might affect many kinds of leisure activities such as sports [1] as well as cause a decrease in school performance [9, 10]. The symptoms may significantly interfere with children’s life in general, which will inevitably affect their entire families as well [11].

Terms such as “ringing” [2, 12], “beeping” or “buzzing,” and a “high-pitched noise” or “whistling” [6] have been used by children to describe tinnitus sounds.

Some hypotheses have been presented regarding why children rarely report tinnitus spontaneously. (1) Children rarely refer to symptoms that are not associated with pain [13]; (2) children have a less-developed body image [14]; (3) there are specific differences in the ascending
auditory pathways in children [15]; (4) children may perceive tinnitus as a familiar experience [16]; (5) children may be more easily distracted by events of the external environments [17]; (6) do not perceive the medical significance of the symptom [18]; and (7) children’s attention process is different from that in adults, and this might also have an effect on how they perceive tinnitus.

In order to diagnose tinnitus in children, it is therefore important to ask children specifically if they have tinnitus.

**Studying Tinnitus in Children**

When studying tinnitus in childhood, it must be kept in mind that a child is not a miniature version of the adult. Children obviously do not possess adult brains. The organs of perception linking the child to the external world are still under maturation. The organization of sensory systems in the brain [15] (see also Chap. 8) and perception and attention in a child are different from adult, promoting a different perception and attitude to the world.

Other obstacles in studies of tinnitus in children are related to the fact that children tend to give positive answers to please the interviewer [19] and it is important to minimize and parents preoccupations that children’s their might have after being aware of tinnitus.

In managing tinnitus in children, it is important to distinguish between the perception of the tinnitus and the impact that the tinnitus has on a person (tinnitus suffering) [5]. Lack of information about the prevalence of tinnitus suffering in children makes it difficult to judge the impact of tinnitus on children.

**Epidemiological Studies**

Although the existence of tinnitus in childhood has been reported since the 1970s there is still great uncertainty regarding the prevalence of tinnitus in children.

**Population Studies**

The few population studies that have been published were done in only a few countries and have shown widely different values of prevalence (from 6% to 59%) (Table 6.1).

Many different factors may have contributed to the discrepancies between the results of the different studies that have been published: (1) the criteria used for defining tinnitus may have been different; (2) hearing criteria may have been different; (3) age range may have been different; (4) methodological factors – interview or questionnaires most likely were different; (5) studies have used different statistical procedures, with different sample sizes; (6) the effect of confounding variables may also have contributed to the variations, for example, social and economic classes, ethnic, and cultural background may have varied; (7) different behavioral factors may have influenced the results such as emotional problems; (8) the effect of environmental factors such as exposure to noise may have been different.

Two studies of the prevalence of tinnitus in children had many participants recruited from otolaryngological clinics. Aust [20] screened children who sought help for otological complaints and Savastano [4] evaluated a general population of children using a specific protocol to investigate tinnitus (Table 6.2). They found tinnitus in 7% and 34%, respectively.

**Factors that may Promote Tinnitus (Risk Factors)**

Risk factors refer to an increase in the chance that an event is going to occur; in the present situation, this means the likelihood that a child will get tinnitus. Identification of risk factors plays an important role toward understand the etiology of tinnitus. Identification of risk factors might help understanding the symptoms and develop strategies for prevention of tinnitus (see Chap. 69). They can be identified by logistic regression models where the risk odds is determined while controlling for irrelevant factors. The OR (OR) is the likelihood that an event will occur; in our case, the likelihood of occurrence of tinnitus versus the chance of absence of tinnitus. The OR for a predictor tells the relative amount by which the odds of the outcome increases (OR > 1.0) or decreases (OR < 1.0). Decrease in OR is a sign that a protective factor against the occurrence of an event is present.
<table>
<thead>
<tr>
<th>Author</th>
<th>Place</th>
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<th>Age</th>
<th>Study design</th>
<th>Diagnosis based on</th>
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<tr>
<td>Nodar [12]</td>
<td>USA</td>
<td>2000</td>
<td>10–18</td>
<td>Longitudinal</td>
<td>Questionnaire &quot;Do you hear a noise in your ears like ringing, buzzing, or a click?&quot;</td>
<td>13.3% normal hearing</td>
<td>58.6% hearing impaired</td>
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<td>Stouffer et al. [19]</td>
<td>Canada</td>
<td>161</td>
<td>7–12</td>
<td>Cross-sectional</td>
<td>Interview &quot;Do you hear a noise in your head for more than 5 min?&quot;</td>
<td>13% in normal hearing</td>
<td>29% in hearing impaired</td>
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<tr>
<td>Holgers and Svedlund [21]</td>
<td>Sweden</td>
<td>964</td>
<td>7</td>
<td>Cross-sectional</td>
<td>Questionnaire &quot;After listening to loud music or other loud sounds/noise, have you afterwards heard aringing, buzzing or other sort of noise in your ears, even if the loud music or noise has been turned off?&quot; &quot;Have you heard a ringing in the ears, without first having listened to loud music or other loud sounds?&quot;</td>
<td>13% in normal hearing</td>
<td>8.8% in hearing impaired</td>
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<tr>
<td>Holgers and Pettersson [29]</td>
<td>Sweden</td>
<td>671</td>
<td>13–16</td>
<td>Cross-sectional</td>
<td>Questionnaire &quot;How often do you experience tinnitus?&quot; &quot;How often is tinnitus annoying?&quot; &quot;Thoughts about tinnitus&quot;</td>
<td>53% tinnitus perception</td>
<td>27% tinnitus annoyance</td>
<td></td>
</tr>
<tr>
<td>Coelho et al. [1]</td>
<td>Brazil</td>
<td>506</td>
<td>5–12</td>
<td>Randomized cross-sectional</td>
<td>Interview Tinnitus sensation &quot;Do you hear a noise inside your ears/head?&quot; &quot;Describe the sounds perceived&quot; and &quot;locate&quot;.</td>
<td>Tinnitus perception 37.7% normal hearing 50%</td>
<td>Tinnitus annoyance 19% normal hearing</td>
<td></td>
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<td></td>
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<td></td>
<td></td>
<td>Hearing impaired 50%</td>
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<td>17.8% hearing impaired</td>
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<tr>
<th>Author</th>
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<tbody>
<tr>
<td>Aksoy et al. [8]</td>
<td>Turkey</td>
<td>1020</td>
<td>6–16</td>
<td>Cross-sectional</td>
<td>Questionnaire</td>
<td>9.2% tinnitus perception</td>
<td>5.8% tinnitus annoyance</td>
<td></td>
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<tr>
<td>Bulbul et al. (2009) [34]</td>
<td>Turkey</td>
<td>428</td>
<td>11–18</td>
<td>Cross-sectional</td>
<td>Questionnaire</td>
<td>33.4% tinnitus</td>
<td>35.2% tinnitus after listening high volume</td>
<td></td>
</tr>
<tr>
<td>Savastano [4]</td>
<td>Italy</td>
<td>1100</td>
<td>6–16</td>
<td>Observational</td>
<td>Interview</td>
<td>34% tinnitus perception</td>
<td>76.4% normal hearing</td>
<td>24.6% hearing impaired</td>
</tr>
<tr>
<td>Aust [20]</td>
<td>Germany</td>
<td>1420</td>
<td>7–12</td>
<td>Observational</td>
<td>Interview</td>
<td>7.2% tinnitus</td>
<td>26.4% normal hearing</td>
<td>75.3% hearing impaired</td>
</tr>
</tbody>
</table>
To our knowledge, only two studies on tinnitus prevalence in children have so far used such statistical analysis.

The following risk factors have been identified on tinnitus in children.

**Age**

The risk for tinnitus sensation and tinnitus annoyance increases with age by 1.1 times, for every year among children in the Brazilian study [1] and by a factor of 1.2 according to Nodar [12]. Aksoy et al. [8] reported a progressive increase on tinnitus incidence around the age of 13–14 years from 10 to 18 years and 6 to 16 years have observed.

**Gender**

Holgers and Svedlund [21] found a higher prevalence of tinnitus among girls, as well as a higher prevalence of depressive and anxiety symptoms. Coelho et al. [1] found that boys had an OR of 0.

To present, tinnitus suffering when compared to girls means that the male gender was a protective factor for the development of tinnitus among children. These findings could be related to: (1) girls present a higher tendency to express symptoms than boys, including those related to affective disorders [22]; (2) spontaneous otoacoustic emissions are more frequent among females [23] and have been described as a possible tinnitus etiology[24]; (3) genetic differences among genders associated with neurotransmitter expressions pursuing an action on auditory pathway, including serotonin [25] and female reproductive hormones affect GABA receptors in the brain [26] (see Chap. 10).

**Hearing Loss**

Tinnitus is more frequent in children with normal hearing [12, 19] than in hearing impaired children, but children with profound hearing loss have lower prevalence of tinnitus than children with moderate loss [27]. Comparison of children with middle ear disease to those with sensorineural hearing loss showed that 43.9% of children with middle ear disease had tinnitus while 29.5% with sensorineural hearing loss had tinnitus [16].

Children with hearing loss had an OR of 3.3 regarding tinnitus that could not be related to sound exposure according to a Swedish study from Holgers and Svedlund [21].

Similar findings were made by Coelho et al. [1] using a regression model where tinnitus was less prevalent in children with moderate to profound sensorineural hearing loss, than in those children with minimum to mild hearing loss. Minimum to mild hearing loss was a risk factor for tinnitus with an OR of 1.8 for tinnitus sensation and 2.4 for tinnitus suffering. Moderate to profound hearing loss (including deafness) was also considered risk factors with ORs of 0.5 for tinnitus sensation and 1.1 for tinnitus suffering.

The fact that a mild loss on hearing is a risk factor for tinnitus in children may be explained by the finding that even a mild hearing loss (thresholds at 30 dB HL) could promote tonotopic reorganization of the auditory cortex [28].

**Temporary Threshold Shifts**

Holgers and Petterson [29] have reported that individuals with temporary threshold shift (TTS) from noise exposure had an OR of 1.4 to present spontaneous tinnitus and 2.0 to noise-induced tinnitus. When comparing participants who sometimes experienced TTS to participants who did not have TTS, the OR was 2.8 to present spontaneous tinnitus and 8.4 to noise-induced tinnitus.

**Noise Exposure**

Holgers and Petterson [29] found that adolescents who attended concerts and discos/clubs had an OR of 1.4 regarding noise-induced tinnitus. Individuals who visited concerts 6–12 times per year had an OR of 4.4, compared to those who never went to concerts. Children who visited discos/clubs had an OR of 3.8.

Coelho et al. [1] reported that history of noise exposure was a risk factor for both tinnitus sensation and tinnitus suffering with ORs of 1.8 and 2.8, respectively. They found that firecrackers were the most frequent kind of noise exposure. Such noise may have
peak levels of 145–165 dB HL at a distance of 2 m or less from the explosion site [30]. Risk of exposure to excessive noise from toys has also been mentioned on the literature [31, 32]. Exposure to high levels of noise from toys and firecrackers were reported by 25% of children who sought medical care because of noise trauma [33].

Tinnitus is also often associated with the use of music players such as the walkman and iPOD devices both in the right ear \((p=0.004)\) and in the left ear \((p=0.000)\) [34].

Activation of neural plasticity by overexposure or reduced impact to the auditory nervous system caused by hearing loss may cause tinnitus (see Chaps. 12 and 13). The reorganization on the tonotopic map of the primary auditory cortex following noise trauma is one sign of activation of neural plasticity that has been documented in several studies [35, 36] and it has been suggested that tinnitus may be related to such reorganization [37–39].

**Motion Sickness**

Motion sickness was found to be a risk factor for tinnitus sensation with an OR of 1.8 [1]. Motion sickness has been highly associated to migraine and vestibular symptoms in children [40].

**Hyperacusis**

Hyperacusis and tinnitus are related symptoms [41] (see Chap. 3). Coelho et al. [1] showed that hyperacusis was the highest risk factor for tinnitus in children, with an OR of 4.2, but tinnitus was not a risk factor for hyperacusis [1, 42].

**Conclusions**

The remedy from some of the shortcomings of present studies is as follows.

The available data regarding the epidemiology of tinnitus have a high degree of variations among different studies. There is therefore a need of more studies to bring down the variability. This chapter has pointed to some factors that have contributed to the variations in the results among different studies. Cross over or cohort studies with randomized samples representative of the whole population should be considered. Participants for such studies could be recruited from schools where stratification and randomization of the participants can be achieved. Participants from a school environment have fewer dropouts; consents from parents can easily be obtained. Multivariate regression models should be used to describe risk factors.

Some of the problems with present studies are related to the definition of tinnitus. Standardized interviews such as: “Do you hear a noise (sound) in your ears or in your head that last more than 5 min?” should be used in evaluation of the tinnitus, and evaluation of the impact on everyday life is important. Questions such as “Does this noise (sound) bother you?” should be included in the questionnaires.

Audiological testing is important for evaluating tinnitus etiology and standardized methodology, and classification of results should be used.

An epidemiological surveillance system would be the basic action to prevent tinnitus. Efficient preventive measures should aim at hearing education and prevention of noise exposure as early as possible (see Chap. 69).

**References**


42. Coelho, CB, TG Sanchez, and RS Tyler, Hyperacusis, sound annoyance and loudness hypersensitivity in children. Prog Brain Res, 2007 166:169–78
Chapter 7
Genetic Risk Factors in Chronic Tinnitus

Philipp G. Sand

Keypoints

1. Individual susceptibility to chronic tinnitus is shaped by the interplay of genetic and environmental factors.
2. Whereas many environmental risks including noise trauma and medication side effects are already well understood, heritable risks remain to be specified.
3. Pilot biometric studies in twins have produced heritability estimates of up to 0.39 in subgroups of affected patients but are still burdened with confounders.
4. The current review addresses the quest for molecular genetic biomarkers of tinnitus and the candidate genes examined so far.
5. Of these, genes encoding neurotrophic factors BDNF and GDNF give promising results that warrant further study.
6. Public attitude toward advances in genetic testing for tinnitus is as yet unexplored and deserves consideration in future research.

Keywords Tinnitus • Association study • Familial clustering • Genetic risk • Heritability • Mutation screening • Tinnitus susceptibility

Abbreviations

BDNF brain-derived neurotrophic factor
CNTF ciliary neurotrophic factor
GDNF glial cell-derived neurotrophic factor
HTTLPR serotonin transporter gene length polymorphic region
HTR1A serotonin receptor 1A
HTR3A serotonin receptor 3A
HUNT-II North-Trøndelag Health Study II (1995–1997)
PRNP Prion protein
SLC6A4 (5-HTT) solute carrier 6A4 (serotonin transporter)

Introduction

Tinnitus is a common clinical syndrome with an estimated 30 million sufferers in the United States [1]. Of those who develop a chronic form of tinnitus, the majority also experience varying degrees of hearing loss. However, the relationship between both conditions is complicated by dissimilar age-specific prevalence rates and interfering risk factors. For tinnitus, these comprise male sex, cigarette smoking, occupational noise exposure, lower income, higher body mass index, and reduced general health status, among others [2]. It is also evident from anecdotal reports that environmental risks interact with heritable susceptibility to chronic tinnitus that is as yet poorly defined [3]. Together, these elements have become a focus of public awareness [4] that is accelerating scientific research into the biological underpinnings of tinnitus. The present review summarizes the latest advancements on innate factors that may help account for interindividual differences in severity and course of symptoms.
Familial Aggregation of Chronic Tinnitus

Limited evidence is currently available on the familial clustering of chronic tinnitus. In siblings of affected individuals, a twofold risk has been noted relative to the general population [2]. A multicentre investigation involving 198 families from six European countries confirmed familial aggregation of tinnitus, albeit to a lesser extent than clustering observed for age-related hearing impairment [5]. A Danish study of 956 monozygotic and dizygotic twins found no evidence of heritability for chronic tinnitus in male individuals, but estimated heritability at .039 in female individuals [6]. Owing to the elderly age of the participants (≥70 years), the impact of other determinants of physical health on prevalence rates is a concern.

Few attempts have been made so far to identify specific features of tinnitus that are highly heritable and that may serve to define subgroups at risk. In 1,147 Belgian individuals with tinnitus, effects of familial loading on the curvature of the audiogram were investigated in an effort to distinguish highly heritable from sporadic conditions [7]. The authors claimed an association of familial forms of tinnitus with flat audiograms, but replication is lacking. Others have proposed that tinnitus in conjunction with hearing loss may predict familial tinnitus [8].

A pilot investigation in Norway has addressed the respective contributions of genetic and environmental factors to chronic tinnitus as part of the Nord-Trøndelag Hearing Loss Study [9]. In this study, epidemiological findings were merged with public registry data on the relatedness between individuals and gave a heritability estimate of 0.106 for a broadly defined tinnitus phenotype. Results were based on data from 51,975 individuals in 17 of 23 municipalities of a rural county (age range: 20–101 years; mean age: 50 years). Participants underwent pure-tone audiometry and completed self-report questionnaires between 1996 and 1998, providing information on occupational and leisure noise exposure, medical history, and symptoms of hearing impairment. Valid audiometry and questionnaire data refer to a core sample of 50,132 individuals [10]. While the authors argue that selection toward individuals with poor hearing is unlikely to have occurred in ~90% of participants who had been recruited as part of an earlier general health survey (HUNT-II), a weakness of the study is the specification of tinnitus-related complaints. Participants were asked whether or not they were “bothered by ringing in their ears,” i.e., tinnitus annoyance was used as a surrogate of tinnitus perception. As has been noted before, reference to the term “bothersome tinnitus” is liable to produce conservative estimates of the syndrome’s actual prevalence [1]. Pending the detailed publication of findings from this Norwegian population, the likelihood of type II errors in future studies may be reduced by a standardized evaluation of severity with attention to wording, the exclusion of known confounders, and further narrowing of the syndrome.

Candidate Gene Studies

A number of candidate genes have been investigated in individuals diagnosed with chronic tinnitus using either a case–control design (e.g., [11]) or a systematic mutation screening approach [12] (Table 7.1). From these investigations, there is little support for the notion of a serotonin-related etiology of tinnitus [13]. Specifically, negative associations of tinnitus with serotonin receptor genes 1A and 3A [12, 14], plus the serotonin transporter gene have been documented [15]. In contrast, positive findings have emerged for two genes that encode neurotrophic factors [11, 16]. Thus, the risk for developing tinnitus in conjunction with hearing impairment was significantly reduced in carriers of a missense variant in the gene encoding brain-derived neurotrophic factor (BDNF) [11]. When information from BDNF variants was combined with data on variants in the gene encoding glial cell-derived neurotrophic factor (GDNF), 16% of the variance in tinnitus severity could be explained [16]. Neurotrophins play key roles in tonotopic organization of the central auditory pathway [17, 18] and have been implicated in defective neuroregeneration in the cortex and hippocampus [19]. Both BDNF and GDNF protect the inner ear against trauma [20] and BDNF expression patterns dynamically respond to traumatic acoustic stimuli [21]. While BDNF expression is decreased in the primary auditory cortex, upregulation occurs within days in the inferior colliculus in what has been considered a putative correlate of ongoing neuronal repair. Together, these experimental data
warrant further exploration of neurotrophin-related signaling in animal models of tinnitus and can serve as a starting point for a targeted search of tinnitus biomarkers in humans.

**Outlook**

It is currently too early to speculate on the possible benefits that can be expected from future genetic tests for tinnitus, or even to predict when such tests may become available. However, genetic testing for other communication disorders (e.g., for deafness) has underscored the importance of providing potential consumers of these services with full information on all aspects of genetic evaluation [22]. Public education will need to address cost effectiveness and the appropriate use of resources. Many related issues, e.g., prenatal testing and gene therapy, are not without controversy and have given rise to individual and societal concerns over recommending tests for heritable syndromes. Adequate understanding of the needs and desires of parents and family members is therefore essential to guide the application of new genetic technologies to clinical practice [23]. In the largest survey of parents of children with hearing loss conducted in the USA to date, positive feelings about advances in the genetics of hearing loss prevailed [24], but no surveys exist to gauge interest in a DNA-based test for tinnitus. Unlike rare monogenic disorders of hearing, tinnitus is presumed to feature a much more complex mode of inheritance [8], which will call for a judicious interpretation of test results that rely on only one or few emerging risk factors.

**References**

Chapter 8
Anatomy and Physiology of the Auditory System

Aage R. Møller

Keypoints

1. The auditory system consists of four anatomically separate structures:
   (a) those that conduct the stimulus to the receptors
   (b) the receptors
   (c) the auditory nerve
   (d) the central auditory nervous system

2. The most important part regarding tinnitus is the auditory nervous system.

3. The auditory nervous system consists of two parallel ascending pathways that project to auditory cortices and two (reciprocal) descending pathways that project to nuclei of the auditory pathways.

4. The nuclei in the ascending auditory pathways process information in a serial hierarchical fashion, and processing occurs in modules with specific functions.

5. Two separate ascending sensory pathways have been identified in the auditory pathways: classical pathways and the non-classical pathways. Also, the somatosensory and visual pathways have two different ascending tracts.

6. The classical pathways are also known as the lemniscal system, or the specific system, and the non-classical pathways are also known as the extralemniscal system, or the unspecific system. The non-classical pathways have been divided into the diffuse system and the polysensory pathways.

7. The classical and non-classical pathways process information differently and have different central targets, especially regarding connections to the thalamus and the cerebral cortex.

8. The non-classical ascending auditory pathways branch off the classical pathways at several levels, the most prominent being the central nucleus of the inferior colliculus.

9. The auditory pathways receive input from the somatosensory system at the external nucleus of the inferior colliculus and from the dorsal cochlear nucleus as well.

10. The auditory pathways are mainly crossed, but there are extensive connections between nuclei at the two sides at two levels: the pontine nuclei (superior olivary complex) and the midbrain level (inferior colliculus). There are also extensive connections between the two sides at the cerebral cortical level.

11. The auditory nerve sends collaterals to cells in all these divisions of the cochlear nucleus. That is the earliest sign of the anatomical basis for parallel processing of information. Parallel processing occurs throughout the ascending pathways by axons branching to connect to more than one group of nerve cells.

12. Descending auditory pathways are abundant, in particular, the cortico-thalamic pathways, but little is known about their function. The descending pathways are largely reciprocal to the ascending pathways. The descending pathways reach as far caudal as the receptors in the cochlea.

13. The classical sensory pathways are interrupted by synaptic contacts with neurons in the ventral parts of the thalamus, which project to the primary sensory cortices.

14. The non-classical sensory pathways use the dorsal and medial thalamus as relay, the neurons of which
project to secondary and association cortices thus bypassing the primary sensory cortices.

15. Neurons in the dorsal and medial thalamus make direct (subcortical) connections with other parts of the CNS, such as structures of the limbic system, while the classical sensory systems connect to other parts of the CNS, mainly via association cortices.

16. There are anatomical connections between the upper spinal cord and the dorsal cochlear nucleus and between the caudal trigeminal nucleus and the dorsal cochlear nucleus. There are anatomical connections between the somatosensory system and midbrain nuclei of the non-classical auditory system.

17. Neurons in the nuclei of the classical pathways respond distinctly to specific sensory stimuli and have distinct frequency selectivity.

18. Sound stimulation may increase the firing rate of auditory nerve fibers, but saturation occurs for most fibers at low sound intensities.

19. Periodic sounds cause many nerve fibers to become locked to the waveform of the sound, and consequently, the firing of such fibers becomes time locked to each other. It subsequently causes the discharge of many neurons in the ascending auditory pathways, which then become time locked to each other.

20. Stream segregation implies that different types of information (for example, spatial and object information) are processed in anatomically different parts of the sensory nervous system.

21. Parallel processing allows the same information to be processed in anatomically different parts of the nervous system, while stream segregation implies that different kinds of information are processed in anatomically different structures.

22. Much less is known about the functional role of the non-classical pathways compared to the classical pathways, but neurons of the nuclei of the non-classical pathways respond less distinctly and are broader tuned than cells in the classical pathways and respond to a broad range of stimuli. They also integrate information on wider spatial scales than the classical pathways.

23. Neurons in the nuclei of the classical auditory pathways, up to and including the primary auditory cortex, respond only to one sensory modality (sound) while neurons of higher order cortices (secondary and association cortices) integrate information from several sensory systems and respond to different sensory modalities. This response can be modulated by input from non-sensory brain areas such as the amygdala.

24. Some neurons in the ascending non-classical pathways respond to more than one sensory modality. Their response to sound can be modulated by other sensory input.

25. The non-classical pathways make direct (subcortical) connections from the thalamus to other parts of the CNS, such as structures of the limbic system, while the classical sensory systems connect to other such parts of the CNS mainly via association cortices.


27. There are no signs of cross-modal interaction in adults, except with some forms of tinnitus and in autistic individuals, indicating that the non-classical auditory pathways are not normally active in adults.

28. Sensory systems connect to motor systems, the limbic system, reticular activating system, and the autonomic nervous system through subcortical and cortical routes.

29. There is considerable interaction between different systems in the brain, such as between different sensory systems and between sensory systems and non-sensory systems.

**Keywords** Ear • Auditory pathways • Anatomy • Non-classical pathways • Physiology • Cross-modal interaction

**Abbreviations**

- **AAF** Anterior auditory (cortical) field
- **AES** Anterior ectosylvian sulcus area
- **AI** Primary auditory cortex
- **AII** Secondary auditory cortex
- **AN** Auditory nerve
- **AVCN** Anterior ventral cochlear nuclei
- **C₂** Upper segment of the cervical spine
- **CN** Cochlear nucleus
Introduction

This chapter reviews anatomy and physiology of the auditory system, emphasizing structures and function that are likely to be involved with tinnitus. For general coverage of auditory anatomy and physiology of the auditory system, see [1]. The following chapter (Chap. 10) discusses pathologies of the auditory system.

Main Components of the Auditory System

The mammalian auditory system consists of four main parts: the apparatus that conducts sounds to the cochlea; the cochlea, where sounds are separated according to their frequency the sensory transduction occurs; the auditory nerve and the ascending auditory pathways, consisting of classical and non-classical including the primary and secondary auditory cortices; and association cortices with two streams (object or “what” and spatial or “where”) and extra-cortical structures that receive projections from the primary and secondary auditory cortices [1].

Anatomy

The most important parts of the auditory system, regarding most forms of tinnitus, are the function of the cochlea, the auditory nerve, and the central auditory nervous system.

The outer and middle ear that conducts sounds to the inner ear becomes important for tinnitus when impairments reduce the sound that reaches the cochlea where the sensory cells are located.

The cochlea is a very complex structure with essentially two fluid systems with different ionic composition (Fig. 8.1). The most important structures for transduction of sound into a neural code in auditory nerve fibers are the sensory cells (hair cells) that are located along the basilar membrane (Fig. 8.2). There are two kinds of sensory cells (inner and outer hair cells) (Fig. 8.1) that are morphologically similar but have complete different functions.

The auditory nervous system involves structures in the pons, midbrain, thalamus and cerebral cortex (Fig. 8.3).

Classical Pathways

The fibers of the auditory nerve terminate in the cochlear nuclei, which has three main divisions, the anterior and posterior ventral cochlear nuclei (AVCN
and PVCN), and the dorsal cochlear nuclei (DCN) (Figs. 8.4 and 8.5). Each fiber of the auditory nerve bifurcates and one of the branches divides again making it possible for each nerve fiber to connect to neurons in each of the three divisions of the cochlear nucleus (Fig. 8.5). This represents the first example of parallel processing allowing the same information to be processed in (three) different populations of nerve cells.

The sensory cells in the cochlea are innervated by the auditory nerve that conducts signals to the cochlear nucleus complex, which is the first nucleus of the ascending auditory pathways.

The cochlear nucleus has three main divisions: the dorsal cochlear nucleus (DCN), the anterior ventral cochlear nucleus (AVCN), and the posterior ventral cochlear nucleus (PVCN).

Cells in the three divisions of the cochlear nucleus project to the central nucleus of the inferior colliculus (ICC) through three fiber tracts, the dorsal (stria of Monaco), medial (stria of Held), and ventral striae (trapezoidal body) that joins into the lateral lemniscus (LL) (Fig. 8.6). Fibers in these three striae give off collaterals to the nuclei of the superior olivary complex and some fibers are interrupted in some of these nuclei.

There are two different ascending pathways from the cochlear nucleus to the cerebral auditory cortices, known as the classical and the non-classical pathways. The classical pathways also known as the lemniscal, or
Anatomy and Physiology of the Auditory System

the distinct pathways, are the best known. The non-classical pathways are also known as the extralemniscal, or diffuse, pathways. The main differences between these two pathways are in the thalamus.

The classical auditory pathways are mainly crossed, but there are extensive connections between the inferior colliculi at the midbrain level (Fig. 8.6) with the result that each one of the two side’s cerebral cortices receives approximately equal amounts of input from both ears. Sounds are thus represented bilaterally at the cortex, despite that the pathways are mainly crossed below the inferior colliculus. This has a clinical importance because lesions (tumors, strokes, traumatic injuries, etc.) on one side’s auditory cortex have only subtle clinical manifestations (normal audiograms, and only impaired speech discrimination for low-redundancy speech [6]). There are no known connections between the two sides’ thalamic nuclei.

The cells of the ICC project to the ventral part of the thalamic auditory nucleus, the medial geniculate body (MGB) that projects to the primary auditory cortex, and several other divisions of the auditory cortex, including the secondary cortex (AII) and the anterior and posterior auditory fields (AAF and PAF) (Fig. 8.7).

Cells in the inferior colliculus project to the auditory thalamic nucleus, the medial geniculate body (MGB). The thalamus plays a fundamental role in auditory processing. It consists of two different cell groups; one is the ventral part, belonging to the classical ascending pathways, which project to the primary auditory cortex. The other parts, the medial and dorsal parts, belong to the non-classical pathways and have fundamentally different functions. The cells project not to primary auditory cortex but bypass the primary cortex and connect directly to the secondary auditory cortex. In addition, these neurons connect to several parts of the brain, such as the amygdala, thus providing as subcortical route to the emotional brain.

Cells in the thalamic nuclei of the classical pathways project to the primary and secondary auditory
The Non-classical Pathways

While all neurons in the classical pathways up to and including the primary auditory cortex only respond to one sensory modality, some neurons in the non-classical pathways also respond to other sensory modalities. Likewise, other modalities of sensory input can change the response to auditory stimulation. This means that the non-classical auditory pathways receive input from other sensory systems, such as the somatosensory system and the visual system.

Anatomical Basis for Cross-modal Interaction

The fact that the non-classical ascending auditory system receives input from more than one system is the basis for sensory cross-modal interaction.

What was originally known as the extralemniscal system begins at the midbrain through connections from the ICC to two other parts of the inferior colliculus (IC), the external nucleus (ICX) and the dorsal cortex of the IC (DC) [7] (Fig. 8.8). The ICX also receives projections from the dorsal part of the spinal cord [8], thus providing input from the somatosensory system to the auditory pathways [9].

The results shown in Fig. 8.8 were based on animal studies with the cat. More recent anatomical studies in the guinea pig have shown that the external nucleus of the inferior colliculus (ICX) receives anatomically verified connections from the trigeminal ganglion¹

¹Some authors use the term “ganglion” for the trigeminal nucleus, but it seems more appropriate to use the term “nucleus” for clusters of nerve cells where synaptic communication occurs. The word “ganglion” should be reserved for clusters of cell bodies, such as the dorsal root ganglia. This is in accordance with the definitions by Stedman’s Electronic Medical Dictionary: A ganglion is: originally, any group of nerve cell bodies in the central or peripheral nervous system; currently, an aggregation of nerve cell bodies located in the peripheral nervous system.
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 mostly from the spinal (caudal) part of the fifth nerve nucleus [11].

The ICX, which is a part of the non-classical auditory pathways, receives anatomically verified connections from the trigeminal ganglion, as shown in studies in guinea pigs [10], mostly from the spinal (caudal) fifth nerve nucleus\(^2\) (Sp5) [11]. For a review of the influence on auditory processing in the brainstem from somatosensory activation, see Dehmel et al. [12].

\(^2\)In neuroanatomy, a nucleus is a group of nerve cell bodies in the brain or spinal cord that can be demarcated from neighboring groups on the basis of either differences in cell type or the presence of a surrounding zone of nerve fibers or cell-poor neuropil.

Cells in the ICX and the DC project to dorsal and medial parts of the thalamic auditory nucleus. These cells do not project to the primary auditory cortex as do the cells from the ventral part of the thalamus, but project to the AII division of the auditory cerebral cortex and to association cortices (Fig. 8.9). This means that the non-classical pathways skip a step in cortical activation compared with the classical pathways.

Fig. 8.6  More detailed drawing of the ascending auditory pathways from the ear to the central nucleus of inferior colliculus (ICC). \(AVCN\) anterior ventral cochlear nucleus, \(PVCN\) posterior ventral cochlear nucleus, \(DCN\) dorsal cochlear nucleus, \(LSO\) lateral superior olive, \(NTB\) nucleus of the trapezoidal body, \(MSO\) medial superior olive, \(SH\) stria of Held (intermediate stria), \(SM\) stria of Monakow (dorsal stria), \(LL\) nucleus of the lateral lemniscus, \(DNLL\) dorsal nucleus of the lateral lemniscus, \(VNLL\) ventral nucleus of the lateral lemniscus. ICC central nucleus of the inferior colliculus. (Modified from Møller, A.R., Sensory Systems: Anatomy and Physiology. 2003, Academic Press, Amsterdam. Reproduced with permission from Elsevier [5])
connect to many other structures such as the hypothalamus (Fig. 8.9).

The connections to limbic structures such as the amygdala are especially important in relation to tinnitus (see Chaps. 21 and 73). The classical and the non-classical pathways make important connections to the lateral nucleus of the amygdala through two different routes, known as the “high route” and the “low route” [13], respectively (Fig. 8.10). The low route uses a subcortical connection from the lateral and medial thalamus while the high route uses a long chain of neurons in the primary–secondary auditory cortices followed by neurons in several parts of the association cortices. Studies have indicated that the low route is not normally active in adults who do not have tinnitus, but there are indications that the non-classical auditory pathways are active in children [14] and in some individuals with tinnitus [15]. This means also that the subcortical connections to the amygdala are active in children and some individuals with tinnitus and also in other possible disorders [16] such as some forms of autism [17, 18].

**Other Anatomical Bases for Interaction between Senses**

These connections to nuclei of the IC from the spinal cord and the trigeminal nucleus were, for a long time, believed to be the only connections between the somatosensory system and the auditory system. Recent studies have shown that connections also exist from the somatosensory system to more peripheral levels of
the ascending auditory pathways. Thus, somatic sensory neurons project to several central auditory structures, including DCN, perhaps the VCN, and parts of inferior colliculus (IC) (the external nucleus). Both first- and second-order somatosensory neurons have been found to project to auditory structures [12, 20, 21]. The projections from primary structures originate from spinal dorsal roots, mainly C₂ but also more caudal roots, and the trigeminal nerve [22]. The projections from secondary somatosensory structures mainly originate from the spinal dorsal column nuclei, the caudal trigeminal nucleus (Sp5), and DRG [11]. Some neurons in the Sp5 and the dorsal column nuclei project to both the CN and the external cortex of IC by way of axon collaterals [23].

Histological studies in guinea pigs have shown that there are connections between the trigeminal nucleus, as well as the marginal cell areas of the cochlear nucleus and the magnocellular portion of the ventral cochlear nucleus. It is mostly the ophthalmic and mandibular divisions of the trigeminal nerve that give rise to these connections [24] and the DCN [21] (for more details see Chap. 9). (For a review of the influence on auditory processing in the brainstem from somatosensory activation see Dehmel et al. [12]).

Early studies involving cats showed direct projections from dorsal column nuclei and the spinal trigeminal nuclei to the cochlear nuclei [25]. Later studies using decerebrated paralyzed cats have shown that electrical stimulation of the dorsal column in the spinal cord and spinal trigeminal nucleus could inhibit the response from cells in the DCN [26]. Direct connections from the C₂ area of the dorsal horn of the spinal cord to the cochlear nuclei have been shown in anatomical studies [27].

The skin around the ears and of the scalp is innervated both by fibers of the C₂ dorsal root and by the trigeminal nerve. This may explain the beneficial effect of electrical stimulation on tinnitus when performed on these areas of skin [28].

There are also anatomically verified connections between the caudal trigeminal nucleus and the dorsal cochlear nucleus [24] (see Chap. 9), and there is also anatomical evidence that the trigeminal nucleus innervates the cochlea [29].

These more recent studies extend the anatomical basis for interaction between the auditory and somatosensory functions, which are the anatomical basis for the cross-modal interaction discussed below.

This also means that the non-classical pathways indeed use the lemniscal system, making the earlier used name “extralemniscal system” irrelevant. The name used in this book (non-classical) seems to be more appropriate.

Projection fibers from these somatic sensory neurons form a laminar pattern of “en passant terminal endings” from ventromedial to dorsolateral within the ventrolateral regions of ICX, including the ventral border of IC and the ventromedial edge of IC (or pericentral regions).
A study has shown a novel projection from the basolateral nucleus of the amygdala to the inferior colliculus in bats [30].

**Descending Systems**

Descending auditory systems have been described as having three different parts. It may be more suitable to describe the descending systems as reciprocal to the ascending systems (see Fig. 8.11) [31].

The axons of the most peripheral parts of the descending pathways (olivocochlear bundle, OCB, Fig. 8.12a, b) terminate on hair cells in the cochlea, mostly outer hair cells. Since these hair cells control mechanical properties of the basilar membrane, the descending pathways can influence the mechanical properties of the basilar membrane and thereby affect auditory sensitivity and frequency selectivity.
The basilar membrane of the cochlea separates sounds according to their frequency in such a way that the population of sensory cells that are activated is a direct function of the frequency (or spectrum) of the sounds that reaches the ear. One kind of hair cells, the inner hair cells, is activated by the motion of the basilar membrane and controls the discharges in the auditory nerve fibers. The outer hair cells are also activated by sound, but they have a mechanical role in that they elongate and shorten in response to sound, thereby acting as “motors” that amplify the motion of the basilar membrane. This action is most pronounced for low sound intensities where the action of the outer hair cells adds approximately 50 dB of sensitivity to the ear. The fact that their action is dependent on the intensity of the sounds that reach the ear makes the function of the cochlea become highly non-linear and the outer hair cells act to compress sounds. The active role of the outer hair cells also causes the cochlea to generate sound under certain circumstances (otoacoustic emissions) (for a review see Kim et al. [34]).

The non-linearity of the outer hair cells manifests by several measures used in clinical diagnosis. The active role of the outer hair cells can be detected by recording the otoacoustic emission of the ear by placing a microphone in the ear canal. There are several kinds of such otoacoustic emission. Spontaneous otoacoustic emission (SOE), the ear producing sound without receiving sound (in silence), is relatively rare. The most commonly studied kinds of otoacoustic emission that are also used clinically are transient elicited otoacoustic emission (TEOAE) and distortion product otoacoustic emission (DPOAE) (Fig. 8.13).

The Nervous System

The physiology of the classical pathways has been studied extensively, mainly by recordings from single auditory nerve fibers of the auditory nerve and from cells in the nucleus and the cerebral cortex. Little is known about the response from cells in the nuclei of the non-classical pathways. Only a few studies have concerned the physiology of the descending pathways and little is known about the function of these anatomically extensive systems.

When studied using pure-tone stimuli, all auditory nerve fibers and cells in the nuclei of the ascending auditory pathways display frequency selectivity. The sharpness of the auditory nerve’s fiber tuning varies systematically with the frequency to which they are tuned (their best frequency; see Fig. 8.14). The sharpness of the tuning also varies with the sound intensity as shown in Fig. 8.15.
that shows tuning of an auditory nerve fiber in response to broad-band noise stimuli, and Fig. 8.15b shows the sharpness of the tuning of an auditory nerve fiber is an almost linear function of the sound intensity. The frequency to which the fiber is tuned decreases with increasing sound intensity. The sharpness of the tuning of the basilar membrane is greater for low-intensity sounds compared with sounds of higher intensity (Fig. 8.15c), thus following a similar pattern as the tuning of auditory nerve fibers, indicating that the cause of the non-linearity seen in the tuning of auditory nerve fibers is some property of the cochlear mechanics.

The shape and sharpness (frequency selectivity) of the frequency tuning of cells in the nucleus and the cerebral cortex vary among cells. Some have a higher degree of selectivity than auditory nerve fibers, other cells are more broadly tuned, and the tuning curves of some cells have more than one peak (Fig. 8.16). When more complex sounds are used, the response pattern becomes different from that obtained to steady test sounds, as has been shown for cells in the cochlear nuclei. These cells show more complex response pattern to sounds, the frequency of which is varied at different rates. The response becomes dependent on rate of tonal change (Fig. 8.17). This means that the response to complex sounds cannot be deduced from knowledge about the response to simple sounds such as pure tones with steady frequency. This non-linear behavior is apparent in the responses from cells in the cochlear nucleus and it becomes more pronounced in the responses from cells of higher order nuclei of the auditory system, including the cerebral cortices.
Moreover, because cells in auditory nuclei and the cerebral cortex receive input from many fibers, the width and the center frequency of their tuning become dependent on the efficacy of the synapses that connect the fibers to the cells. Since synaptic efficacy can be altered by activation of neural plasticity, the tuning of cells is not static but subject to change.

**Non-classical System**

Much less is known about the physiology of the non-classical pathways. Few studies have been published on the response from cells in the nuclei that belong to the non-classical pathways as opposed to those belonging to the classical pathways. It is known, however, that the responses of cells in the non-classical pathways are generally less distinct, and tuning is broader in neurons of the non-classical pathways compared with the classical pathways. Perhaps the most important aspect of the auditory non-classical pathways for understanding the pathologies of tinnitus is the interactions between signals from other sensory systems that occur causing cross-modal interactions; the effect of the use of the medial and dorsal thalamic nuclei with their subcortical connections; and the absence of connections to the primary cortices.

**Cross-modal Interaction**

It has become more and more evident that there is considerable interaction between systems that earlier were regarded as being separate. The old concept that certain functions of the brain are contained in certain areas of the brain has gradually been eroded. It was earlier regarded as an axiom that the information from the different sense organs was processed in specific and separate parts of the brain. That somatosensory signals can interfere with hearing is discussed in several parts of this book. The anatomy of sensory systems as described above involves both the non-classical pathways and connections between the dorsal column nuclei and the cochlear nucleus, which also receives connections from the trigeminal ganglion. Physiological
Fig. 8.15 (a) Estimates of frequency transfer function of a single auditory nerve fiber in a rat at different stimulus intensities (given in dB SPL), obtained by Fourier transforming cross-correlograms of the responses to low-pass–filtered pseudorandom noise (3,400 Hz cutoff). The amplitude is normalized to show the ratio (in dB) between the Fourier-transformed cross-correlograms and the sound pressure and the individual curves would have coincided if the cochlear filtering and neural conduction had been linear. Modified from Møller, A.R. 1983. Frequency selectivity of phase-locking of complex sounds in the auditory nerve of the rat. Hear. Res. 11, 267–284. [37]. Reproduced with permission from Elsevier. (b) Shift in the center frequency (solid lines) and the width of the tuning of a single auditory nerve fiber (dashed line) in the auditory nerve of a rat as a function of the stimulus intensity. The width is given a “Q10dB” which is the center frequency divided by the width at 10 dB above the peak (Reprinted from Møller, A.R. 1977. Frequency selectivity of single auditory nerve fibers in response to broadband noise stimuli. Reproduced from J. Acoust. Soc. Am. 62, 135–142, with permission from the American Institute of Physics [38]) (c) Vibration amplitude at a single point of the basilar membrane of a guinea pig obtained using pure tones as test sounds at four different intensities. The amplitude scale is normalized, and the individual curves would have coincided if the basilar membrane motion had been linear. From Johnstone, B.M., Patuzzi, R., Yates, G.K. 1986. Basilar membrane measurements and the traveling wave. Hear. Res. 22, 147–153 [Johnstone, 1986 #1116] based on results from Sellick, P.M., Patuzzi, R., Johnstone, B.M. 1982. Measurement of basilar membrane motion in the guinea pig using the Mossbauer technique. J. Acoust. Soc. Am. 72, 131–141. [39]. Reproduced with permission from the American Institute of Physics
Fig. 8.16 Examples of frequency tuning curves with different shapes obtained from neurons of the superior olivary complex of the cat. From Guinan Jr., J.J., Norris, B.E., Guinan, S.S. 1972. Single auditory units in the superior olivary complex. II. Location of unit categories and tonotopic organization. Int. J. Neurosci. 4, 147–166. [40]. Reproduced with permission from Elsevier

Fig. 8.17 Period histograms of a cell in the cochlear nucleus of a rat in response to tones the frequency of which was varied between 5 and 25 kHz at different rates. (a) and (c): Slow rate, (b) and (d): Fast rate. The top histograms (a and c, slow rate) show the responses obtained when the duration of a full cycle was 10 s and the lower histograms (b and d, fast rates) show the responses obtained when the duration of a complete cycle was 156 ms. The change in the frequency of the stimulus tone was accomplished by having a trapezoidal waveform control of the frequency of the sound generator (e). The two left-hand graphs (a and b) are histograms of a full cycle of the modulation and the right-hand graphs (c and d) show the details between the vertical lines in the left-hand graphs. From Møller, A.R. 1974. Coding of sounds with rapidly varying spectrum in the cochlear nucleus. J. Acoust. Soc. Am. 55, 631–640. [41]. Reproduced with the permission from the American Institute of Physics
A.R. Møller

studies have shown that the non-classical auditory pathways are active in children [14] and in some individuals with tinnitus [15].

As mentioned above, it has been shown in several studies that there are anatomically verified connections between the trigeminal nucleus and the dorsal cochlear nucleus, as well as between the upper part of the spinal cord and the dorsal cochlear nucleus.

Physiological studies in animals (guinea pigs) have shown that multisensory integration occurs in the DCN, thus confirming that the anatomical connections discussed above are also physiologically active. This means that the synapses that connect the axons to the cells in the DCN are effective. Electrical stimulation of the trigeminal ganglion elicited unit responses in the VCN with latencies from 5 to 17 ms, indicating that the anatomically verified pathway from the trigeminal ganglion to the VCN is functionally active [42, 43].

These connections between the somatosensory system and the cochlear nucleus are most likely the basis for findings that electrical stimulation of the skin around ears suppresses tinnitus in some individuals [44] (see Chap. 43). Such stimulation would therefore activate both the dorsal gray of the upper spinal cord and the spinal nucleus. The observed effect on tinnitus may therefore come from either spinal connection to cochlear (DCN) nuclei or the trigeminal nucleus.

The effect of electrical stimulation of the skin around the ears has been verified in animal experiments.

After exposure to loud tones, recordings from cells in the DCN of hamsters showed that such exposure caused hyperactivity. When the skin at the base of the ears was stimulated electrically, the response would be either depressed in both control animals and exposed animals; excited in controls and depressed in tone-exposed animals; depressed and excited; or excited in both controls and tone exposed [45]. The suppression was significantly higher during and after stimulation than in controls, with the effect slightly greater after the stimulation than during stimulation.

The pathways for this cross-modal interaction have been studied in animals and it has been shown that there is both physiological and anatomical evidence for cross-modal interaction in the dorsal cochlear nucleus [43].

In one study in hamsters using tracing experiments, it was found that the DCN received input from the spinal trigeminal nucleus (that is commonly regarded to be involved in face pain), but also the dorsal raphe nucleus seems to be involved along with the locus coeruleus [46].

These studies indicate that the cross-modal interaction in the dorsal cochlear nucleus may involve a direct and an indirect pathway. Therefore, relieving tinnitus through somatosensory electrical stimulation may involve manipulations of both auditory and non-auditory neural circuits.

It is mainly somatic receptors in the upper body that can modulate activity elicited by sound stimulation in the neurons of the non-classical auditory pathways [7]. In order for this to occur, the non-classical auditory pathways must be active. The non-classical auditory system has the ability of cross-modal interactions because neurons in that system receive signals from more than one sensory modality. There are no signs of the non-classical auditory system being active in adults who do not have tinnitus. Adults who do not have tinnitus do not have signs that the non-classical pathways are active. This means that some forms of cross-modal interactions require re-routing of information, so that the non-classical auditory system becomes active. As mentioned above, there are signs of such cross-modal interaction between the somatosensory system and the auditory system in young children [14], in some individuals with tinnitus [15], and in some autistic individuals [17, 18].

This means that there is physiologic evidence that multisensory integrations occur in the DCN in guinea pigs [43, 47], in the ICX in cats [7], and the ICX in rats [48]. There may be differences between the animals that have been studied and humans; it is not known how these results from animals can be applied to humans. Studies in humans of the ability of somatosensory stimulation to affect sound perception, in terms of loudness, seem to show that such interaction occurs normally in children but not in adults [15]. Stimulation of the somatosensory system can affect the loudness and character of tinnitus in some individuals [15], and there are other studies that show evidence that somatosensory activation can affect the loudness and the character of tinnitus [49–51]. Studies have also shown evidence of interaction between the somatic and visual systems [52].

Conclusions

The description of the anatomy and physiology of the auditory system in this chapter covers what was regarded to be relevant regarding tinnitus. There are
many aspects of the anatomy and physiology that are not included, and which were judged to be less related to the topic of this book. A more detailed description of the anatomy and physiology of the auditory system can be found in Møller, A. R. (2006) Hearing: Anatomy, Physiology, and Disorders of the Auditory System, 2nd Ed. Academic Press: Amsterdam [1].

References


Chapter 9
Interaction Between Somatosensory and Auditory Systems

Aage R. Møller and Susan Shore

Keypoints

1. Studies in animals (guinea pigs) have shown projections from the dorsal column nuclei and the caudal trigeminal nucleus to cells in the cochlear nucleus (CN).
2. Recordings from single cells in the DCN and evoked potentials indicate that the pathways from the trigeminal nucleus are functional.
3. Electrical stimulation of the dorsal column and the cervical dorsal root ganglia elicits short and long latency inhibition separated by a transient excitatory peak in DCN single units.
4. Electrical stimulation of the trigeminal nucleus elicits excitation in some DCN units and inhibition in others.
5. Dorsal cochlear nucleus neurons show greater sensitivity to somatosensory stimulation, and the interaction between somatic stimulation and sound stimulation is greater after exposure to loud sounds that cause hearing loss and probably tinnitus. These findings may be explained by increased innervation of the cochlear nucleus by somatosensory fibers after noise exposure.

Keywords
Tinnitus • Cross-modal interaction • Cochlear nucleus • Trigeminal system • Dorsal column system

Abbreviations

<table>
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<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>ANF</td>
<td>Auditory nerve fibers</td>
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<td>CN</td>
<td>Cochlear nucleus</td>
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<td>DCN</td>
<td>Dorsal cochlear nucleus</td>
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<td>GCD</td>
<td>Granule cell domain</td>
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<td>DRG</td>
<td>Dorsal root ganglion</td>
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<td>PSTH</td>
<td>Post stimulus time histogram</td>
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<td>Sp5</td>
<td>Spinal trigeminal nucleus</td>
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<td>Sp5C</td>
<td>Caudal spinal trigeminal nucleus</td>
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<td>TG</td>
<td>Trigeminal ganglion</td>
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<td>TN</td>
<td>Trigeminal nucleus</td>
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<td>VCN</td>
<td>Ventral cochlear nucleus</td>
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Introduction

It has been known for many years that information from different senses are coordinated in many parts of the cerebral cortex, but it is only recently that it has become evident that different senses interact with each other at subcortical levels. One of the routes of sensory interaction is the connection between the dorsal root ganglion or dorsal column nuclei [1] and the cochlear nucleus (CN); another is the connection between the trigeminal ganglion or caudal trigeminal nucleus (Sp5C) and cochlear nuclei (CN) [1–3]. These connections are the anatomical basis for some of the cross-modal interactions observed in animal experiments, in humans, and in some individuals with tinnitus (see Chap. 10).

It is of particular interest that the interaction between stimulation of the somatosensory system and sound stimulation is enhanced by previous intense sound overstimulation of the kind that normally results in tinnitus [4]. In this chapter, we will discuss the anatomical and physiological bases for cross-modal...
interaction and the possible relationship to tinnitus by
the subcortical connections between the somatosensory
system and the auditory system. Histological and phys-
iological studies in animals will be reviewed. Interactions
between the auditory system and other systems in gen-
eral were discussed in the preceding chapter (Chap. 8).

Anatomical Basis for Interaction
Between the Somatic System and the CN

Histological studies in guinea pigs have shown that
there are connections between the trigeminal system
(trigeminal ganglion and trigeminal nucleus) and the
marginal cell areas of the CN, and the magnocellular
portion of the ventral cochlear nucleus (VCN), and the
dorsal cochlear nucleus (DCN) [5]. Studies of the
functional role of these connections using electrical
stimulation of the trigeminal ganglion (TG) showed
responses from cells in the CN, indicating that these
anatomically verified pathways from the trigeminal
ganglion to the VCN can be activated [6, 7]. The laten-
cies varied considerably, from 5 to 17 ms, indicating
that different pathways may be involved.

Several studies have found connections between the
dorsal column of the spinal cord and cells along the
medial edge of the VCN, the dorsal ridge of the anterior
ventral cochlear nucleus (AVCN) (i.e., subpeduncular
corner between the AVCN and the inferior cerebellar
peduncle), and lamina of the granule cell domain
(GCD) [1, 3]. These axons of the dorsal column origi-
nate from the C2 dorsal roots of the spinal cord.
Likewise, it has been found that electrical stimulation
of the dorsal root ganglion elicits responses from cells
in the DCN [8]. The projection from somatosensory
nuclei to the CN is particularly abundant in granule
cell regions in layer 2 of the DCN. The small cell cap
region of CN and also larger cells in deep DCN receive
projections from cells in the Sp5 [5, 9] and the dorsal
column nuclei [5, 10–14].

The dorsal column nuclei receive innocuous soma-
tosensory input and proprioceptive sensory input
[15, 16]; caudal trigeminal nucleus (Sp5C) receives pain
and temperature information from regions of the face
and mouth. This nucleus also receives proprioceptive
signals from the vocal tract, including the temporo-
mandibular joint and tongue muscles [17]. The sub-
stantia gelatinosa layer of the Sp5C that is analogous
to the lamina II in the dorsal horn of the spinal cord
primarily receives nociceptive afferents.

Effects of Trigeminal Nerve
Activation on CN Activity

Studies in the guinea pig have shown that electrical
stimulation of the ophthalmic/mandibular divisions of
the trigeminal ganglion where neurons project to the
CN [18] elicit responses from neurons in the VCN [6]
and excite (Fig. 9.1) or inhibit neurons in the DCN
(Fig. 9.2) [7, 19].

In the post-stimulus histograms of the response from
cells in the DCN shown in Figs. 9.1 and 9.2, the stimulus
artifact is seen at 25 ms, indicating the time at which
the electrical stimulus was applied to the TG. The four chan-
nels of recordings were obtained simultaneously and, in
the example illustrated in Fig. 9.2, the electrical stimula-
tion of three of these had an inhibitory effect on sponta-
neous activity. In this recording, the latency of the
inhibition was approximately 12 ms; in other similar
recordings, the latency varied between 5 and 20 ms, indi-
cating multisynaptic pathways would be the underlying
neural circuits producing these responses [7].

The results of these studies show that the trigeminal
pathways can alter the spontaneous activity in both the
VCN and the DCN. This change in spontaneous activ-
ity of CN neurons might explain tinnitus’ loudness
modulation from trigeminal system stimulation
observed in some individuals with what is known as
somatic tinnitus.

Other studies [8] using recordings of evoked poten-
tials have shown that electrical stimulation of spinal
nerves in the neck can elicit a response from cells in the
DCN. The C2 root mediates mechanoreception and
proprioception of the pinna and the surrounding skin.
The evoked potentials were largest in response to stimu-
lation of the cervical nerves corresponding to mecha-
noreception and proprioception in the pinna (C2), neck
(C3), and forelimbs (C7) [8]. These results show a simi-
larity with the effect of TG stimulation [6]. Other stud-
ies have shown that stimulation of the femoral nerve
can activate cells in the DCN as evidenced by fos
expression [21].

Combining electrical stimulation of the trigeminal
systems with sound stimulation shows that these two
kinds of activation of CN neurons interact in a nonlinear
Interaction Between Somatosensory and Auditory Systems

One modality can inhibit or enhance the response to another modality, depending on the time interval between the sound and the electrical stimulations (Fig. 9.3).

It is seen from Fig. 9.3 that the interaction between these two stimuli depends on the interval between the stimuli. When the acoustic stimulus precedes the TG stimulation, the response to the acoustic stimulus is suppressed, the effect being the greatest when the two stimuli are presented within a small interval. Somatosensory stimulation can alter the firing rate to acoustic stimulation, even when it precedes the acoustic stimulation by as much as 90 ms and stimulation and the effect still persist for the duration of the sound stimulation [7].

The interaction between these two kinds of stimulation is different in different cells as illustrated in Fig. 9.4b, which shows the discharge rate as a function of the interval between sound stimulation and the stimulation of the TG for three different cells in the DCN. The firing rate of one of the cells increases when the interval between the stimuli is increased from 20 to 95 ms. Another cell shows a depression that is the largest when the interval is 60 ms; the third cell depicted in Fig. 9.4 has very little effect of the combination of the two stimuli.

Other studies of the interaction between stimulation of the somatic sensory system and sound [8, 22, 23] have shown that sound-evoked responses are influenced by electrical stimulation of the dorsal column of the spinal cord in a similar way as stimulation of the trigeminal system. The responses from cells in the DCN to electrical activation of dorsal
root ganglia (DRG) are similar to those obtained by stimulation of dorsal column nuclei, but they have longer latencies [8].

**Interaction Between the Somatosensory System and the DCN After Sound Over Stimulation**

It is of particular interest regarding tinnitus that the interaction between the somatosensory system and the auditory system is larger after sound overstimulation that causes hearing loss and probably tinnitus [4].

Studies in the guinea pig have shown that the bimodal interaction was enhanced after noise exposure. The interaction between somatic and auditory stimulation in animals that have not been exposed to loud sounds varies among different cells, some showing enhancement and some showing suppression as shown above (Figs. 9.1 and 9.2). This is also the case in noise-exposed animals. However, two weeks after noise exposure, more cells showed suppression than in animals that were not exposed to noise, 75% compared with 49%. This difference developed over time after noise exposure, and it was very small, one week after noise exposure.

Noise exposure also decreased the threshold to trigeminal stimulation, and the spontaneous discharge

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rates of cells in the DCN increased [4]. The increase in the responses from cells in the DCN to somatosensory stimulation may be a compensation for lost auditory input because of the hearing loss caused by the sound exposure. The duration of the inhibitory response to trigeminal stimulation decreased and its amplitude increased after noise exposure (Fig. 9.4).

The observed changes in the response properties of DCN cells after noise exposure might be caused by activation of neural plasticity that has strengthened the synaptic coupling to the somatic source of input to some cells in the DCN. There are other examples that show that loss of one kind of sensory input has enhanced input from other senses. The most profound example might be invasion of visual fibers into the auditory cortex after interruption of auditory input [24].

We believe that the observed suppression of the responses to broad band noise is a result of the summation of weak responses from cartwheel cells to the noise and stronger and long-lasting activation of these cells by the input from the trigeminal nucleus, which in turn leads to inhibition of fusiform cells. Facilitation of the response to noise stimuli, on the other hand, may occur because of long-term potentiation of direct activation of fusiform cells by granule cells. Cartwheel cells excite each other and inhibit fusiform cells (see Fig. 9.6). Stimulation of the TG may excite cells in the ventral cochlear nucleus (multipolar onset cells), which can inhibit vertical cells and fusiform cells [6].

The circuits of the DCN relevant to interactions from somatosensory nuclei are shown in Fig. 9.6, which illustrates the situation before (a) and after (b) noise exposure.

Based on their findings that the vesicular glutamate transporters (VGLUT1 and VGLUT2) are differentially associated with auditory nerve and somatosensory inputs to the CN, respectively [25], Zeng et al. [26] examined the relative distributions of VGLUT1 and 2 after unilateral deafening. After unilateral intra-cochlear injections of kanamycin (1 and 2 weeks), VGLUT1 immunoreactivity in the magnocellular CN ipsilateral to the cochlear damage was significantly decreased, reflecting decreased auditory nerve input as expected. On the other hand, VGLUT2, which is associated with the non-auditory inputs, including somatosensory inputs, increased in regions that received non-auditory input 2 weeks after deafening, suggesting the possibility of axonal sprouting of these somatosensory inputs to the CN. These morphologic changes may be the cause of the observed increase in the response to trigeminal stimulation (Fig. 9.5).

These results should be viewed in the light of earlier studies that have shown that noise exposure causes morphologic changes in the cochlear nucleus [27]. Other studies [5] have shown enhanced cartwheel cell activity after noise damage, which could be responsible for the increase in suppressive bimodal integration demonstrated above [4].
Fig. 9.4 Quantification of the spike rates achieved in (a) for unit 16. Spike rates for two other units, 5a and 5b, are also shown. Insets: Poststimulus time histograms for responses to BF tone bursts indicate unit types: Unit 16 is a P-Buildup, Unit 5b may be a cartwheel cell (Reproduced from Shore, S., Zhou, J. and Koehler, S. (2007). Neural mechanisms underlying somatic tinnitus. In: Tinnitus: Pathophysiology and Treatment, Progress in Brain Research. pp. 107–123. Eds. B. Langguth, G. Hajak, T. Kleinjung, A. Cacace, A. R. Møller. Elsevier: Amsterdam. [20])

Fig. 9.5 Responses to trigeminal stimulation are redistributed at 1 and 2 weeks after noise overexposure. The percentage of single units with excitatory, inhibitory, or excitatory/inhibitory responses after trigeminal nucleus stimulation at 80 μA is shown. Following noise exposure, inhibitory responses predominate, whereas the normal animals show more excitatory than inhibitory responses. The increased incidence of inhibition by trigeminal stimulation in noise-damaged animals may signify a change in the distribution of trigeminal inputs to the cochlear nucleus granule cells following cochlear damage. (Reproduced from: Shore, S. E., Koehler, S., Oldakowski, M., Hughes, L. F. and Syed, S. (2008) Dorsal cochlear nucleus responses to somatosensory stimulation are enhanced after noise-induced hearing loss. Eur J Neurosci. 27, 155–168). [4] Reproduced with permission of Wiley
Fig. 9.6 Schematic of dorsal cochlear nucleus circuitry putatively involved in bimodal integration in normal and noise-damaged animals (a) Normal system. Trigeminal nucleus (Tg) stimulation excites cochlear nucleus granule cells (gr), which, in turn, excite stellate (St), cartwheel (Ca), and fusiform (Fu) or giant (not shown) cells. DAS dorsal acoustic stria, p.f. parallel fibers. (b) Noise-damaged system. ANF input to basal dendrites of Fu cells is weakened. (Reproduced after Shore, S. E. (2005) Multisensory integration in the dorsal cochlear nucleus: unit responses to acoustic and trigeminal ganglion stimulation. *Eur J Neurosci* 21, 3334–3348. [7] and from: Shore, S. E., Koehler, S., Oldakowski, M., Hughes, L. F. and Syed, S. (2008) Dorsal cochlear nucleus responses to somatosensory stimulation are enhanced after noise-induced hearing loss. *Eur J Neurosci.* 27, 155–168). [4]

**Conclusion**

Studies in the guinea pig have shown that stimulation of the dorsal column of the upper spinal cord or the trigeminal ganglion or nucleus can elicit excitation or inhibition of neurons in the cochlear nuclei. These studies also indicate that such stimulation can modulate the sound-driven activity in the cochlear nuclei. The effect of somatosensory stimulation on the neural activity in the cochlear nuclei is greater in animals that have noise-induced hearing loss and probably tinnitus when studied 2 weeks after the noise exposure, but not immediately after noise exposure (Fig. 9.5).

**References**

Keypoints

1. Symptoms such as tinnitus can be caused by damage and diseases that affect the conductive apparatus of the ear, its receptor organs, the auditory nerve, and nerve cells in the nuclei of the auditory system, including the cerebral auditory cortex.
2. Tinnitus can also be caused by activation of neural plasticity (causing plasticity diseases), which can cause altered function at the cellular level in the brain and re-routing of information.
3. The brain is not a fixed system but it is continuously shaped and reshaped by signals it receives from the outside world.
4. Neural plasticity is a property of the nervous system that becomes apparent only when turned on. Activation of neural plasticity can be beneficial or harmful.
5. Activation of beneficial neural plasticity facilitates recovery from damage to the nervous system (such as from strokes). In sensory systems, it may serve to compensate for loss of function or to adapt the nervous system to change in demand. Expression of neural plasticity can make the nervous system adapt to changing demands (prostheses such as cochlear and cochlear nucleus implants).
6. Activation of harmful neural plasticity is involved in creation of symptoms of disease (plasticity disorders) such as some forms of tinnitus, central neuropathic pain, and some forms of muscle spasm.
7. Activation of neural plasticity can change processing of information and cause:
   (a) Reorganization and re-routing of information in the central nervous system.
   (b) Change in the balance between inhibition and excitation.
   (c) Increased synchrony of activity of single nerve cells.
   (d) Increased temporal coherence of activity in populations of nerve cells.
8. Deprivation of input, overstimulation, injuries, and unknown intrinsic factors can promote expression of neural plasticity.
9. Many forms of tinnitus are phantom sensations caused by activation of neural plasticity and similar to phantom sensations in other sensory systems causing central neuropathic pain, paresthesia, and spasm in motor systems.
10. Many forms of tinnitus are associated with changes in processing of information that may involve hyperacusis and distortion of sounds.
11. Abnormal (pathologic) changes in connectivity may occur because of activation of neural plasticity that opens (unmask) dormant synapses or close (mask) synapses that are conducting normally.
12. Activation of non-classical pathways is an example of change in connectivity.
13. Tinnitus is often accompanied by cross-modal interaction, which may be explained by an abnormal activation of non-classical sensory pathways through re-routing of information.
14. Involvement of the non-classical pathways may explain symptoms of mood disorders, phantom sensations, improved perceptual capabilities, or atypical sensory experiences that often accompany severe tinnitus.
Introduction

Since tinnitus appears as a sound, it is often referred to the ear. For many years, the ear was therefore assumed to be the anatomical location of the pathology that caused subjective tinnitus. Jürgen Tonndorf was one of the first investigators who proposed a neurophysiologic cause for some forms of tinnitus and suggested a model for generation of tinnitus involving the central nervous system [1]. It was a major progress in understanding the pathology of many forms of tinnitus when it became more generally accepted that most forms of tinnitus are phantom sounds caused by abnormal function of neural circuits in the brain [2]. It is now evident that while the pathology that causes tinnitus may start with an event involving the ear, the pathology that causes most forms of persistent subjective tinnitus is in the central nervous system where some abnormal neural activity is generated and interpreted in a similar way as activity generated when sound reaches the ear.

Many different factors have been suspected to be involved in causing tinnitus, such as pathologies of the ear, the auditory nerve, and various parts of the central auditory nervous system. Damage to the auditory nerve from trauma, including surgical trauma and ionized radiation, are common causes of tinnitus, as are viral infections. It is not known exactly how trauma to the auditory nerve can cause tinnitus. One hypothesis has been that ephaptic transmission between denuded auditory nerve fibers (see Chap. 84) occur after trauma and, more recently, it has been suggested that such transmission may occur between nerve cells in the auditory nervous system.

Factors such as overstimulation and deprivation of signals to the auditory nervous system have also been suggested as causing tinnitus through activation of neural plasticity.

Subjective tinnitus has many similarities with phantom sensations from other senses, such as paresthesia of the somatosensory system, and in particular, with central neuropathic pain, which we will discuss in another chapter (see Chap. 14).

Similar “phantom sensations” as tinnitus rarely occur in vision (phosphene), olfaction (phantosmia), olfactory hallucinations, or abnormal taste (metallic taste). In the vestibular system, some forms of vertigo may be phantom sensations.

There is considerable evidence that activation of neural plasticity is involved in many forms of tinnitus [3–6] (see Chap. 12). We are, however, far from fully understanding the nature of the abnormalities that cause many forms of tinnitus, and how it is brought about is still being investigated; hypotheses regarding the pathology of tinnitus are constantly created and abandoned [7].

It has been pointed out in several places in this book that tinnitus is not a single disorder but many different ones. This is a major obstacle in attempts to understand its pathology, as it is for developing effective treatments. This means that the pathologies that can cause tinnitus are likely numerous and the anatomical location of the pathologies of these different kinds of tinnitus varies between the ear, the auditory nerve, and many different parts of the brain.

The pathological processes involved are poorly understood, but they may be different for varying kinds of tinnitus. Imbalance between inhibition and excitation has been suggested. Ephaptic transmission between axons in the auditory nerve has been suggested [7, 8], but some forms of ephaptic transmission between nerve cells in the central nervous system may also be involved. It is known that such ephaptic transmission plays a role in some forms of epilepsy [9]. Increased temporal coherence may be promoted by abnormal (ephaptic) transmission between auditory nerve fibers mimicking sound activation or between cells in the central nervous system.

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1Ephaptic transmission is transmission between axons and nerve cells without synaptic transmission. It may occur when bare axons or nerve cells are in close contact with each other so that one can activate the other electrically.
Anatomical Location of the Physiologic Abnormalities

The symptoms and signs of most diseases occur not because of a single event or because of damage to a single structure; the symptoms and signs of most disorders are caused by a cascade of events that occur at the same time, although there may not have been any signs or symptoms if they had occurred alone or one at a time. Most disorders of the nervous system involve a cascade of structures in the brain. While only one structure is pathologic, many structures may behave abnormally because they receive abnormal signals from the pathologic structures. It is an obstacle to research and treatment of many sensory disorders such as tinnitus and pain that several brain structures may behave abnormally even when only one of the structures is pathologic [10, 11]. The reason is that a faulty structure of a sensory system sends abnormal signals to other structures, which then behave abnormally, either because they relay abnormal activity or because the abnormal activity has affected the function of the structure in question.

Many mistakes in the diagnosis and the treatment are done because the treatment has been directed to the wrong structure.

There are other reasons why focus can be directed to the wrong structure. Phantom sensations are by definition not referred to the structure where the abnormal neural activity causing the sensations is generated. This is most obvious from the phantom limb, where a person feels pain in a leg that has been amputated. In a similar way, the pain in central neuropathic pain may be referred to a specific body part, but the abnormal neural activity is generated in the brain without any input from sensors in the body. Naturally, tinnitus is often referred to the ear and that can be the case even when the individual is deaf or has had the auditory nerve severed.

It was a major progress in understanding the pathophysiology of tinnitus when it became accepted that most forms of subjective tinnitus are caused by abnormal neural activity in the nervous system occurring without input from the ear. The fact is that tinnitus can occur in deaf individuals and after severance of the auditory nerve lends strong evidence to the hypothesis that the anatomical location of the physiologic abnormality is the central nervous system in many individuals with tinnitus, and input from the ear is not involved, at least not in later stages of the disorder.

In the hypothetical example in Fig. 10.1, the abnormal activity was caused by neural plasticity that was activated by deprivation of input and which activated a chain of structures. The abnormal activity in this chain of structures will subsequently reach neurons in the parts of the brain that generate the symptoms of tinnitus.

It is important to identify the structure in the beginning of the chain that is pathologic. Aiming treatment at neural structures that are functioning normally but produce abnormal input because they receive abnormal input is not effective in treating the disease in question.

Aiming treatment at structures that behave abnormally because they receive signals from structures that are pathologic may affect the perception of tinnitus because it may interrupt the flow of the abnormal signals that cause the tinnitus but will not yield permanent relief because the pathology is still unaffected. Such treatment may, however, ameliorate the symptoms as long as the treatment is applied. Similar situations exist regarding other common diseases, such as central neuropathic pain (see Chap. 14) and diabetes type 2 (Fig. 10.2).

Often two factors (or more) must be present at the same time to cause symptoms and signs of disease (Fig. 10.3). One example is diabetes type 2, where a cascade of events in a chain of structures result in changes that produce symptoms of diabetes neuropathy characterizing the disease.

**Fig. 10.1** Hypothetical flow chart of events in a series of structures as a result of deprivation of input. From Möller AR. *Neural plasticity and disorders of the nervous system*. Cambridge: University of Cambridge Press, 2006. Reproduced with permission from Cambridge University Press. Neurological Research with permission from W.S. Maney and Son Ltd [10]
The Ear

There are two ways the ear can be involved in causing tinnitus; one is by producing the kind of neural activity in the auditory nerve that is interpreted by the nervous system as a sound. The other way the ear can cause tinnitus is by depriving input to the nervous system that will turn on neural plasticity. Injury to the ear (see Chap. 34) may cause tinnitus because of the deprivation of input to the auditory nervous system it causes and which is known to be able to cause hyperactivity in specific structures through activation of neural plasticity [6, 12, 13]. Cochlear damage such as from noise exposure (see Chap. 37) or age-related changes (see Chap. 36) can also cause deprivation of input to the auditory system. Deprivation of input is a strong promoter of neural plasticity, which in turn causes changes in the nervous system that involves the generation of neural activity that may be interpreted as sound-evoked neural activity.

Studies have shown that passing electrical current through the cochlea in some patients with tinnitus can ameliorate their tinnitus [14, 15]. This is a strong sign that the ear is involved in causing and maintaining some forms of tinnitus.

Another sign that the ear is involved in causing some forms of tinnitus is the success of severing the auditory nerve in treatment of the disorder [16, 17]. This is, however, a controversial matter and other investigators have found that severing or damaging the auditory nerve does not affect tinnitus and may, in fact, make tinnitus worse [18].

If tinnitus is caused by deprivation of input to the nervous system, passing electrical current through the cochlea [15, 19] that may activate either hair cells or auditory nerve fibers and may thus compensate for deprivation.

Such electrical stimulation applied to the cochlea may, however, also activate the trigeminal nerve fibers, of which innervate the mucosa that lines the middle ear cavity including the cochlea capsule. Other studies have shown that electrical stimulation of the cochlear capsule can alleviate some forms of tinnitus [20]. Again, that may not have been caused by stimulating auditory receptors but instead by stimulating receptors in the mucosa of the middle ear cavity. These receptors are innervated by the trigeminal nerve fibers that terminate in the trigeminal nucleus. The cells these fibers terminate on project to cochlear nuclei known to be involved in tinnitus [21–23] (see Chaps. 8 and 9).
The ear is probably the cause of the tinnitus in only a few of those individuals who have severe chronic tinnitus, and severance of the auditory nerve is rarely done now. Pulec has emphasized [16, 17] that the auditory nerve must be sectioned centrally to the spiral ganglion in order to relieve tinnitus. This may mean that disconnecting the ear may not be the (entire) reason for the success of auditory nerve sectioning in tinnitus, and it indicates that the cause of the tinnitus may be the auditory nerve rather than the ear in some of the individuals who benefitted from this procedure.

**The Auditory Nerve**

Traumatic injury to the auditory nerve almost always causes tinnitus. This may occur in surgical operation and head trauma. Surgical trauma and ionized radiation used to treat vestibular schwannoma may cause loss of hearing and tinnitus, but moderate injury to the auditory nerve may cause only little change in the hearing threshold with a large decrease in speech discrimination and tinnitus. Viral infections, such as the herpes zoster virus, can affect the auditory nerve and cause hearing loss and tinnitus. The Ramsay Hunt syndrome is caused by the herpes zoster virus, and although it primarily affects the facial nerve, it can also affect the auditory-vestibular nerve and cause tinnitus [24]. Other viral infections, such as the Coxsackie B virus, have also been reported to cause tinnitus [25].

The neural activity in slightly injured nerves is altered in different ways and the results on its ability to activate its target cells can vary widely and have different consequences (Fig. 10.4). Injuries to the auditory nerve may make some nerve fibers unresponsive and may cause changes (slowing) in the propagated conduction in the nerve fibers; the change is normally not the same for all nerve fibers. Such temporal dispersion (decreased coherence) in the nerve impulses that arrive at the target neuron can have widely different effects in the excitation of the target neuron: it can fail to activate the target neuron: it can activate it in a similar way as before the injury occurred, or the injury can actually cause an increase in the excitation of the target neuron because it causes a prolonged excitatory postsynaptic potential (EPSP). This means that the effect of slight injury to the auditory nerve could have widely different effects on the neurons in the cochlear nucleus, including increased excitation that could be associated with tinnitus.

Altered time pattern of discharges in the auditory nerve from continuous type pattern may occur as a result of damage to the ear or, more likely, from damage to the auditory nerve may induce tinnitus.

Bursting neural activity that often results from injuries can activate target cells that were not activated by steady firing, even if the mean discharge rate is not altered after changing to burst mode of firing (Fig. 10.5). Such bursting activity can open synapses not normally conducted when the incoming neural activity is a continuous stream of impulses. Bursting activity has been linked to tinnitus but not to hearing loss.
There are many ways that pathways of the central nervous system can be involved in tinnitus, hyperacusis, and other symptoms that occur together or independently.

Sound causes both excitation and inhibition in the nervous system and if this balance is upset toward less inhibition, the “amplification” in neural networks may be so high that self-oscillations occur, and thus, may explain some forms of tinnitus. Reduced inhibition in the auditory nervous system is assumed to promote the development of some kinds of tinnitus. The role of altered balance between inhibition and excitation has been studied in the dorsal cochlear nucleus (DCN) in animal models of tinnitus [5, 13].

The changes in the nervous system that cause the abnormal activity perceived as tinnitus may be initiated by abnormal activity from the ear or an injured auditory nerve. However, it often progresses over time when structures of the central nervous system become involved. That the pathologic processes that cause the tinnitus develop further over time is supported by the observation that tinnitus becomes more resistant to treatment, the longer a person has had the tinnitus [27]. This means that the role of the ear in the pathology of tinnitus may diminish over time.

The changes in the function of the central nervous system through activation of neural plasticity may start by abnormal or absent input from the ear, but after some time the abnormal functioning may have reached a stable pathologic state. This may occur in accordance with Hebb’s principle (“neurons that fire together wire together”). This means that the function of the sensory system may have two stable states: one that is the normal state and the other that is the pathologic state. If this hypothesis is correct, it means that there is a similarity with the bilateral state of the spinal cord in some forms of central neuropathic pain [28] (see Chap. 14).

Various forms of injuries to the central nervous system, and in particular changes in function, caused by activation of neural plasticity can be involved in causing tinnitus. This means that tinnitus can be regarded as being caused by harmful neural plasticity, thus a plasticity disease [29]. Change in synaptic efficacy through activation of neural plasticity, changes in the relation between inhibition and excitation, and re-routing of information to another part of the brain than normally activated by sound are the most important causes of many forms of subjective tinnitus. Establishment of ephaptic transmission is another change in function that has been suggested to be involved in some forms of tinnitus.

Activation of neural plasticity is the most common cause of tinnitus (see Chap. 12). The central nervous system consists of many subsystems that are connected in a complex way. When these connections change, which may occur when neural plasticity is activated, a manifold of symptoms from different parts of the brain may result. The most important is perhaps that activation of neural plasticity may switch on the non-classical pathways, which has several implications: one of which is a form of cross-modal interaction and another is establishment of a subcortical route to the amygdala (see Chap. 8). Other possible effects are redirection of signals to other parts of the brain than those that are normally activated by sound. Misophonia and phonophobia (see Chap. 4) are examples of signs of pathologies that may result from such redirection of information.

Changes in the function of the central nervous system through activation of neural plasticity can also cause diseases that have similarities with some forms of tinnitus, such as central neuropathic pain (see Chap. 14).

**The Central Nervous System**

There are many ways that pathways of the central nervous system can be involved in tinnitus, hyperacusis, and other symptoms that occur together or independently.

Sound causes both excitation and inhibition in the nervous system and if this balance is upset toward less inhibition, the “amplification” in neural networks may be so high that self-oscillations occur, and thus, may explain some forms of tinnitus. Reduced inhibition in the auditory nervous system is assumed to promote the development of some kinds of tinnitus. The role of altered balance between inhibition and excitation has been studied in the dorsal cochlear nucleus (DCN) in animal models of tinnitus [5, 13].

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The changes in the function of the central nervous system through activation of neural plasticity may start by abnormal or absent input from the ear, but after some time the abnormal functioning may have reached a stable pathologic state. This may occur in accordance with Hebb’s principle (“neurons that fire together wire together”). This means that the function of the sensory system may have two stable states: one that is the normal state and the other that is the pathologic state. If this hypothesis is correct, it means that there is a similarity with the bilateral state of the spinal cord in some forms of central neuropathic pain [28] (see Chap. 14).

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Changes in the function of the central nervous system through activation of neural plasticity can also cause diseases that have similarities with some forms of tinnitus, such as central neuropathic pain (see Chap. 14).
Neural Plasticity

Neural plasticity is the ability of the nervous system to change its function on the basis of experience [26]. Neural plasticity is a normal property of the nervous system that only becomes apparent when turned on.

Activation of neural plasticity can be purposeful and beneficial, or it can be purposeful but not beneficial. Activation of neural plasticity can make it possible to adapt to changing demand, after damage to a region of the brain or spinal cord; activation of neural plasticity can re-route information to regions that are undamaged; or it can be harmful, creating symptoms and signs of disease. We have called the diseases that are caused by activation of such harmful plasticity, *plasticity diseases* [29].

Neural plasticity has similarities with learning as well as differences. The neural mechanisms are similar, involving change in synaptic efficacy causing long-term depression (LTD) and long-term potentiation (LTP) [30], but learning is different from neural plasticity in many ways. What is learned must be recalled, while the result of plastic changes is always available. The difference can be illustrated by learning to pronounce uncommon word and names, which is a matter of neural plasticity, while learning what to say must be actively recalled, thus a matter of memorizing and learning. Activation of neural plasticity is involved in the normal childhood development and aging processes.

Changes in the function of the nervous system that occur because of activation of neural plasticity are mainly changes in synaptic efficacy [31], it can also include other changes in the function of nerve cells such as change in protein synthesis [12]. These changes occur with little delay and cannot be detected using common clinical tests. Changes in synaptic efficacy may start a sequence of events that include formation of new synapses and elimination of other synapses. Later, change in function may cause morphological changes such as creation of new connections according to the Hebb’s principle [32]. The plastic changes may include sprouting of axons, programmed cell death, etc. There are few physiological tests that can reveal such morphological changes and cannot be detected by available clinical diagnostic methods.

The immediate effect that is caused by change in synaptic efficacy is reversible, and the function may recover when the factors that initiated the plasticity are no longer present. The effects that follow, and which involves morphological changes, are more difficult to reverse (see Chaps. 10 and 15). This has an immediate clinical effect in that it causes plasticity diseases such as tinnitus and central neuropathic pain to become more difficult to successfully treat after being present a long time.

There are two fundamentally different effects of activation of neural plasticity: one is beneficial and one is harmful. The beneficial effect is the best known; it makes it possible to adapt to changing demands, and change functions to function areas of the brain after injuries such as from strokes. The other effect of activation of neural plasticity is harmful, causing symptoms and signs of diseases such as some forms of tinnitus and neuropathic pain. We have called such diseases “plasticity diseases.” Some forms of tinnitus belong to a group of diseases where activation of neural plasticity plays an important role for creating symptoms. Many forms of tinnitus are thus plasticity diseases.

The pathology of plasticity diseases is complex, and several factors are often involved in causing a plasticity disorder [26]. The effect of several factors may add up, and in some disorders two or more factors must be present at the same time in order to cause the symptoms of the disease in question. Hemifacial spasm (HFS) is such an example of more than one factor being necessary for causing the symptoms of the disorder [33, 34]. Irritation of the facial nerve root seems to be the cause because moving the blood vessel off the facial nerve root is a very effective treatment. The fact that blood vessels in close contact with the nerve root is common while HFS is a very rare disorder shows that another (unknown) factor must also be present to cause the symptoms of HFS.

In such a situation, it may be sufficient to treat both the factors if both are necessary to cause symptoms. An example of that is HFS that is caused by hyperactivity in the facial motoneuron, but a blood vessel in close contact with the facial nerve root is one of the two (or more) factors that are necessary for causing the spasm that is the characteristic symptom of HFS. It is therefore sufficient to treat one of the causes, and naturally the selection in moving the blood vessel off the nerve root is typically chosen. Some forms of tinnitus may also have several factors that all need to be present in order for symptoms to manifest but where it would be sufficient to treat just one of these factors.

Other studies have shown evidence that the symptoms (spasm in one side of the face) are not caused by pathologies of the facial nerve but instead hyperactivity of the facial motoneuron, probably caused by activation of neural plasticity [34]. This means that HFS is also a plasticity disorder with similarities such as tinnitus and central pain.
There is now considerable evidence that activation of harmful neural plasticity is the cause of many forms of tinnitus, and perhaps also other symptoms that often occur together with tinnitus, such as hyperacusis and perhaps phonophobia and misophonia. Tinnitus is therefore one of several forms of “plasticity disorders.” It may be appropriate to speculate how a common property of the nervous system, such as neural plasticity, can be harmful to an individual person.

**The Role of Ephaptic Transmission in Tinnitus**

One of the first hypotheses regarding hyperactive disorders such as face pain (trigeminal neuralgia) and HFS [35] was ephaptic transmission between the axons of the trigeminal nerve, the myelin of which had been damaged. The theory about ephaptic transmission between injured (denuded) nerve fibers has been further developed and applied to peripheral nerves and dorsal spinal roots [36, 37]. A similar hypothesis was proposed for the auditory nerve to explain the cause of tinnitus [8].

Such direct communication between denuded axons (ephaptic transmission) was hypothesized to promote temporal coherent firing of many nerve fibers. Ephaptic transmission [8, 36, 37] may occur after injuries, such as in connections with surgical trauma and perhaps in conjunction with vestibular schwannoma [38] – conditions that are known to often be associated with tinnitus. However, the question about ephaptic transmission between nerve fibers in cranial nerve roots is controversial, and the hypothesis has never been proven. For another disorder that can be cured by moving a blood vessel off a cranial nerve root, HFS, it was shown to be unlikely to occur for any long period [39].

From having been the established theory in disorders such as trigeminal neuralgia, HFS, and some vestibular disorders (disabling positional vertigo), it was shown in studies of patients undergoing microvascular decompression operations for HFS that the signs of this disorder are related to hyperactivity of the facial motonucleus rather than ephaptic transmission in the nerve root [39]. However, the fact that the HFS can be effectively treated by moving a blood vessel off the facial nerve root implicates the nerve root with the disease. Likely, the explanation is that development and maintaining the symptoms of these diseases involve several steps where the vascular irritation of a nerve root is just one.

Another form of ephaptic transmission, direct and synchronous activation of many nerve cells, has been hypothesized to be involved in causing epileptic seizures by synchronizing the firing of many neurons in a neuron pool, without being caused by transmission by chemical synapses [9].

Not only that, such “non-synaptic” mechanisms may occur not only from electrotonic coupling through gap junctions but also that the effect of the electrical field outside cell bodies can activate neighboring cells, thus causing ephaptic transmission and thereby causing many nerve cells to fire in synchrony. Ionic interactions (e.g., increases in the extracellular concentration of K+) may have similar effects. This form of ephaptic transmission is believed to be involved in causing some forms of epileptic seizures [9]. It has been shown that some forms of tinnitus are caused by localized epileptike activity [4].

Higher packing density favors that kind of abnormal transmission between nerve cells and may be a contributing cause of symptoms in, for example, autistic individuals where it is known that the packing density of cells in the brain is greater than normal [40]. Similar phenomena may be involved in forms of tinnitus that often occur in conjunction with traumatic brain damage, such as closed head injuries (see Chap. 67).

**The Role of the Non-Classical Auditory Pathways**

There are many ways the central nervous system can be involved in causing and maintaining tinnitus. The most common cause is the change in function that occurs through activation of neural plasticity, which can change the excitability of synapses (synaptic efficacy), activate dormant synapses, and make synapses dormant which can re-route information (see Chap. 12).

One example is re-routing information to the non-classical auditory pathways. The non-classical auditory pathways are normally active in children, as indicated by the presence of a cross-modal interaction, where perception of sound is affected by electrical stimulation of the somatosensory system (the median nerve at the wrist) [41]. Such cross-modal interaction
Pathology of the Auditory System that Can Cause Tinnitus

does not normally occur in adults but has been shown to occur in some individuals with tinnitus [42–44], indicating that the non-classical auditory pathway is involved in causing tinnitus in some individuals with tinnitus [42, 43]. The non-classical pathways (see Chap. 8) [45] have input from other sensory systems and provide a direct subcortical route to the amygdala, which may explain why some individuals with tinnitus also have affective symptoms such as depression and phonophobia.

It has been shown that female reproductive hormones influence neural transmission and enhance among other things GABA receptors [46]. The difference in the incidence of tinnitus in males and females may be related to this effect of female reproductive hormones on GABAergic inhibition.

**Involvement of Non-Auditory Parts of the Central Nervous System**

There is evidence from animal experiments that parts of the brain other than auditory pathways may be functioning abnormally when tinnitus is induced through activation of neural plasticity by, for example, stimulation with loud sounds [13, 47]. For instance, it has been shown that place cells in the hippocampus function abnormally in animals exposed to intense noise, which would have caused tinnitus in humans [48].

Since many systems are connected to each other, pathology in one part may result in pathologic activity in many systems of the brain with wide and unexpected consequences.

**Causes of Hearing Loss and Tinnitus**

There are many ways the nervous system can be involved in causing hearing loss and tinnitus from factors such as noise exposure and administration of ototoxic substances. The cause of the tinnitus could be deprivation of signals that the ear sends to the auditory nervous system. The fact that tinnitus only occurs in some individuals with hair cell injuries indicates that factors other than injuries to hair cells are necessary for the development of tinnitus, emphasizing the complexity of tinnitus. More than one factor must often be present for tinnitus to manifest.

**Effect of Exposure to Loud Noise and Ototoxic Substances**

Noise exposure and administration of ototoxic drugs are common causes of hearing loss. It has been assumed that this is caused by damage to hair cells. However, it has become evident that matters are more complex and the auditory nervous system is also involved (see Chap. 37). Tinnitus often accompanies hearing loss caused by noise exposure or administration of ototoxic antibiotics and other ototoxic drugs (see Chap. 42) [49]. These agents have mostly been assumed to cause damage to hair cells in the cochlea. However, these assumptions may also have to be revised as more information about the involvement of the nervous system is gained.

Noise exposure or administration of ototoxic drugs causes hearing loss. Noise exposure and administration of ototoxic drugs have been assumed to primarily have these adverse effects by damaging hair cells (especially outer hair cells) in the cochlea, thus impairing the cochlear amplifier (see Chap. 37) [45]. However, animal studies have provided evidence that the effect is more complex [50, 51].

Animal experiments have shown that exposure to high-intensity sounds causes changes to occur in the response from structures of the ascending auditory pathways such as the cochlear nucleus [52], as well as in non-auditory structures such as the hippocampus [48]. Exposure to loud sounds may cause tinnitus because of the resulting hearing loss and deprivation of input to the auditory system, activating neural plasticity.

**Deprivation of Input to the Auditory Nervous System**

There is evidence that severance of the auditory nerve can in fact cause tinnitus or make existing tinnitus worse [18]. Lack of input to the auditory nervous system as a cause of tinnitus is supported by the finding that tinnitus can be ameliorated by cochlear implants [53, 54] (see Chap. 77) or by applying high-frequency (4,800 Hz) impulses to the round window of the cochlea [55].

Hearing loss causes deprivation of input to the auditory nervous system and can activate neural plasticity.

Another reason that deprivation of input from certain parts of the cochlea, such as the high-frequency (basal) part, causes tinnitus may be that such input
normally provides inhibitory influence on neurons in the auditory nervous system.

There is some evidence that the effect of overexposure to sound is not only on the cochlea but also affecting the auditory nervous system, especially the cochlear nucleus where noise exposure has been reported to cause morphological changes [56]. Other investigators have reported physiological changes after exposure to loud noise [48, 52, 57, 58].

Neural plasticity may also be activated by abnormal input, including overexposure to noise (see Chap. 12). The central nervous system may act on the hair cells through the descending pathways olivocochlear bundle. We discussed earlier age-related hearing loss (presbycusis) and showed that it can have influence from the central nervous system [59] in addition to the well-known degeneration of cochlear hair cells (Chap. 36).

There is evidence from other studies that deprivation of input to a structure such as the sensory cerebral cortex may result in the unused part being taken over by other systems. For example, an unused auditory cortex can be taken over by the visual system [60]. It is now well accepted that children who are born with hearing deficits or acquire hearing deficits early in life must have input to their auditory system re-established in order to ensure a normal development of the auditory system. Little is known about the possibility of early deficits in hearing causing tinnitus.

**Signs of Tinnitus**

There are few external signs of tinnitus except the individual’s testimony. Cross-modal interaction occurs in some individuals with tinnitus (discussed in Chap. 9). Some can modulate their tinnitus by muscle contractions or change of gaze and some individuals cause tinnitus by stimulation of the skin [43, 44, 61]. There are possibly some signs of electroencephalography (EEG) or magnetoencephalography (MEG) changes that are specific to tinnitus [62].

**Interaction between the Auditory and Somatosensory System**

Animal experiments have shown that axons from one sense can grow into the brain territory of another sense if activation of that sense is suppressed or not existing such as in congenitally deaf individuals [60].

There is evidence from several studies that an abnormal cross-modal interaction occurs together with tinnitus in some individuals [42–44, 61, 63]. Several different explanations for such cross-modal interaction have been presented. One explanation involves the non-classical auditory pathways. It was mentioned in Chap. 8 that the non-classical auditory pathways receive input from other sensory systems. This means that activation of the non-classical pathways can make interaction from other sensory systems possible, but signs of function of the non-classical pathways (cross-modal interaction) have been shown only in children [41] and under some pathologic conditions such as tinnitus and autism [64]. While the non-classical pathways seem to be active in some animals used in auditory experimentation, such interaction does not seem to be normal in adult humans, but it is a normal phenomenon in children [41], which may be one of the reasons that children seem to react differently to tinnitus (see Chap. 6). When cross-modal interaction occurs in adults, it may be regarded as pathologic and can be involved in symptoms such as tinnitus or can occur together with tinnitus [42].

Other possibilities for cross-modal interaction involve the anatomical connections between the caudal trigeminal nucleus and the DCN dorsal column nuclei. The connections between the DCN and the dorsal column nuclei and the trigeminal nucleus (as described in Chaps. 8 and 9) provide the anatomical substrate for one form of cross-modal interaction. These connections, shown to be active in animals, may not be active in humans under normal circumstances. The synapses on neurons in the DCN through which they receive somatosensory input may normally be dormant, but reduced auditory input to these neurons may activate the synapses that mediate somatosensory input to these cells [63, 65]. This assumption is supported by the finding that the response of DCN neurons to somatosensory input is increased in animals that have noise-induced hearing loss [66]. It is known from other studies that decreased use of synapses cause some of the synapses to become dormant, and some may become eliminated and their place on a cell membrane becomes taken over by other synapses (see [26]).

There are other mechanisms that could contribute to these effects. One example is the activation of GABA\(_A^b\) receptors in the DCN that regulate dendritic excitability and excitatory inputs [67, 68]. GABA\(_A^b\)
receptors in CN could modulate glutamatergic neurotransmission. The GABAergic inputs could come from several cell types, some of which also contain glycine receptors [69, 70]. Similar processes could occur in the ICX that receive direct input from DCN neurons. Effects that are present at the level of the DCN may become enhanced within the IC.

Some studies [71, 72] found that tinnitus often occurs together with TMJ problems and often resolves when the TMJ problems have been successfully treated [73, 74]. A population study involving 989 consecutive patients [75], however, did not find a higher prevalence of tinnitus in patients with TMJ problems (7.28%) than what occurs in the general population (10–14%). These anomalies may be explained by interaction between the auditory system and the somatosensory system.

A different and perhaps an even stronger indication of an abnormal interaction between sensory systems is the observation made by some individuals with tinnitus that stimulation of the skin results in perceiving a sound. A person with tinnitus mentioned that when he dried his back with a towel, he would hear a swishing sound. Some individuals can turn their tinnitus on and off or make it change its pitch or loudness by performing certain motor or sensory manipulations [76]. Also, a change in eye gaze, from a neutral head-referenced position, is one such behavior that can evoke or modulate tinnitus [61]. These phenomena are signs of interactions between the auditory system and the somatosensory system.

Tinnitus that Occurs as a Part of the Symptoms of Other Diseases

Tinnitus, and especially hyperacusis, may occur with other symptoms of some diseases such as Ménière’s disease, Wilson’s disease, and some forms of autism.

Ménière’s Disease

Tinnitus is one of the three (or four) symptoms of Ménière’s disease (see Chaps. 38 and 60). Since the two other symptoms of this disorder (fluctuating hearing loss and vertigo) are closely related to the function of the ear, it has been assumed that tinnitus that occurs as a part of the symptoms of Ménière’s disease is also caused by a malfunction of the structures in the ear. There are indications that the tinnitus in Ménière’s disease is different from other kinds of tinnitus. For example, it can be shown to be affected by sympathectomy [77], not known to be effective in other kinds of tinnitus.

The symptoms — fluctuating hearing loss, tinnitus, and vertigo — of Ménière’s disease (see Chaps. 42 and 60) have been regarded to be caused by local changes in the fluid system of the inner ear. The finding that stimulating vestibular receptors with air puffs applied to the middle ear cavity can ameliorate at least some of the symptoms of Ménière’s disease [78] is an indication that the nervous system is involved in causing the pathology of Ménière’s disease.

Applying air puffs to the inner ear, thus stimulating the receptors in the inner ear, may have an effect by activating neural plasticity. These observations indicate that the pathology of the disease is not limited to the inner ear, but the nervous system may also be involved in creating the symptoms of Ménière’s disease.

This observation was developed into a clinical method using a device named the “Meniett.” The results of clinical tests have shown beneficial effects of such treatment [79, 80].

Migraine

It has been hypothesized that symptoms such as phonophobia, tinnitus, fluctuation in hearing perception, and increased noise sensitivity [81] that often occur during migraine attacks are caused by an effect on cochlear blood vessels as a component of basilar artery migraine. Evidence has been presented that trigeminal neurogenic inflammation is involved in the development of vascular migraine with its components such as tinnitus and phonophobia [81].

Disorders that Affect the Auditory Nerve

Different kinds of injury can cause tinnitus. Thus, vestibular schwannoma, although these tumors often originate from the superior vestibular nerve, cause some destruction of the auditory nerve, and tinnitus
almost always occurs in individuals with vestibular schwannoma (see Chap. 39) [43, 80]. Physical trauma to the auditory nerve, which may occur in head injuries caused by accidents, explosions (see Chap. 67), or surgery, may cause injuries to the auditory nerve by manipulations and heat from electro-coagulations. Temporomandibular joint disorders are often accompanied by tinnitus [74, 75] (see Chap. 96). Cerebrovascular diseases may also be accompanied by tinnitus (see Chap. 41). The common headache is often accompanied by tinnitus (see Chap. 61). Also, viral infections can result in tinnitus. Very little is known about how several different kinds of these injuries to the auditory nerve may cause tinnitus.

**Head Injuries**

It is known that head injury is often associated with tinnitus (see Chap. 67). Head injuries of various forms, such as from blast injuries, also result in many other symptoms such as epileptic seizures [9]. Close contact between many nerve cells, such as from abnormally dense packing of nerve cells, promotes ephaptic transmission. There are reasons to believe that ephaptic transmission between nerve cells may contribute to tinnitus in head injuries, which means nerve cells can be activated by field potentials in adjacent cells.

Some case reports have shown that disorders of the cerebral cortex (temporal lobe) may be associated with tinnitus [82], supporting the hypothesis that tinnitus can originate from the cortex.

Rare diseases such as Williams syndrome [83] and some forms of autism [64] are associated with hyperacusis.

**Tinnitus and Stress**

Stress seems to influence the tinnitus of people who already have the disorder, and stress may cause tinnitus [82, 84]. This could occur in many ways, one is direct effect on the hair cells through secretion of norepinephrine. Another cause of stress could be through an effect on the nervous system. It has been reported that tinnitus sometimes occurs just before a person faints [85]. This has been related to an effect on the inner ear fluid systems from a hemodynamic imbalance [85]. It could also naturally be caused by change in sympathetic activity that accompanies syncope or it could be caused by the effect of low blood pressure on the brain. Syncope may be regarded as a situation where systemic blood pressure has become lower than the range where the auto-regulation can keep blood flow in the brain constant, independent of systemic blood pressure.

The abundant innervation of both cochlear and vestibular sensory cells by the efferent fibers of the olivo-cochlear bundles (see Chap. 8) could affect the function of the hair cells and possibly the balance of the fluid volumes in the inner ear.

Many sympathetic nerve fibers terminate close to hair cells in both the cochlea and the vestibular apparatus [86]. These fibers secrete norepinephrine in response to activation of the sympathetic nervous system. Epinephrine secreted from these fibers can alter the function of the hair cells and increase their sensitivity; that may explain why sympathectomy has a beneficial effect on tinnitus in Ménière’s disease [77]. Sympathetic blockage (stellate ganglion block) has also been shown to reduce sudden hearing loss [87], thus, a further sign of an effect of the sympathetic innervation of the cochlea.

**What is the Neural Code of Tinnitus?**

It is not known which properties of the discharge in single auditory nerve fibers are interpreted as a sound. It was earlier thought that increased firing rates would signal the presence of sound, but the discharge rate of auditory nerve fibers in animals that have been treated in ways that would cause tinnitus in humans (such as by using administration of ototoxic antibiotics) is not elevated but rather reduced [88–91]. The same is the case for acute injury to the cochlea [92]. Other animal experiments have shown that administration of salicylate, in dosages that are known from humans to give tinnitus, can cause an increase in the spontaneous discharge rates of single auditory nerve fibers [93, 94].

Eggermont and coworkers have found that long-time (4 months) stimulation with frequency bands (two octave wide and 80 dB sound pressure level [SPL]) of sounds of moderate intensity decreases the
responsiveness of cells in the primary auditory cerebral cortex that are tuned to the frequencies of the stimulation and increases the responsiveness of cells that are tuned to frequencies of the edge of the spectrum of the stimulus sounds [95]. More recently, it was found that similar effects could be obtained using lower stimulus intensity (68 dB SPL) and shorter time of exposure (6 weeks) [96].

Recent research has shown evidence that synchrony of the firing (temporal coherence) in large groups of nerve cells is more likely to be the neural code that causes the sensation of the presence of sounds, and most likely many forms of tinnitus [49, 97, 98], than discharge rate.

It has been hypothesized that phase-locking discharge in many auditory nerve axons is the neural code that signals the presence of sound [8] and thus, also probably the code that falsely communicates a sound is present when not causing tinnitus. Animal studies have indicated that the coherence of temporal patterns of activity in individual neurons in a pool of neurons is important for providing the normal awareness of the presence of a sound, including tinnitus. This means that correlation of neural activity in populations of nerve cells in the auditory nervous system is most likely what causes tinnitus [49, 98] (see Chap. 16).

Thus, there is considerable evidence that tinnitus is not directly related to the discharge rate of auditory nerve cells, which seems surprising because tinnitus is regarded as a hyperactive disorder. These observations, however, are in agreement with other studies that indicate that the discharge rate of auditory nerve fibers does not communicate information about the strength (intensity) of sounds [99].

Such phase locking occurs normally in response to sounds because the same source drives the nerve activity. It can also occur pathologically through an abnormal coupling between nerve fibers or nerve cells, known as ephaptic transmission.

**Which Structures Function Abnormally in Individuals with Tinnitus?**

Which parts of the auditory pathways have abnormal function in individuals with tinnitus is not known. Many studies have focused on the primary auditory cortex but there is also evidence that subcortical connections from the dorsal and medial thalamus to the lateral nucleus of the amygdala may convey sensation of tinnitus. The sensory cerebral cortices are just another processing station. The anatomical location where perception occurs is much higher, but it is unknown exactly in which structures neural activity causes sensory perception.

The amygdala nuclei have connections to most parts of the brain, which means that this route could be important in some forms of tinnitus. The medial and dorsal parts of the auditory thalamic nuclei are parts of the non-classical auditory pathways. Signs of activation of these pathways have been shown as a constant phenomenon in children below the age of 12 years [41]. Adults thus do not normally have such signs, but some individuals with tinnitus have similar signs of involvement of the non-classical auditory pathways as young children [42]. Other studies have shown an increased activation of the amygdala in some individuals with tinnitus [100].

**Detectable Changes in Function of the Auditory Nervous System in Tinnitus**

There are a few objective signs that can provide information about the magnitude of tinnitus and its character, and indeed there are no available objective tests that can determine if a person has tinnitus at all. The character and the magnitude of the annoyance and distress caused by tinnitus can only be assessed by interviewing the person. While there are no clinically recognized methods for objectively assessing the severity of tinnitus or helping distinguish between different forms of tinnitus, some laboratory methods have shown promise for being of clinical value. It has, however, recently been shown that some forms of tinnitus are associated with specific abnormalities in EEG and MEG recordings. The future will tell whether these techniques will be applied in clinics for diagnosing tinnitus.

Another promising method is to use transcranial magnetic stimulation (TMS) as a test of the ability for electrical stimulation of specific parts of the cerebral cortex to beneficially affect an individual’s tinnitus [101–103].
Change in Tonotopic Maps

Studies have shown indications of abnormalities in the tonotopic organization of the cerebral auditory cortex in patients with tinnitus using magnetoencephalographic (MEG) recordings [104]. The functional importance of the tonotopic organization is, however, unknown; it is not known if the observed changes are primary to the pathology of tinnitus or caused by the effect of abnormal neural input to the auditory cerebral cortex.

Much attention has been paid to these observed changes in the tonotopic organization in individuals with tinnitus [104, 105]. The functional importance of normal tonotopic organization is unknown. Therefore, it is not known what the implications of altering the tonotopic organization might be.

Tonotopic map changes and change in tuning acuity of neurons in the auditory system can be explained by change in synaptic efficacy that may occur because neural plasticity has been turned on. Many of the synapses that connect these inputs to a cell are normally dormant. By unmasking synapses and masking other synapses, it is possible to change the tuning of cells, shift their center frequency, and change the width of the tuning. The width of the tuning is determined by how many inputs are active, which thus determines the shape of the tuning curve of the cell in question. This means that masking and unmasking of synapses can change the tonotopic organization of auditory nuclei and in the auditory parts of the cerebral cortex.

Pathology of a specific structure can make many other structures behave abnormally without having a pathologic function simply because they receive abnormal input [29]. For example, the observed change in tonotopic organization in the cerebral cortex may not be caused by pathologies of the cerebral cortex, but the reorganization and abnormal response of cortical structures can equally be caused by pathologic neural activity generated by more peripheral structures and delivered to a normally functioning cerebral cortex.

Conclusion

Of the different pathologies that affect the auditory system, tinnitus is the most complex disorder and it affects many people. It is also the one least known about and has the least effective treatments of common disorders of the ear. It has been falsely regarded as an ear disease for many years, and not until relatively recently has it become generally accepted that the anatomical location of most forms of tinnitus is the central nervous system. Most forms of tinnitus are phantom sensations caused by activation of harmful neural plasticity; it is thus a plasticity disorder that has many similarities to central neuropathic pain. Its management requires a multidisciplinary approach.

References


Chapter 11
The Role of Auditory Deprivation

Aage R. Møller

Keypoints

1. Deprivation of input to the auditory system can cause two kinds of change in function: It can alter the balance between inhibition and excitation and can activate neural plasticity.
2. Hearing loss of any kind, such as conductive hearing loss or cochlear hearing loss, causes decreased input to the auditory nervous system.
3. Noise-induced hearing loss is an example of deprivation of auditory stimulation and overexposure, which in itself may activate neural plasticity.
4. Altered balance between inhibition and excitation can change the gain in the auditory system. If the gain is increased, it may cause hyperactivity in the form of tinnitus.
5. The effect on the balance between inhibition and excitation is likely to abate when normal input to the auditory system is established.
6. Activation of neural plasticity, which may occur because of sensory stimulation, may last after restoring normal sensory stimulation.
7. Plastic changes may become permanent, and reversal of neural plasticity may require special actions.

Keywords Tinnitus • Sound deprivation • Neural plasticity • Inhibition • Temporal coherence

Introduction

The effect on the nervous system of sensory deprivation can be profound, and is different when it occurs at birth or shortly thereafter, compared with occurring during adult life. The anatomical and functional development of the nervous system depends on sensory stimulation. Therefore, sound deprivation can have a stronger effect on young individuals than on adults. The fact that there are indications that the nonclassical pathways are normally active in children [1, 2] while not normally active in adults, may influence the way children react to deprivation of sound compared with adults.

Deprivation of input to the auditory system can mainly cause two different kinds of change in the function of the auditory nervous system, both of which can cause tinnitus: (1) It can decrease or shift the balance between excitation and inhibition and thereby increase the gain in the auditory nervous system and (2) deprivation of sensory stimulation can activate neural plasticity involving change in synaptic efficacy and sprouting of axons [3]. The effect of sound deprivation may not be easily observed because children do not complain of tinnitus in the same way as adults (see Chap. 6).

There are many ways that the auditory nervous system can be deprived of normal stimulation. Any form of hearing loss can cause some degree of sensory deprivation, whether it occurs through obstruction of the ear canal, disorders of the middle ear (see Chap. 34), or from disorders of the cochlea (see Chap. 35), it may have the same effect on the nervous system.

Tinnitus is common after noise-induced hearing loss (see Chap. 37). The reduced hearing may activate...
neural plasticity, causing the form of tinnitus that occurs after exposure to loud sounds; however, overexposure in itself may also activate neural plasticity and thereby cause tinnitus. The tinnitus that occurs after exposure to noise, which causes hearing loss, or after a brief period of deprivation of sound often disappears after some time. In some instances, however, exposure to loud noise, especially impulsive or high-frequency sounds such as fire alarms, can cause permanent tinnitus.

It was earlier believed that increased neural firing was the cause of tinnitus, but more recent studies seem to indicate that temporal and spatial coherence of activity is more important for eliciting a sensation of the presence of sound including tinnitus [4, 5]. Noreña and Eggermont [6] showed a slight increase in spontaneous firing in cells in the auditory cortex after acoustic trauma.

Many different parts of the nervous system have been implicated in tinnitus. Some investigators have found evidence of altered spontaneous activity that is different at different levels of the auditory system [7]. There is evidence that tinnitus may be associated with less neural excitation in the periphery of the ascending auditory pathway but greater activity in more central structures. Some have hypothesized that increased synchrony of neural firing can cause tinnitus. Other investigators have hypothesized that temporal coherence of firing in large groups of nerve cells is the cause of some forms of tinnitus [4, 8].

The frequency tuning in the cochlea is the basis for the tuning of nerve cells throughout the auditory nervous system. The acuity of tuning to sounds depends on the intensity of the sound; the higher the intensity, the broader the tuning [9, 10]. Increased temporal coherence of firing in many nerve cells may be caused by the broadening of the cochlea’s tuning that occurs at higher sound intensities, thus causing a greater degree of overlap of different cells’ response areas in the cortex. Unmasking of dormant synapses of interneurons, which often occur as a result of activation of neural plasticity, may also cause increased coherence of neural firing [3]. Changes in the relation between excitation and inhibition may likewise cause increased coherence and increased spontaneous firing. Both such changes may therefore be caused by reduced sensory stimulation.

Change in Balance Between Inhibition and Excitation

Single auditory nerve fibers have both excitatory and inhibitory response areas that mainly surround the excitatory areas [11]. The inhibition that is present in the response of single auditory nerve fibers is not caused by synaptic inhibition, but it is instead a form of suppression that is a result of cochlear nonlinearities [8]. Similar arrangement of suppression and excitation is present throughout the auditory nervous system, where the suppression is caused by synaptic inhibition. This means that a sound such as a tone will activate both inhibition and excitation (see Chap. 15). This suppression or inhibition is similar to what is in the visual system known as lateral inhibition. If pathologies of the cochlea result in a greater reduction of inhibition than excitation in a population of neurons, they may become sufficiently active to produce awareness of sound without sound reaching the ear, thus tinnitus. Tinnitus can be suppressed by proper arrangement of sound stimulation. Thus, sound in certain frequency regions can suppress some forms of tinnitus, and that may occur because such sounds contribute more to inhibition than excitation of specific populations of nerve cells. There are some indications that high-frequency sounds elicit stronger inhibitory influence on neurons in the cochlear nucleus more than low frequencies. This means that high-frequency hearing loss, which is common, may cause tinnitus because it reduces normally occurring inhibition. This can also explain why high-frequency stimulation can be effective in reducing some forms of tinnitus.

The interaction between inhibition and excitation is present along the ascending pathways including the cerebral cortices. Lateral inhibition is especially prevalent in the inferior colliculus where interaction between excitation and inhibition is especially prevalent. It has been shown that selective damage to sensory cells (acoustic trauma) in the cochlea that reduces the evoked potentials recorded from the auditory nerve, in fact, increases the discharge rate of many neurons in the inferior colliculus [12], indicating that the deprivation stimulation caused by cochlear trauma has decreased inhibition in these third-order neurons of the ascending auditory pathways. The observed changes suggest that these cells receive inhibitory input from
high-frequency regions of their response areas and this inhibition has been reduced by deprivation of stimulation caused by cochlear trauma.

Josef Syka and his collaborators have shown that acoustic trauma causes increased activity in central auditory structures [13, 14]. This means many investigators agree that neural activity in the auditory periphery is decreased by acoustic trauma while it is increased at central levels such as the inferior colliculus and the cerebral cortex. Reduced inhibitory activity could explain these changes that go in opposite directions in the periphery and the central auditory structures. Also, the edge effect [15] may be a consequence of lateral inhibition. The fact that tinnitus is more prevalent in elderly individuals may be explained by the reduction in inhibition that normally occurs with age [16], thereby shifting the balance between excitation and inhibition toward excitation.

The hypothesis that deprivation of high-frequency sounds is involved in many forms of tinnitus is supported by animal (chinchilla) studies that show that auditory nerve fibers tuned to high frequencies tend to have elevated spontaneous activity [17]. This indicates that normal suppression was reduced after the noise exposure assumed to have caused tinnitus. Mathematical modeling predicts that the response of cells in the dorsal cochlear nucleus after noise trauma depends on the cell type, one type of cells become hyperactive, whereas another type is not affected by acoustic trauma [18].

Studies of temporal integration in the inferior colliculus of rats have shown decreased signs of GABAergic inhibitory activity in the cochlear nucleus after acoustic trauma [19]; administration of GABA_A receptor agonists (benzodiazepines) reversed these changes [20].

Experience from treatment of people with tinnitus has also supported the hypothesis that deprivation of auditory stimulation decreases inhibition in the auditory nervous system.

Watanabe et al. [21] in a study of 600 individuals with tinnitus found that therapy with narrow band noise could suppress tinnitus in 66% of individuals, more so in individuals with presbycusis than sudden deafness. Souliere et al. [22] studied the effect of cochlear implants on loudness, annoyance, daily duration, location, and residual inhibition of tinnitus in 33 postlingual deafened individuals. Eighty-five percent of these individuals had tinnitus. The study showed a significant reduction in both loudness and annoyance. Fifty-four percent of the individuals who had tinnitus before implantation had a loudness decrease of 30% or more; 43% had a decrease in annoyance of 30% or more. The duration of the tinnitus decreased 30% or more in 48% of the individuals who had tinnitus before implantation. The fact that many of the participants in these studies experienced contralateral residual inhibition and tinnitus suppression suggests that a central mechanism contributed to their tinnitus.

Using a computational model of a lateral inhibition neural network, Kral and Majernik [23] showed evidence that lateral inhibition might be involved in some forms of tinnitus. These investigators suggested that the spontaneous activity in the auditory nerve, when subjected to lateral inhibition, can cause phantom perceptions in the absence of auditory stimulation that many individuals experience when placed in silence, such as in an acoustically shielded chamber used for audiologic testing. Kral and Majernik [23] suggested that neural noise normally generated in neural networks is generally masked by a sound stimuli or ambient broadband acoustic noise. Inhibition may balance excitation in response to broadband noise, but the spectrum of other kinds of noise determines what extent the response will be suppressed by inhibition or whether excitation dominates.

Rubinstein et al. [24] have shown that stimulation with electrical impulses at a high rate applied to the cochlea can reduce tinnitus in some individuals. The fact that especially high-frequency electrical stimulation of the cochlea has a beneficial effect on tinnitus in both deaf individuals [22] as well as individuals who do not have much hearing loss (see Chap. 77) [22] supports these hypotheses. Other investigators [25, 26, 27] found that electrical stimulation of the cochlea can reduce some forms of tinnitus by counteracting the effect of reduced activation of the auditory nervous system.

Activation of Neural Plasticity

Considerable evidence has been presented that activation of neural plasticity is involved in many forms of tinnitus (see Chaps. 12, 13, and 14) [28]. Many forms of tinnitus are therefore “plasticity disorders” [29].
The strongest promoter of neural plasticity is deprivation of sensory stimulation [3].

The effect of activation of neural plasticity can be changes in the function of the nervous system that occur with a short delay and last for just a short period for a long time.

It is assumed that acoustic trauma causes deprivation of input to the auditory nervous system because of the hearing loss it causes. There is, however, also the possibility that overstimulation may activate neural plasticity, which in turn can cause changes in the function of the nervous system that may result in tinnitus.

Studies in animals have shown evidence that auditory deprivation can cause cortical map modifications, and such cortical plasticity is associated with decreased inhibition [30].

There are several ways that deprivation of sensory stimulation can immediately affect functions of the auditory nervous system. These matters are discussed in Chap. 12.

References

Chapter 12
The Role of Neural Plasticity in Tinnitus

Aage R. Möller

Keypoints

1. There is evidence from many studies that plastic changes in the central nervous system are involved in causing many forms of tinnitus.
2. Expression of neural plasticity may cause symptoms of sensory system disorders by changing neural processing and rerouting of information.
3. Rerouting of information through activation of neural plasticity may explain the occurrence of affective symptoms (mood disorders), phantom sensations, improved perceptual capabilities, or atypical sensory experiences such as phantom sensations, tinnitus, and neuropathic pain.
4. Changes in the processing of information may cause hyperacusis and distortion of sounds in connection with some forms of tinnitus.
5. There is evidence that the nonclassical auditory pathways in adults may be activated through expression of neural plasticity, causing cross-modal interaction in some individuals with tinnitus.

Keywords  Tinnitus • Neural plasticity • Hyperacusis • Plasticity disorders • Nonclassical pathways • Extralemniscal pathways

Abbreviations

CNS  Central nervous system
MVD  Microvascular decompression

Introduction

Plastic changes in the brain are involved in many forms of tinnitus and also in abnormal perception of sounds such as hyperacusis, as discussed in Chap. 10. Neural plasticity may also be involved in the affective disorders that often accompany tinnitus, such as phonophobia and depression.

Many forms of tinnitus are phantom sensations; the sensation is caused entirely by activity in the central nervous system that is maintained without any signals from the body, including the ear. Therefore, tinnitus has similarities with the phantom limb syndrome and central neuropathic pain. These symptoms belong to a group of adverse and harmful effects that can occur when neural plasticity is turned on and they have been termed “plasticity disorders” [1]. Deprivation of input to the nervous system is the strongest factor that can activate neural plasticity. Activation of neural plasticity is involved in many forms of tinnitus, and the topic has been reviewed recently by many authors in journal articles [2–5] and books [6, 7]. Since deprivation of sensory signals in general is a strong promoter of plastic changes, deprivation of sound is also an important factor in tinnitus as it can worsen existing tinnitus and cause the disorder in individuals who do not experience it in normal-sound environments.

Effects of Activation of Neural Plasticity

There are thus two fundamentally different effects of activation of neural plasticity; one being beneficial and the other being harmful, causing symptoms and signs of disease. The harmful effects are thus called “plasticity disorders” [1].

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A.R. Möller
The University of Texas at Dallas, School of Behavioral and Brain Sciences, GR 41, 800 W Campbell Road, Richardson, TX 75080, USA
e-mail: amoller@utdallas.edu
Plastic changes can alter the processing of sounds, cause hyperactivity that may cause tinnitus, and re-route signals in the CNS, which may cause hyperacusis and affective symptoms and promote coherent firings of many neurons in a pool of neurons. It has been hypothesized that such increased coherence may be the cause of tinnitus.

Neural plasticity is a property of the nervous system that can change its function in various ways. The changes in function occur only when neural plasticity is activated. Neural plasticity may be activated, or turned on, by sensory experience such as reduced sensory signals (deprivation of input), by overstimulation, and by injuries of various kinds [7]. Intrinsic factors may turn on neural plasticity without any known cause.

The immediate effect of activation of neural plasticity would explain why a person who is placed in a silent room experiences tinnitus immediately. The effect that occurs later would explain why a certain treatment is more successful early in a disease.

**Rerouting of Information**

Activation of neural plasticity can open routes that are normally blocked because of ineffective synapses [8] by unmasking such dormant synapses [7, 9] (see Chap. 10). This property may be especially important for tinnitus. There are indications that the nonclassical pathways are becoming activated in that way in some individuals with tinnitus [10]. Activation of the nonclassical pathways may result in a different processing of sounds and can explain cross-modal interaction because the nonclassical auditory pathways receive input not only from ear but also from sensory receptors of other sensory systems, such as the somatosensory system. The nonclassical pathways also involve subcortical routes to the amygdala and other limbic structures by activating what has been called the “low route” [11] from the dorsal and medial thalamus [12]. Limbic structures have been shown to be abnormally activated in some individuals with tinnitus [13] using imaging techniques.

Several studies have shown evidence that tinnitus may be associated with affective syndromes such as depression and fear (phonophobia) which may be explained by an abnormal establishment of a subcortical route from the dorsal and medial thalamus to the lateral nucleus of the amygdala.

**What Can Activate Neural Plasticity?**

Many factors can activate neural plasticity including sensory experience. In connection with tinnitus, reduced sensory input from the ear to the auditory nervous system is an important factor for developing tinnitus. However, overstimulation can also activate neural plasticity causing tinnitus. Other factors that can activate neural plasticity are intrinsic factors such as inflammation; unknown factors may also be involved in turning on neural plasticity and causing or contributing to tinnitus.

**Deprivation of Signals to the Nervous System**

Most individuals who have tinnitus suffer from hearing loss, but a few have very little hearing loss or actually possess normal hearing.

There is considerable evidence that tinnitus may occur after damage to the ear or the auditory nerve that may cause reduced input to the central nervous system. This can have two different effects: It can change the balance between inhibition and excitation and it can promote activation of neural plasticity. Damage to the ear can reduce input to the central auditory system and shift the balance between inhibition and excitation. Deprivation, as such, is unlikely to be the cause of tinnitus but can start a sequence of events of central modifications of the function of the auditory system and likely other parts of the central nervous system. Decreased input to the nervous system can activate neural plasticity.

Sounds that reach the ear cause both inhibition and excitation; decreased input to the nervous system may also change the relation between inhibition and excitation. Sound deprivation can occur because of pathologies of the ear or the auditory nerve, as well as a lack of environmental sounds.

The effect of deprivation of sound increases with age. In a study of 120 normal hearing young adults, it was found that 64 experienced tinnitus when seated in a silent room used for audiologic tests after a short time period (4 min). None of the participants experienced tinnitus in ordinary environments [14]. Placed in a room of silence is also likely to increase the tinnitus of individuals who have tinnitus in ordinary environments.

 Interruption of input from the ear to the auditory system (deafferentation of the auditory periphery) can
induce neural plasticity, causing changes in the function of the central nervous system from deprivation of input that may cause some forms of tinnitus. Age-related hearing loss (see Chap. 36) is the most common situation where deprivation of sound causes tinnitus and also hearing loss. Cochlear hearing loss from noise exposure (see Chap. 37) is another example of decreased signals from the ear that may cause tinnitus because it activates neural plasticity.

Middle-ear diseases can lead to tinnitus through activating neural plasticity by reducing the sound that reaches the cochlea (see Chap. 34). When appropriately treated or compensated by amplification, the tinnitus is likely to improve or disappear completely.

Age-related hearing that often occurs in elderly individuals mainly affects high-frequency hearing loss. Hearing loss is more likely to cause tinnitus in older individuals than when occurring in younger individuals. This may be because inhibition in the nervous system decreases with age [15], which just adds to the effect of reduced sound that normally causes inhibition.

Altered time pattern of discharges in the auditory nerve from continuous type of pattern to burst pattern that may occur as a result of damage to the ear or more likely from damage to the auditory nerve may induce tinnitus. Another factor that seems more plausible as a cause of tinnitus is synchrony of the discharges in many nerve fibers [16, 17].

The fact that there are many individuals who have considerable hearing loss but no tinnitus means deprivation or reduced input to the auditory system does not always turn on harmful plasticity. This means neural plasticity turned on by deprivation of input to the auditory system is not the only cause of tinnitus.

**Overexposure**

Exposure to loud sounds, especially impulsive noise, can cause immediate tinnitus [18, 19] in addition to hearing loss (see Chap. 37). In most people, the tinnitus decreases with time after the end of the exposure. In some individuals, a single exposure can cause a lifetime of tinnitus, thus creating a best-able situation where the nervous system has two stable conditions; one normal and one that is pathological. A one-time exposure may cause tinnitus because permanent damage has occurred to the ear. Changes in function of the nervous system induced by activation of neural plasticity may have two permanent states: one normal and one pathologic, which can explain why permanent tinnitus may occur after a single exposure. The cause of the tinnitus from noise exposure may be changes in the ear or the nervous system through activation of neural plasticity, or by the deprivation of input to the auditory nervous system that occurs because of the damage caused by the overexposure. There are indicators that overexposure in itself can turn neural plasticity on.

In order to correct this bi-stable property, it is not only necessary to reverse the cause of the pathologic changes but the state of the neural circuitry must also be flipped back to restore normalcy (see Chap. 14). Conditions caused by plastic changes can be permanent because of Hebb’s principle: neurons that fire together will eventually also connect morphologically together (“wire together”) [7].

**Treatment of Plasticity Disorders**

Similarly to what is experienced from neuropathic pain, tinnitus becomes difficult to treat the longer it persists. This was shown in a study of the efficacy of MVD of the auditory nerve for treatment of tinnitus in patients who had had their tinnitus for different lengths of time [20]. There was a marked difference when the participants were divided according to the time they had had their tinnitus. Of the 72 patients – 40 men and 32 women – who underwent MVD for tinnitus, 18.2% had total relief, 22.2% showed marked improvement, 11.1% had slight improvement, and 2.8% (two patients) became worse. Those who experienced total relief or marked improvement had had their tinnitus for a shorter period than those who had had their tinnitus for a longer period, 2.9 and 2.7 years, respectively; those who only achieved slight improvement or no improvement at all had had their tinnitus much longer, 5.2 and 7.9 years, respectively.

The finding that there was a marked difference between the outcome in men and women is more difficult to explain. Of the women, 54.8% had relief or some improvement of their tinnitus, but only 29.3% of the men had such improvements. The participants in this study all had severe tinnitus (it can serve as a reminder of the seriousness of severe tinnitus that two of the participants who did not have any improvement committed suicide within a year after the MVD operation).
Development of Beneficial and Harmful Plasticity

It seems reasonable to assume that beneficial neural plasticity has developed as a result of natural selection (Darwinian), but development of harmful plasticity causing plasticity disorders seems contradictory to natural selection that is normally assumed to favor the development of favorable functions such as beneficial plasticity. Activation of beneficial neural plasticity that causes adaptation to changing demands and redirection of signals from injured parts of the brain to functional parts seems in accordance with common hypotheses about Darwinian development. These functions are likely to have developed according to the principle of natural selection of the fittest [21] and are purposive and beneficial to an individual person. It is less obvious how harmful plasticity has developed, but it could be argued that plasticity disorders are caused by maladaptive plasticity.

If this hypothesis is accepted, one can ask if nature tries to correct its mistakes and if harmful plasticity (plasticity disorders) can be expected to vanish as development goes forth. Mistakes in evolution may disappear, if they cause disadvantages to survival or the accomplishment of reproduction. It is questionable whether tinnitus is a disadvantage in reproduction. It may be assumed that the goals of natural selection are to improve reproduction. It is questionable if plasticity disorders such as tinnitus can affect the ability to reproduce, in particular, since they mainly occur late in life after the end of the reproductive period, which for women is around the age of 40, but much higher for men. Plasticity disorders that occur late in life do not affect the ability to reproduce and thus there is little evolutionary pressure to eliminate such disorders by natural selection.

References

Chapter 13
Neural Synchrony and Neural Plasticity in Tinnitus

Larry E. Roberts

Keypoints

1. Most individuals with chronic tinnitus have high-frequency hearing loss, induced by noise exposure, otological disease, or the aging process. Physiological evidence suggests that in such individuals, tinnitus is likely caused not by irritative processes that persist in the ear after cochlear injury, but by changes that occur in central auditory pathways when the ear is partly disconnected from the brain.

2. In animals, hearing loss induced by experimental noise trauma leads to a reorganization of tonotopic maps in the primary auditory cortex, such that frequencies near the edge of normal hearing come to be overrepresented at the expense of frequencies in the hearing loss region. Neurons show increased spontaneous firing rates in cortical and subcortical auditory structures, and in the auditory cortex, increased synchronous activity in the region of hearing impairment.

3. Evidence from physiological, psychoacoustic, and human brain imaging studies suggests that increased neural synchrony (temporally coupled neural activity) in the hearing loss region may be an important mechanism contributing to tinnitus. Tinnitus spectra and residual inhibition functions overlap the region of auditory threshold shift, consistent with this hypothesis.

4. Several forms of neural plasticity may contribute to changes in spontaneous firing rates and neural synchrony that develop after hearing loss. Because the tuning of auditory neurons can be modified by acoustic training procedures throughout the lifespan, it may be possible to reverse some of the neural changes underlying tinnitus.

5. For this goal to be achieved, it must be possible to modify auditory representations by acoustic training in individuals with tinnitus, and the neural modifications induced by training must intersect with the underlying tinnitus mechanisms. Auditory plasticity in normal hearing individuals and people with tinnitus requires further study.

Keywords  Mechanism of tinnitus • Neural synchrony Cortical reorganization • Neural plasticity • Tinnitus spectrum • Residual inhibition

Abbreviations

HL  Hearing level
CF  Center frequency
RI  Residual inhibition
ASSR  Auditory steady-state response
AM  Amplitude modulation
EEG  Electroencephalogram
MEG  Magnetoencephalography

Introduction

Although our understanding of the mechanisms of tinnitus comes from many sources, two recent lines of research, in particular, have provided insight into the question of how the sensation of tinnitus is generated. The first line of research has shown that hearing loss induced by noise exposure in animal models leads to a reorganization of tonotopic maps in the primary auditory cortex, such that frequencies near the edge of nor-
mal hearing come to be overrepresented at the expense of frequencies in the hearing loss region [1–3]. Because hearing loss is a putative cause of tinnitus, it was suggested that this overrepresentation, or changes in neuron response properties associated with it, may underlie tinnitus percepts [4, 5]. The second line of research demonstrated that neural representations for sound in the primary auditory cortex are not fixed after early development but can be modified over the lifespan by procedures such as deafferentation or auditory training that alter the organism’s experience with sound [6, 7]. This phenomenon is called “neural plasticity” (see Chap. 12). These two lines of research have converged to ask whether neural plasticity may be involved in the generation of tinnitus, and if so, whether acoustic training procedures might be designed to reduce tinnitus or prevent its development when hearing loss occurs.

This chapter reviews evidence from animal models of hearing loss, human psychoacoustic studies, and brain imaging experiments that suggests that tinnitus is generated by abnormal synchronous (temporally coupled) neural activity that develops in the auditory cortex when central auditory structures are deafferented by cochlear pathology. It is useful to formulate a perspective on the neural basis of tinnitus, because treatment procedures designed to reduce tinnitus must interact with this mechanism if tinnitus is to be altered. I also briefly review evidence for neural plasticity in the auditory system and ask whether the rules that describe auditory plasticity in normal hearing individuals apply as well to individuals with tinnitus. This cannot be assumed, because people with tinnitus experience not only some degree of hearing loss but also an auditory sensation that may interfere with the remodeling process. In a later chapter (Chap. 72), we discuss current approaches to sensory training from the perspective of research on these two questions.

**The Neural Synchrony Model of Tinnitus**

It is widely recognized that most individuals who have tinnitus also have sensorineural hearing loss caused by injury, otological disease, noise exposure, or the aging process. Even when auditory thresholds are in the normal range (≤25 dB HL), tinnitus sufferers often have evidence for restricted cochlear dead regions [8] or show threshold elevations in the audiogram on the order of 10 dB in the tinnitus frequency range compared to age-matched controls [9] suggesting that some degree of hearing impairment is present. In most cases, however, it is doubtful that chronic tinnitus is generated by irritative processes that persist in the cochlea damaged by hearing loss. Damage to the cochlea caused by lesioning or noise exposure typically leads not to an increase in spontaneous activity in auditory nerve fibers, which might be expected from such processes, but rather to a decrease in auditory nerve activity, pointing to a reduction of input to central auditory structures [5]. These observations suggest that the sensation of tinnitus in the majority of individuals is generated not in the ear but by changes that have occurred in central auditory pathways when the brain has been partly disconnected from the ear by hearing loss (deprivation of input, see Chap. 11). Consistent with this understanding, most individuals who had tinnitus before removal of a vestibular schwannoma with sectioning of the auditory nerve also had tinnitus after the operation. Tinnitus is also a predictable outcome after sectioning of the auditory nerve in individuals who did not have tinnitus before their operations for vestibular schwannomas or other conditions [10] (see Chap. 39).

Animal models of hearing loss have begun to give a picture of the changes that occur in central auditory pathways following auditory deafferentation. The understanding supported by these studies is summarized in Fig. 13.1a (from [5]), which depicts the primary auditory cortex of a cat that has sustained a high-frequency hearing loss induced by noise trauma. The left side of the figure shows the undamaged region, including thalamocortical afferents synapsing on input neurons followed by feed-forward (i) and lateral (ii) inhibition after one synaptic delay. Feed-forward inhibition is functionally dissociable from lateral inhibition [11] and quenches target neurons after their depolarization, which may protect thalamocortical synapses from down-regulation (and preserve their cochleotopic tuning) when the neurons are driven by uncorrelated input from horizontal fibers in the tonotopic map. Animal studies have shown that when a region of the tonotopic map is disconnected from the ear by cochlear damage (right side of Fig. 13.1a), auditory neurons in the affected region begin to respond preferentially to input conveyed by horizontal fibers as their thalamocortical input is impaired or lost. As a consequence, the cortical tonotopic map “reorganizes” when the affected neurons begin to express the tuning preference of their neighbors, leading to an overrepresentation of edge frequencies in the tonotopic gradient.
Neural Synchrony and Neural Plasticity in Tinnitus

It has been proposed that this overrepresentation of edge frequencies may correspond to the tinnitus percept, which was thought to be confined to the edge of normal hearing. However, this is doubtful not only because of evidence to be presented below but also because it is not obvious how the activity of the affected neurons would be heard in terms other than their original cochleotopic tuning.

Other changes in the response properties of auditory neurons documented by animal studies of hearing loss are more likely to contribute to the tinnitus percept. One such change is that neurons in cortical and subcortical auditory structures (but not auditory nerve fibers) increase their spontaneous firing rates as input from the ear is diminished. This effect could reflect an adaptive rescaling of neuron input/output functions by homeostatic plasticity [12] when afferent input to central auditory structures is impaired, or inhibitory deficits consequent on deafferentation, or most probably both factors. At the level of the cortex, increased spontaneous firing has been observed to occur across the tonotopic map, including tonotopic regions that are affected by hearing loss (typically high-frequency regions) as well as regions that are less affected (typically low-frequency regions). Increased spontaneous neural activity is likely to be an important factor in the development of tinnitus, although it has been suggested by several investigators that uncorrelated neural activity may not be sufficient to generate a coherent sound percept. A second change that may occur is an increase in the temporally synchronous activity of a population of
neurons, which is expressed as an increase in cross-correlated neural firing when compared to control animals [13]. This change is more closely confined to the hearing loss region and appears to reflect synchronous network activity that is forged over lateral connections by neuroplastic mechanisms operating in this region [14], possibly because the quenching effect of feed-forward inhibition is lost. It should be noted that although thalamocortical input to the affected tonotopic region is affected by cochlear injury, the output of the synchronously active neurons remains intact. The neural synchrony model of tinnitus suggests that this output (which is conveyed to the thalamus by nerve fibers more numerous than the forward path) is processed by other brain regions and generates the tinnitus percept (see Chap. 12).

This picture of the neural mechanism of tinnitus has implications for the psychoacoustic properties of tinnitus. One implication is that when participants in a study are asked to rate sounds of different frequencies for similarly to their tinnitus, ratings should not be restricted to the region of the audiometric edge (although contrast enhancement at the edge may contribute [15]), but should instead span the region of hearing loss, increasing in proportion to the depth of hearing impairment. This result should be obtained for individuals with tonal tinnitus as well as tinnitus with wider bandwidths because audiometric function is similar among these tinnitus types [16]. Independent studies by laboratories in France [17], Canada [9], and New Zealand [18] have confirmed this prediction (see Fig. 13.2). A further implication is that post-masking suppression of tinnitus by band-limited noise maskers (called “residual inhibition,” or RI, in the tinnitus literature) should increase proportionately as the center frequency (CF) of the masking sound enters the tinnitus frequency region. This is because these masking sounds (which are presented at intensity levels exceeding the hearing threshold and the tinnitus sound) should reinject feed-forward inhibition into the affected regions of the cortical tonotopic map, temporarily disrupting the synchronous activity underlying tinnitus and weakening the tinnitus percept. This prediction has also been confirmed (Fig. 13.2; from [9]). It should be noted that RI does not appear to be caused by habituation of the affected neurons to frequencies contained in the masker. On the contrary, these neurons are actually more easily driven by amplitude-modulated sounds presented to the tinnitus frequency region during RI than during tinnitus (see Fig. 13.3, from [19, 20]), possibly because their capture by synchronous network activity underlying tinnitus has been disrupted. Rapid rescaling of subcortical auditory input to the frequencies contained in the masker could also contribute to this effect [21]. Other brain imaging results that support the neural synchrony model include evidence for (1) a degraded frequency (tonotopic) representation above ~2 kHz in the region of primary auditory cortex in individuals with tinnitus compared to controls [22] (this reorganization resembling that seen in animal models of hearing loss) and (2) increased spontaneous oscillatory brain activity in individuals with tinnitus [23]. The latter effect tracks the laterality of the tinnitus percept and may reflect augmented network underlying this condition.

As described here, the neural synchrony model accords an important role to the primary auditory cortex in the generation of tinnitus percepts. However, neuron response properties, including increased spontaneous activity and map reorganization, are also altered by hearing loss in subcortical auditory structures [24, 25], although neural synchrony in these regions has not yet...
been studied. Changes occurring in subcortical structures could be projected to the primary cortex and determine some of the effects seen there, as well as some distinct properties of tinnitus including its modulation by somatosensory inputs in many patients [26, 27]. Alternatively, the changes seen in subcortical nuclei could be sculpted by returning output from the auditory cortex, which may recruit a brain network supporting tinnitus percepts. Functional brain imaging studies have implicated several brain areas in tinnitus [28–31], including frontal and limbic areas that may subserve, respectively, the attentional and emotional aspects of tinnitus described by Jastreboff [32] in a comprehensive model of tinnitus published more than a decade ago.

These lines of evidence pointing to a role for neural synchrony in tinnitus have implications for how sensory training might best be conducted (see Chap. 72). The neural synchrony hypothesis implies that the goal of training should be to disrupt the synchronous neural activity believed to underlie tinnitus percepts.

When significant residual hearing is present (delivered by surviving on-target thalamocortical projections or thalamocortical radiations), this goal could be attempted by training suprathreshold sounds in the tinnitus frequency region. These sounds may reinject feed-forward inhibition into the tonotopic map and/or rescale neuron transfer functions in subcortical structures to represent the trained frequencies, thereby disrupting neural synchrony and strengthening thalamocortical synapses previously down-regulated by abnormal synchronous network behavior. Maskers that induce RI may operate in a similar fashion, although repeated induction of RI does not appear to convey a lasting benefit [33], at least in the absence of active auditory training. Alternatively, acoustic training in the region of normal hearing could convey uncorrelated inputs into the affected map region via lateral connections, disrupting neural synchrony or suppressing it by lateral inhibition. Before considering research on various approaches (see Chap. 72), it is...
useful to briefly consider what is known about how auditory remodeling works in individuals with normal hearing, and how it may contribute to the development of tinnitus.

**Neuroplastic Remodeling in Tinnitus**

A feature common to the neural synchrony model and the wider framework of Jastreboff [32] is a role for neural plasticity in the generation of tinnitus percepts. Although direct evidence is lacking and not easily procured, there are compelling reasons to propose a role for such mechanisms in tinnitus. Spike-timing-dependent plasticity [34] appears to be a general property of cortical neurons, and this mechanism, acting in concert with increased spontaneous firing rates consequent on inhibitory deficits and homeostatic plasticity [12], would be expected to facilitate the formation of synchronous networks in regions of the primary auditory cortex affected by hearing loss. Synchronous activity appears to be expressed over cortical distances that exceed those expected from thalamocortical radiations, which implicates temporal coincidence mediated by horizontal fibers as a driving mechanism [14]. From the limited data available, it appears that cross-correlated activity develops within hours of hearing loss and grows over time [13], although the limit of this growth is not known. Neural plasticity has the potential to explain the variability that is seen in tinnitus percepts among affected individuals, with the addition of no new principles.

In the last 15 years, much has been learned about how neural plasticity remodels auditory representations in normal hearing animals. Experience with sound has a profound effect on tonotopic organization and the tuning properties of auditory neurons in the developing brain [35, 36] and after maturity as well [37, 38]. Neural modeling during development appears to be driven largely by the spectrotemporal statistics of the acoustic input, such that neural representations become tuned to the sounds present in the animal’s environment. After maturity, top–down mechanisms begin to play an additional role, preferentially gating neural plasticity in the auditory cortex for sounds that are important for behavioral goals [6, 39]. Several response properties are affected by acoustic training in mature organisms, including shifts in the tuning preference of auditory neurons toward the trained stimuli [6, 7], spike rates induced by these stimuli [40, 41], tuning bandwidth [42, 43], response latency in post-stimulus time histograms [41, 42], and tonotopic map expansions for the trained sounds [44]. However, passive immersion in a distinctive acoustic environment can still have profound effects on neuron response properties and neural organization in the adult brain [38], which may reflect, at least in part, changes in subcortical auditory nuclei that are driven unselectively by stimulus input. These broad principles derived from animal studies appear to be applicable to humans as well [45–48], although much remains to be discovered about the specific rules that guide remodeling in both domains and the mechanisms that underlie them.

Whether these principles apply as well to individuals with tinnitus is less well established. A brain response that is relevant to this question is the stimulus-driven “auditory steady-state response” (ASSR, shown earlier in Fig. 13.3b). This response is evoked in the electroencephalogram by sounds that are amplitude modulated (AM) near 40 Hz, localizes tonotopically to cortical sources in the region of primary auditory cortex, and gives a picture of changes occurring in or projecting to this region during auditory training (see Fig. 13.4a). In individuals with normal hearing, acoustic training to detect single pulses of enhanced amplitude in a 40-Hz AM 2 kHz sound of 1-s duration has been found to modify temporal population activity expressed in the primary auditory cortex. This effect is expressed as an advance in the phase of the ASSR, which reflects a reduction in the time delay between the 40-Hz response and stimulus waveforms (see Fig. 13.4b). The phase advance is a robust phenomenon that consolidates after 24–72 h, increases with continued training, relates perceptual performance, and does not require explicit behavioral training for its appearance [48]. ASSR amplitude is also increased by auditory training, implying more neurons depolarizing synchronously to represent the trained sound [48]. However, the training effect on ASSR amplitude lags that on phase, does not correlate well with perception, and is not observed when multiple sound frequencies are presented during training [49].

These results are from individuals with normal hearing who were studied in order to discover rules that guide remodeling in the human brain. What happens when individuals with tinnitus are trained? The answer to this question is presently not well established.
In a preliminary study [20], we found that while a group of control participants age matched to a tinnitus group showed the expected phase advance when trained on a 5-kHz 40-Hz AM sound ($n=11$, $p=0.006$), only two of eight participants with tinnitus did so, resulting in a nonsignificant group effect overall ($p=0.44$, see Fig. 13.4c, upper panel). It is possible that synchronous neural activity underlying tinnitus may have obstructed or reset training effects in the primary auditory cortex of the participants who had tinnitus (5 kHz was chosen for study because it is in the tinnitus frequency range). However, remodeling of secondary auditory cortical areas appeared to be normal in those who had tinnitus. The P2 (latency ~180 ms) auditory-evoked potential, which localizes to cortical sources in this region and is known to be highly plastic [49], showed a normal enhancement in both groups after auditory training (Fig. 13.4c, lower panel). Several other long latency (>100 ms) auditory evoked potentials localizing to secondary cortex or beyond are known to increase with acoustic training in the laboratory in normal hearing individuals (in order of increasing latency: N1 [50], N1c [49], Ta [51], P2 [47–49, 52], N2 [53], MMN [54]), or to be enhanced for musical sounds in trained musicians (N1c [55], P2 [55–57], anterior frontotemporal sources [58], induced frontotemporal...

**Fig. 13.4** Effects of auditory training on auditory-evoked potentials. (a) Response evoked by a 2-kHz tone amplitude modulated at 40 Hz (ASSR). The stimulus waveform and the response waveform recorded at electrode Cz are shown, together with the bipolar scalp topography (128 sensors). In inverse modeling, the cortical generators for an ASSR evoked by a carrier frequency of 4,100 Hz localized medial to those for an ASSR evoked by a carrier frequency of 250 Hz, in the region of primary auditory cortex. (b) Compass plots showing the amplitude (vector length) and phase (vector angle) of the ASSR at each of 128 sensors, before (left panel) and after (right panel) seven sessions of acoustic training. Individuals with normal hearing who did not have tinnitus ($n=9$) were trained to detect a single 40-Hz AM pulse of enhanced amplitude in a stimulus of 1-s duration (carrier frequency 2 kHz). A phase shift of $23^\circ$ was observed ($p<0.001$, advance of the response waveform toward the stimulus waveform), but the amplitude enhancement did not reach significance. (c) Upper panel: The phase shift (over seven sessions of training) did not reach significance in the participants who had tinnitus ($p=0.44$) but was present in their age-matched controls ($p=0.006$). In both groups, the carrier frequency was 5 kHz (in the tinnitus frequency region of the individuals with tinnitus). Negative values indicate a shift of ASSR phase toward the stimulus waveform. Lower panel: The P2 transient-evoked response (latency ~180 ms) increased with training in both groups, suggesting that secondary auditory areas are remodeled normally in individuals who have tinnitus (cf. [49]).
gamma oscillations [59]). These evoked potentials reveal a distributed neural system for auditory (and perhaps other) learning in the human brain that may overlap with neural structures involved in tinnitus. However, the behavior of the responses during acoustic training in tinnitus is unknown.

Most studies of human auditory learning have employed active training procedures in which adults attended to and processed the sound stimuli while making discriminative decisions. However, there is growing evidence that remodeling of equal magnitude occurs when the sounds are presented as background cues, even when individuals are engaged in watching a subtitled film and have no knowledge of auditory task structure [47, 48, 60]. The ASSR and P2 effects described above were remodeled equally by active training, compared to when the auditory stimuli were presented passively as background sounds to individuals with normal hearing [48]. Animals housed in distinctive sound environments with no processing demands also display significant auditory remodeling, even in adulthood [37, 38]. A working hypothesis based on animal data is that these effects are produced by a rescaling of neuron input/output transfer functions in subcortical auditory structures by fundamental mechanisms that are stimulus driven and expressed in the auditory cortex throughout the lifespan. Explicit auditory training may produce additional changes mediated by attention, but this more mature mechanism is not a prerequisite for remodeling. The fact that auditory representations are modified by passive as well as active exposure could be good news for tinnitus, to the extent that arduous training regimens may be avoided.

Overview and Conclusion

Animal research in the last two decades has established that neural plasticity is a fundamental property of neurons in the auditory system. Evidence has also accumulated that hearing loss leads to changes in central auditory pathways, including tonotopic map reorganization and increased neuron firing rates in primary auditory cortex that may be forged by neuroplastic mechanisms into abnormal synchronous network behavior that generates tinnitus. In this Chapter, I have summarized physiological, psychoacoustic, and brain imaging evidence pointing to a role for neural synchrony in tinnitus.

Also reviewed were results from animal research indicating that cortical representations for sound in the primary auditory cortex are not fixed after early development as was once believed, but can be modified by auditory training well into adulthood. The findings have spawned renewed research into the question of whether tinnitus can be reduced or eliminated by acoustic training designed to normalize aberrant auditory neural representations. For this goal to be achieved, it must be possible to modify auditory representations by acoustic training in individuals with tinnitus, and the neural modifications induced by training must intersect with tinnitus mechanisms. Preliminary research suggests that areas of secondary auditory cortex remodel normally in individuals with tinnitus compared to normal controls, although whether this is true of the primary auditory cortex requires further study.

Acknowledgments

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References


18. Kay F (2008) Towards improving the assessment of tinnitus pitch. Section of Audiology, Faculty of Medical and Health Sciences, University of Auckland.


Chapter 14
Similarities Between Tinnitus and Pain

Aage R. Møller

**Keypoints**

1. Both pain and tinnitus have many different forms.
2. Tinnitus and central neuropathic pain are phantom sensations similar to the phantom limb symptoms that occur without any physical stimulation of sensory receptors.
3. Tinnitus and neuropathic pain are typical examples of “plasticity disorders” where the symptoms are caused by plastic changes that are not beneficial to an individual person.
4. Central neuropathic pain and tinnitus have no physical signs.
5. The severity of pain and tinnitus are difficult to assess quantitatively even under laboratory circumstances. Only the patients’ own perception is a true measure of the severity of central pain and subjective tinnitus.
6. The perception of pain and tinnitus is affected by many factors such as actual circumstances, expectation, stress, and a person’s emotional state.
7. Many forms of pain are best described as suffering; the same is the case for severe subjective tinnitus.
8. Pain and tinnitus can have strong emotional components, it often prevents or disturbs sleep, and it can interfere with or prevent intellectual work.
9. It is difficult to get reliable data on epidemiology of tinnitus and central neuropathic pain because of their subjective nature and large variability.
10. Activation of neural plasticity is involved in causing and maintaining central neuropathic pain and many forms of subjective tinnitus.

11. The nervous system is the site of the anomalies that cause central neuropathic pain and many forms of tinnitus. Both tinnitus and pain involve a cascade of neural structures.
12. The pathology of the nervous system in some forms of central neuropathic pain is stable in the pathologic state. It may be similar for some forms of tinnitus.
13. Pain that is perceived as escapable uses a different part of the periaqueductal gray than pain that is perceived as inescapable. It is not known if tinnitus also has such distinctions.
14. Severe tinnitus is often accompanied by hyperacusis (lowered tolerance to sounds); pain may be accompanied by allodynia (pain from normally innocuous touch of the skin) hyperpathia (exaggerated reaction to acute pain), and hypersensitivity (lowered threshold for painful stimulation).
15. Some forms of tinnitus and pain can be modulated by electrical stimulation of the skin.
16. Electrical stimulation of several cortical structures can modulate both pain and tinnitus.
17. The sympathetic nervous system can modulate pain and some forms of tinnitus.

**Keywords**  Tinnitus • Pain • Central neuropathic pain • Hyperacusis • Allodynia

**Abbreviations**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>DCN</td>
<td>Dorsal cochlear nucleus</td>
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<tr>
<td>NST</td>
<td>Nucleus of the tractus solitaries</td>
</tr>
<tr>
<td>PAG</td>
<td>Periaqueductal gray</td>
</tr>
<tr>
<td>TENS</td>
<td>Transderm electrical nerve stimulation</td>
</tr>
<tr>
<td>VCN</td>
<td>Ventral cochlear nucleus</td>
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<tr>
<td>WDR</td>
<td>Wide dynamic range neurons</td>
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Introduction

It was Jürgen Tonndorf [1] who first drew attention to the similarities between tinnitus and pain. Other investigators have later elaborated on the many similarities between tinnitus and severe chronic pain (central neuropathic pain) [2–5]. Activation of neural plasticity is involved and both are examples of “plasticity diseases” [6]. Pain gets far more attention than tinnitus. The fifth edition of the Wall and Melzack’s Textbook of Pain has over 1,200 pages; Weiner’s Pain Management has over 1,500 pages. Textbooks devoted to tinnitus are essentially non-existent (this book is the first textbook on tinnitus). The research literature on pain is far greater than that on tinnitus; a search in PubMed came up with approximately 400,000 articles about pain vs. approximately 6,000 for tinnitus. Literature about hyperacusis and phonophobia is sparse.

Many forms of tinnitus have similarities with central neuropathic pain, in that activation of neural plasticity is involved in creating the symptoms. Central neuropathic pain is a particular condition where the symptoms are caused by abnormal activity in populations of neurons in the spinal cord and brain that occurs without signals from receptors in the body. Subjective tinnitus and central neuropathic pain are phantom sensations where the sensations are not elicited by activation of receptors. Central neuropathic pain and some forms of tinnitus are symptoms with very few, if any, objective signs. Despite that, both central neuropathic pain [7] and severe tinnitus [8] can affect a person’s entire life, the entire family, as well as social and working relationships. Both these disorders may prevent or disturb sleep and interfere with intellectual work. There are examples of people who like their work but retire because of tinnitus.

Other similarities include the lack of effective treatment, diverse etiology, and sparse knowledge about the anatomical and physiologic bases for these disorders. The treatments a patient with either one of these disorders may receive depend on the specialty of the physician they choose to consult, and the specific interest of the physician or surgeon. In only a few forms of central neuropathic pain and tinnitus can any underlying disease be found.

There are many forms of tinnitus and many forms of pain. Some common forms of pain such as headache and back pain can be managed by simple analgesics. No such general treatment is known for tinnitus. Peripheral neuropathic pain, migraine, and fibromyalgia are complex pain conditions that have less satisfactory treatments. Central neuropathic pain can often be managed by medication. Brain and spinal cord injuries can cause both pain and tinnitus that are difficult to treat.

A separate chapter (Chap. 15) describes the basic anatomy and physiology of pain. Here, we will discuss the similarities between some forms of subjective tinnitus and central neuropathic pain, both being phantom sensations, with activation of neural plasticity playing a central role in their cause. The similarities between treatment of pain and tinnitus are discussed in chapter 94.

Common Features of Subjective Tinnitus and Central Pain

Both central neuropathic pain [9, 10] and subjective tinnitus [11] have many different forms (see Chap. 2). Different forms of disorders with the same name cause difficulties in studying their pathologies and treatments. It would be more appropriate to consider both central and subjective tinnitus as groups of different disorders rather than a single disorder.

Most forms of subjective tinnitus and central neuropathic pain are phantom sensations, which mean that the symptoms are not caused by physical stimulation, but are similar to symptoms that occur after amputations. This is known as phantom limb syndrome, where pain and other sensations are felt as if coming from the limb that no longer exists [12]. The symptoms of many forms of tinnitus and central neuropathic pain are felt as coming from a different anatomical location than the actual pathology and physiological anomaly that cause the symptoms. The anatomical locations of the pathologies of most forms of central pain and most forms of tinnitus are in the brain, although the pain is often referred to a specific part of the body. This is well known from studies of central pain and it is also evident from some observations regarding tinnitus. Tinnitus is often referred to the ear, although tinnitus may occur in deaf people and after severance of the auditory nerve, thus similar to the pain that is felt as coming from an amputated leg. Neural plasticity activated by the absence of input from receptors in an amputated limb is the main cause of the phantom limb syndrome.
Some forms of tinnitus are caused by deprivation of auditory inputs as evidenced from the fact that tinnitus can be caused by middle ear disorders (see Chap. 34) and disappears when sound conduction to the ear is restored, either by treating the conductive pathology such as otosclerosis or by a hearing aid (see Chaps. 56 and 76) or cochlear implants (see Chap. 77). Many people get tinnitus when placed in a silent environment [13].

Neural plasticity plays an important role in creating central tinnitus and central pain, and activation of neural plasticity also plays a role in tinnitus that is caused by pathology in the ear and in acute pain caused by stimulation of nociceptors.

Both tinnitus and central neuropathic pain are examples of harmful effects of plastic changes, thus forms of “plasticity disorders” [6] caused by neural plasticity going awry. Activation of such maladaptive neural plasticity causes abnormal neural activity and re-routing of information. Activation of neural plasticity is involved in many forms of subjective tinnitus and central neuropathic pain as has been discussed in other parts of this book (see Chaps 10 and 15). The way plasticity is turned on is often unknown and probably more complex than what it is for creating phantom limb symptoms.

Both pain and tinnitus can cause suffering; that may be a different condition than tinnitus and pain that does not cause suffering. Tinnitus that causes suffering, or is “bothersome” [14], may activate other neural circuits than tinnitus that does not have these qualities. It has been shown that pain that is “escapable” and pain that is perceived as being “inescapable” activate different parts of a neural structure, the periaqueductal gray (PAG) [15], and different parts of the hypothalamus and midbrain [16].

Peripheral processes can contribute to the initiation of chronic neuropathic pain as well as many forms of tinnitus. Peripheral and central sensitization have been shown to play an important role in the creation of hyperactivity that is the cause of central neuropathic pain [9, 10]. The same is probably the case for tinnitus, although it has not been studied to the same extent as pain [17]. The fact that different mechanisms can initiate processes that result in changes in the central nervous system that cause many forms of tinnitus may explain some of the differences in the symptoms that patients experience [9, 18]. The same is the case for central pain.

Another similarity between tinnitus and pain is that the severity of these disorders cannot be substantiated by objective tests. Even health care professionals may sometimes misjudge the severity of these diseases. Individuals with tinnitus as well as individuals with pain have no attributes of illness and therefore do not attract much attention and sympathy. Relatives and friends may doubt the seriousness of their diseases. In the absence of objective test results, health professionals may even sometimes think that their patients may be malingering. This makes both tinnitus and central pain disorders some of the most challenging disorders for clinicians.

Prevalence of Central Pain and Tinnitus

One of the problems in getting reliable epidemiologic data is similar for pain and tinnitus, namely that the definition of the severity varies among individuals with these conditions. These problems are greater for central neuropathic pain than other neuropathic pain conditions, and it is greater for tinnitus than for other hearing disorders, such as hearing loss from exposure to noise (see Chap. 37), which have been studied extensively as has age-related hearing loss (presbycusis) (see Chap. 36). No reliable information about the epidemiology of central neuropathic pain is available, nor is the prevalence of other chronic pain conditions such as peripheral neuropathic pain that commonly occurs in individuals with diabetes neuropathy completely known [18, 19].

The prevalence of chronic neuropathic pain may be greater than commonly assumed and its prevalence is likely to increase in the future. This is very similar to tinnitus, where the prevalence seems to increase. The prevalence of both central pain and tinnitus increases after middle age, which means that age-related changes add to the factors that cause tinnitus and pain (see Chap. 36).

Tinnitus is estimated to effect 13–20% of the overall population of the United States [20]. These complaints, often associated with hearing loss, increase with age to 27–34% of the population older than 70 years reporting significant tinnitus [21]. Twenty percent to 45% of tinnitus sufferers also have hyperacusis; a few individuals only have hyperacusis [22] (see also Chap. 5).

One reason for the increased incidence of tinnitus is the increased occurrence of head injuries (see Chap. 67), which also is associated with pain conditions. From 10 to 30% of people with spinal cord injuries have central
pain. Individuals with head injuries often have central pain and tinnitus [23, 24] (see Chap. 67). After strokes, 1–8% have central pain [19, 25].

The prevalence of post-surgical neuropathic pain has been estimated to be 2–3% of the population in the developed world [18]. This problem is poorly recognized. Equally poorly recognized is postoperative tinnitus. It occurs often after surgical removal of vestibular schwannoma where the concerns are about preserving facial function and hearing, which has improved after introduction of intraoperative neurophysiologic monitoring [26]. However, little is known about how to reduce the risk of tinnitus.

**Neuroanatomical Similarities Between Tinnitus and Pain**

The neuroanatomy of hearing and pain has many similarities. The neural pathways for acute pain have similarities with the classical and non-classical ascending auditory pathways. The medial tract of the spinothalamic system may be regarded as the non-classical pathways of the somatosensory system (see Chap. 15). The fibers of the lateral spinothalamic tract terminate in neurons in the ventral thalamus corresponding to the classical pathway, whereas the medial spinothalamic tract terminates in the dorsal and medial thalamus and thus resembles the non-classical pathways of other sensory systems. The lateral spinothalamic tract provides information about the location of the pain and the medial tract provides information about the nature of the pain (see Chap. 15).

The medial and dorsal thalamus have subcortical connections to several regions of the brain, such as the limbic system, and the neurons in the cortical projections of the dorsal thalamus bypass the primary somatosensory cortex. These neurons terminate directly on neurons in the secondary and association cortices while the classical pathways project to primary cortices.

**Functional Similarities Between Pain and Tinnitus**

Tinnitus has similarities with several characteristics of central neuropathic pain. Repeating painful stimulations causes increasing intensity of pain, known as the “wind up” phenomenon [27]. When a noxious stimulation is repeated at a short interval, the pain from the second presentation feels stronger. This is thus a form of abnormal temporal integration of painful stimulation. In other studies, it has been shown that temporal integration of pain signals is different in individuals with signs of neuropathic pain and individuals without central neuropathic pain [28].

A few similar studies have been done regarding temporal integration of sound in individuals with tinnitus [29], but animal experiments indicate that strong sound stimulation changes the temporal integration in the inferior colliculus as assessed using evoked potential techniques [30].

**Sensitization and Modulation of Pain and Tinnitus**

It is well known that peripheral and central sensitization can play important roles in creation of pain. Together with re-organization of neural circuits, this is regarded as the cause of central neuropathic pain. Evidence is accumulating that similar processes affecting the auditory system may play important roles in some forms of tinnitus.

**Peripheral Sensitization**

There are several ways in which peripheral sensitization of receptors in the body and the ear can contribute to pain and tinnitus. One way is through activation of the sympathetic nervous system, which can cause sympathetic nerve fibers that terminate near receptors to secrete norepinephrine, which increases the sensitivity of the receptors. Epinephrine secreting nerve fibers have been identified near receptors in the skin and close to the receptors (hair cells) in the cochlea [31]. Thus, the fact that sympathectomy is an effective treatment for tinnitus when it is a symptom of Ménière’s disease [32] indicates that the sympathetic nervous system is involved in at least the kind of tinnitus that occurs in Ménière’s disease.

The sympathetic nervous system may even activate the receptors without external stimulation, so they send information to the nervous system similar to when
normal stimulation of the receptors occurs with physical stimuli. The most extreme of such sympathetically induced pain is reflex sympathetic dystrophy (RSD), now known as complex regional pain syndrome type I [33].

Central Sensitization

Certain kinds of neurons in the dorsal horn of the spinal cord (and the trigeminal nucleus), known as the wide dynamic range (WDR) neurons, are believed to have important roles in central sensitization of pain circuits ([9, 34, 35], see also [36]). Activation of neural plasticity that can change synaptic efficacy also plays an important factor in creating the abnormal states of the neural circuits in the dorsal horn associated with central pain [37] (see Chap. 15).

Activation of neural plasticity in the neural circuits of the dorsal horn is important because it can change the excitability of neurons (central sensitization) and re-route information by making dormant synapses become active [38]. Also, it can make synapses that are normally active become dormant.

It has been hypothesized that reorganization of the neural circuits in the spinal cord plays an important role for creation and maintaining central neuropathic pain. Doubell has proposed that the pain circuits in the dorsal horn of the spinal cord (and the trigeminal nucleus) can operate in four main different states [34] (see Chap. 15).

Similar hypotheses may apply to some forms of tinnitus, but hypotheses about the pathology of tinnitus are less uniform and less detailed. Studies in animals in which tinnitus conditions were induced by deprivation of input to the auditory system [39] or by overstimulation [40] have shown evidence that some neurons in the inferior colliculus have the ability to change their function in a similar way as the WDR neurons.

Interaction Between Sensory Systems

The old concept that certain functions of the brain are done in specific parts of the brain has gradually been eroded. It has become more and more evident that considerable interaction between many systems in the brain and the spinal cord occurs normally, as well as in diseases where certain interactions have adverse and harmful effects. It was earlier regarded as an axiom that the information from the different sense organs was processed in specific and separate parts of the brain.

Anatomical Aspects

We have discussed in Chap. 10 how somatosensory signals can interfere with hearing when the non-classical auditory pathways are active such as it is in children [41] and in some individuals with tinnitus [42]. This can occur in two different ways. One way is through connections that neurons in the dorsal column nuclei and the trigeminal nucleus make with neurons in the dorsal cochlear nucleus (DCN) [43–45] (see Chap. 9). The other way is through activation of the non-classical ascending auditory pathways, which receive input from other senses through connections to the inferior colliculus [46] (see Chap. 8).

Physiologic Signs of Cross-Modal Interaction

Certain anomalies of sensory systems in individuals with central pain have similarities with anomalies that occur in connection with some forms of tinnitus. One such anomaly is cross-modal sensory interaction, which means that the perception of one sensory modality can be affected by stimulation of another sense.

It has been known for a long time that acute pain sensations elicited by stimulation of pain receptors can be modulated by stimulation of nerve fibers, which innervate receptors that mediate innocuous sensory stimuli (touch, etc.). This is a normal phenomenon involving Aβ fibers in the spinal cord, which have inhibitory influence on cells that receive nocuous input from pain receptors (Fig. 14.1). This fact is used in treatment of pain, using electrical stimulation of the skin. This method is in routine use under the name of transderm electrical nerve stimulation (TENS) [47], and it has shown effectiveness for acute pain [48] as well as central pain [49]. It relies of stimulation of large sensory fibers, which can have an inhibitory influence on pain cells in the spinal cord and activate neural plasticity and thereby is effective in reducing pain not caused by activation of pain receptors (central neuropathic pain).
Modulation of tinnitus by activation of the somatosensory system [50, 51] has been demonstrated by electrical stimulation of the median nerve at the wrist [42], manipulation of neck muscles [52, 53], from temporomandibular problems [54, 55], and from changing one’s gaze [50, 56, 57]. Functional imaging studies indicate that gaze-evoked tinnitus is caused by neural activity associated with eye movements that enters the auditory system [58]. These effects seem to be mediated through cross-modal interaction between the auditory system and the somatosensory system.

The anatomical and physiologic bases for these interactions are not as well known as the modulation of pain by somatosensory stimulation. Electrical stimulation of the median nerve at the wrist [42], manipulation of neck muscles [52, 53], from temporomandibular problems [54, 55], and from changing one’s gaze [50, 56, 57]. Functional imaging studies indicate that gaze-evoked tinnitus is caused by neural activity associated with eye movements that enters the auditory system [58]. These effects seem to be mediated through cross-modal interaction between the auditory system and the somatosensory system.

The other way that somatic stimulation can affect the auditory system is through the non-classical ascending auditory pathways [64]. The non-classical auditory pathways receive input not only from the ear but also from other sensory systems such as the somatosensory system [45, 46] (see Chap. 8).

A different kind of interaction on pain [65] and possibly tinnitus is that from the vagus nerve. Earlier, little attention was paid to the vagus nerve; the focus has been on the motor functions of the vagus nerve. However, approximately 80% of the nerve fibers are afferent fibers. The discovery that electrical stimulation of the vagus nerve could treat epilepsy renewed the attention to the vagus nerve. Electrical stimulation of the vagus nerve is an approved treatment for epilepsy in the US [66] and is in clinical use for controlling epileptic seizures. Electrical stimulation has also been used for treatment of depression.

Electrical stimulation of the left vagus nerve has been shown to suppress some forms of pain [65] (see Chap. 94), and research is now aimed at other applications such as treatment of depression and control of severe tinnitus.

Afferent vagus nerve fibers terminate in the nucleus of the tractus solitaries (NST), which connect to many parts of the brain. The vagus nerve supplies cholinergic input to many structures in ways that have similarities to that of the basal nucleus of Meynert, which provides arousal and promote cortical plasticity [67, 68].

Central neuropathic pain is often associated with allodynia (pain from light touch stimulation). This may be similar to the rarely reported perception of sound by rubbing the skin, such as by a towel.

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Chapter 15
Anatomy and Physiology of Pain

Aage R. Møller

Keypoints

1. Pain is a subjective sensation that has no objective correlates.
2. Pain has many forms, and the perception of pain is affected by many factors including actual circumstances, expectation, stress, and the emotional state of the person.
3. Many forms of pain are best described as suffering.
4. Pain may be divided into two large groups: pain that is caused by direct stimulation (physical or chemical) of specific receptors (nociceptors) and pain that is not caused by stimulation of nociceptors. There is also a third kind of pain in which activation of neural plasticity plays an important role.
5. Stimulation of nociceptors that are located in the skin, the cornea, tooth pulp, muscles, joints, peripheral nerves, the respiratory system, and viscera causes acute pain that has both a fast and a slow component.
6. Pain can also be caused by trauma or inflammatory processes generated in nerves or in the nervous system and not from activation of specific pain receptors.
7. Expression of neural plasticity can create pain (central neuropathic pain) that is caused by neural activity in the brain without peripheral input.
8. Transmission of pain in the dorsal horn of the spinal cord (and the trigeminal nucleus) can be modulated by input from skin receptors (A\textbeta fibers) and descending activity from supraspinal sources.
9. The sympathetic nervous system can modulate the sensitivity of nociceptors and the transmission of pain signals in the spinal cord and the trigeminal nucleus.
10. Activation of neural plasticity can also cause change in processing of nociceptor-elicited pain signals causing hyperpathia (exaggerated and prolonged response to painful stimuli) and allodynia (painful sensation from light touch of the skin).
11. The vagus nerve is involved in some forms of pain, and electrical stimulation may reduce pain.

Keywords Somatic pain • Visceral pain • Neuropathic pain • Central neuropathic pain • Neural plasticity

Abbreviations

CNS Central nervous system
DLPT Dorsolateral pontomesencephalic tegmentum
IASP International Association for the Study of Pain
NA Norepinephrine
NST Nucleus of the solitary tract
PAG Periaqueductal grey
RVM Rostral ventromedial medulla
SI Primary somatosensory cortex
STT Spinothalamic tract
VPI Ventral posterior inferior (thalamus)
VPL Ventral posterior lateral (thalamus)
VPM Ventral posterior medial (thalamus)

Introduction

Some forms of pain have similarities with tinnitus, as discussed in another chapter (Chap. 14). There are further anatomical and physiologic aspects of pain that
make it appropriate to include a chapter on the anatomy and physiology of pain in a book on tinnitus. Pain can have many forms; it can be constant or intermittent. It can cause little or moderate discomfort or it can be disabling, preventing sleep and intellectual work. Pain can cause fear of a serious disease, and its perception can change just from assurance that it is not a sign of a serious disease. Pain that is not a sign of a serious disease, thus, mainly affects quality of life. Pain that is not a sign of a serious disease usually receives little attention from health care personnel.

Pain is related more to suffering than to any other quality; but again, there is large individual variation. Helplessness and expectations are important.

Pain is the most common reason for visits to the emergency room and plays an important role in diagnosis of many forms of diseases. However, training of physicians in this particular area is often inadequate in the US and falls short of providing the basis for effective treatment and care for patients with various degrees of pain.

Pain is a subjective sensation that lacks objective signs; it cannot be measured with any clinical methods, only the patient’s own description can provide information about its strength, character, and the location on the body to which it is referred. Estimates of the intensity of pain can be obtained using a visual analog scale, but it still depends on the individual’s judgment about the pain.

The basic research that is devoted to pain is much less than what seems justified by the degree of suffering from idiopathic pain. Many forms of pain are in many ways an enigma. Many forms of pain are not caused by diseases. Available treatments are often ineffective, and some treatments cause severe side effects. Different kinds of narcotics are effective pain treatments but are restricted by legal measures because of fear for addiction or because physicians hesitate to prescribe them because of fear of legal actions.

Pain is often regarded as a somatosensory sense and is often discussed in textbooks together with sensory systems. However, the sensation of pain is more complex than somatosensory sensations; it is a much more variable sensation than somatosensory perceptions such as touch, vibrations, and warmth and cool.

It has been said that the only pain that is tolerable is someone else’s pain. This chapter provides a brief description of the anatomy and physiology of pain. More detailed descriptions can be found in Wall and Melzack “Textbook of pain” and in Møller “Neural plasticity and disorders of the nervous system”.

### Different Kinds of Pain

Pain can be divided into two broad classes of acute and chronic pain according to how long the pain has lasted. The term “chronic pain” is usually used for pain that lasts more than 3 months. However, chronic pain is not related to the pathology of the pain or to its etiology [1–3] and thus is an arbitrary distinction. The International Association for the Study of Pain (IASP) regards pain that lasts more than 6 months to be chronic pain. Chronic pain may be caused by diseases such as rheumatoid arthritis but is often caused by re-organization of the central nervous system (CNS) (central neuropathic pain) or a combination of these two causes.

Another way of categorizing pain is as somatic and visceral pain. Somatic pain is caused by activation of pain receptors in the skin, muscles, joints, etc. Visceral pain originates in viscera from mechanical or chemical stimulation, including inflammation. Another way of dividing pain is in somatic pain, visceral pain, and neuropathic pain. Devor and Boive [4, 5] have defined three main types of pain (see Fig. 15.1). There is considerable overlap between these kinds of pain [4]. The term “neuropathic pain” describes pain that originates in the nervous system. It can be divided into peripheral and central neuropathic pain. The term “neuropathic pain” theoretically covers all pain caused by nerves, the spinal cord, and the brain, but the term is used by neurologists to describe pain caused by peripheral nerves and cranial nerves. The term “central neuropathic pain” is used for pain caused by abnormal neural activity in the central nervous system (spinal cord and brain). This kind of pain occurs without input from pain receptors. Central neuropathic pain is a phantom sensation that

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1International Association for the Study of Pain (IASP) definition of pain: An unpleasant sensory and emotional experience associated with tissue damage or potential damage or described in such terms.

2René Leriche, French surgeon, 1879–1955

3Neuropathic pain: Neurologists use the term “neuropathic pain” only in reference to pain from peripheral and cranial nerves, although the term relates to pain from the nervous system in general.
has similarities to some forms of tinnitus. All other forms of pain are caused by activation of pain receptors. Pain receptors are localized in the skin, muscle tendons, fascia, and viscera. Heat, chemicals, and inflammatory processes can activate pain receptors.

Central pain is caused by a lesion or dysfunction in the CNS [3], according to the IASP.

Central neuropathic pain is caused by abnormal neural activity in the CNS that may be caused by functional re-organization of the nervous system, most likely elicited through activation of neural plasticity and thus not caused by activation of pain receptors. Central neuropathic pain is a “phantom” sensation similar to that experienced from amputated limbs and tinnitus. Phantom sensations are caused by the expression of neural plasticity [7, 8] and are therefore “plasticity diseases” similar to tinnitus [9]. The term “central neuropathic pain” is used to distinguish pain that is not caused by morphologically verifiable lesions of nerves in the CNS [10] from causes that have morphological or chemical correlates.

These different categories of pain may overlap and interact with each other, and, in particular, central neuropathic pain may begin with somatic pain of some kind.

Pain causes many different reactions and is often associated with activation of other parts of the nervous system than those that are traditionally regarded as

\[\text{Normal} \quad \text{Inflammation} \quad \text{Neuropathic}\]

**Fig. 15.1** A classification of pain that defines two main overlapping groups of pain namely, nociceptive pain that can occur as a normal condition and as a result of inflammatory processes; and pathophysiological pain that includes neuropathic pain and pain caused by inflammatory processes [4]. From Møller AR (2006) Neural plasticity and disorders of the nervous system [6]. Reproduced with permission from Cambridge University Press

\[\text{Normal} \quad \text{Inflamation} \quad \text{Neuropathic}\]

**Somatic Pain**

Somatic pain is caused by activation of pain receptors in the body. It can occur from tissue damage, traumatic injuries, and surgical operations. It is a common cause of acute pain. Ischemia also causes pain by stimulation of certain pain receptors. These kinds of pain occur rapidly and are normally short lasting. Inflammation of the skin, joints, and muscles are other common causes of somatic pain, but pain can last a long duration of time if the inflammation is chronic. Unmyelinated axons can grow into scar tissue, such as from operations of the spinal cord and cause central pain [17]. Muscle and joint pain, such as in rheumatoid arthritis, are common causes of chronic pain.

**Visceral Pain**

Visceral pain is not perceived in the same way as somatic pain. The pain is not felt at the anatomical location of the cause of the pain. Pain that originates in the viscera and the heart is often referred to locations on the surface of the body [1, 18–20]. Such pain is known as referred pain. The location of the pain is less specific than somatic pain and varies among individuals [1]. Visceral pain often has an emotional component such as being perceived as inescapable [21].

\[\text{Normal} \quad \text{Inflammation} \quad \text{Neuropathic}\]

\[\text{Normal} \quad \text{Inflammation} \quad \text{Neuropathic}\]

\[\text{Normal} \quad \text{Inflammation} \quad \text{Neuropathic}\]

\[\text{Normal} \quad \text{Inflammation} \quad \text{Neuropathic}\]

\[\text{Normal} \quad \text{Inflammation} \quad \text{Neuropathic}\]

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\[\text{Normal} \quad \text{Inflammation} \quad \text{Neuropathic}\]

\[\text{Normal} \quad \text{Inflammation} \quad \text{Neuropathic}\]
The reason why pain from viscera is poorly localized is most likely related to the fact that none of the secondary neurons in the spinal cord receive only visceral input; there are much fewer visceral afferent fibers than somatic afferent fibers [22]. Another reason may be that the vagus nerve can mediate pain from viscera [23].

**Pathways**

Peripheral nerves that carry signals from pain receptors are layer I and II of the dorsal horn. In the head, pain fibers travel in the fifth, ninth, and tenth cranial nerves to terminate in the caudal (spinal) part of the trigeminal nucleus. Central pain pathways include both ascending and descending pathways. Ascending pain pathways are often regarded as part of the somatosensory system, known as the anterior lateral system.

The different parts of the anterior lateral tracts carry pain information from the spinal cord and the trigeminal nucleus to the reticular formation, the periaqueductal gray (PAG), and to the dorsal and ventral thalamus. From here, pain information can reach several parts of the brain. Some forms of visceral pain are carried in the vagus nerve that terminates in the nucleus of the solitary tract, and from there travel to several parts of the brain [23].

The trigeminal nucleus has similarities with the dorsal horn of the spinal cord. It is an elongated structure in the brainstem that reaches from the midbrain into the upper part of the spinal cord (Fig. 15.2). Its rostral parts are concerned with innocuous stimulation of the skin in the face and mucosa in the nose and

![Fig. 15.2 Pain pathways from the head of the trigeminal nucleus (indicated by dashed rectangle). RF: Reticular formation. Adapted from Sessle [25]. From Møller AR (2006) Neural plasticity and disorders of the nervous system [6]. Reproduced with permission from Cambridge University Press](image-url)
mouth. The caudal part is mostly involved with noxious stimulation and thus pain. The connections to the dorsal cochlear nucleus make this a structure of importance for tinnitus (see Chap. 9). The trigeminal nucleus is the site of pathologies that cause a particular kind of pain, trigeminal neuralgia, that consists of attacks of excruciating pain in one of the three radiations of the trigeminal nerve [24].

**Neural Circuitry in the Spinal Cord**

Dorsal root fibers from pain receptors make synaptic contact with cells in layer I and II of the dorsal horn (Fig. 15.3); most of the axons of these cells cross the midline at the segmental level and ascend in the anteriorlateral tracts [26]. C fibers terminate mainly on cells in lamina II of the dorsal horn (Fig. 15.3) (Rexed’s classification [27]). The axons of these cells make synaptic contact with cells in lamina I. (Lamina I is also known as the substantia gelatinosa.) Aδ fibers terminate on cells in layer I, and collateral fibers of these Aδ fibers terminate in lamina IV and V of the dorsal horn [26].

Some of the interneurons in lamina I send collaterals to segments above and below their own segment. These fibers travel in the tract of Lissauer (dorsolateral fasciculus), forming part of the anteriorlateral tract, mainly the spinothalamic tract (STT) [26] (Fig. 15.4). These cells receive input from nociceptors that respond to different modalities of noxious stimuli [28].

The anterior lateral tract consists of the spinoreticular, the spinotectal, and spinothalamic tracts, the latter being the best known and probably the most important.

Cells in lamina VI, VII, and VIII in the so-called “intermediate zone” receive input from large diameter fibers that innervate receptors for innocuous and noxious (painful) stimulations from large areas of skin. Some cells receive input from viscera.

The neurons in lamina I send axons crossing the midline to form the lateral tract of the STT that ascends toward the thalamus. The lateral STT is crudely organized somatotopically and mediates the magnitude and quality of pain (“What” in Fig. 15.3). The anterior portion of the STT communicates awareness and spatial information about pain (Fig. 15.5 “Where”). The fibers of this tract originate in cells in deeper layers of the dorsal horn, layers V and VII and fibers from cells in the intermediate and also from layer VI, VII, and VIII of the intermediate zone of the dorsal horn.

![Fig 15.3](image1.png)  
**Fig 15.3** Illustration of the termination of Aδ fibers and C fibers in the dorsal horn. DRG: Dorsal root ganglia. Lamina I and II are also known as substantia gelatinosa. From Møller AR (2006) Neural plasticity and disorders of the nervous system [6]. Reproduced with permission from Cambridge University Press

![Fig 15.4](image2.png)  
**Fig 15.4** Schematic illustration of the connections through which innocuous sensory input mediated by large myelinated (Aβ) fibers can inhibit pain neurons in lamina I that receive noxious input from Aδ fiber and C fibers via interneurons and which give rise to axons of the STT. From Møller AR (2006) Neural plasticity and disorders of the nervous system [6]. Reproduced with permission from Cambridge University Press
Large diameter (Aβ) fibers from sensory receptors in the skin respond to innocuous stimuli and project to cells in lamina III, IV, and V [28] (Figs 15.3 and 15.4). The axons of these fibers form the anterior STT tract. Large Aβ fibers that terminate on cells in the dorsal horn can inhibit cells that respond to noxious stimulation, and innocuous sensory input can thereby modulate (inhibit) conduction of pain impulses in the dorsal horn.

The fibers of the STT terminate in several parts of the thalamus; as many as six areas have been identified [28]. The anterior STT targets the ventral thalamus (ventral posterior lateral (VPL), the ventral posterior medial (VPM), ventral posterior inferior (VPI) nuclei, and several nuclei in the medio-dorsal thalamus) [28]. The neurons of the VPL and VPI project to the primary somatosensory cortex (SI). This pathway probably gives rise to the sensation of the fast phase of pain, which is clearly localized. The fibers of the lateral portion of the STT (Fig. 15.3) originate mainly in cells in lamina I of the dorsal horn that receive input from C fibers via neurons in lamina II. This part of the lateral portion of STT mediates the burning sensation of pain.

The lateral tract projects to the dorsal and medial portion of the thalamus from where axons travel to the insula and limbic structures. The cells of these structures project to secondary somatosensory cortices on both sides and to other non-sensory structures, and to some extent to area 3a (SI) [28] (Fig. 15.3) [29] as well.

The lateral tract communicates object information ("What" in Fig. 15.5) about pain and it is responsible for affective qualities of painful stimulation, thus similar to the non-classical pathways of the auditory system (see Chap. 8) that may evoke fear and other emotional reactions to sound in individuals with tinnitus.

The fact that pure C-fiber activation reaches the SI cortex means that C fibers may produce a sensation of pressure or touch, in addition to a sensation of burning pain.

There is considerable individual variation in the pain pathways [30], however, and many of the studies of the neuroanatomy of pain have been done in animals; the results may not be directly applicable to humans.

The spinoreticular tract is mainly bilateral, and its main target is the reticular formation of the brainstem. The spinomesencephalic tract has as its main target the periaqueductal gray (PAG). This means that only the STT has connections to the ventral thalamus, and from there connects to the SI. The spinoreticular and spinomesencephalic tracts are important for control of pain processing. These structures, through descending systems, can modulate traffic in ascending pain pathways and thereby cause suppression and enhancement of pain sensations (see p. xx).

The fibers of many parts of the anterior lateral tracts send collateral fibers to many locations along their ascending paths. Many of these collaterals terminate in the reticular formation of the brainstem, thus affecting wakefulness.

**Neural Plasticity in the Spinal Cord**

Neural plasticity, regarding processing of pain signals in the dorsal horn of the spinal cord (and the trigeminal...
nucleus), is extensive and has been studied in detail. Evidence has been presented that the dorsal pain circuits in the dorsal horn can operate in four different states. Doubell [31] has described these states in the following way:

State 1 is the normal state, where low-threshold mechanoreceptors mediate sensations such as that of touch, vibration, pressure, warmth, or cool. When the spinal cord is in this state, stimulation of high threshold receptors causes localized sensations that are clearly recognized as painful without emotional engagement.

State 2 represents a change in function that is characterized by suppression of transmission of both normal innocuous somatosensory information and the neural activity that normally elicit painful sensations. In this state, descending signals from the brain cause reactions such as “flight or fight”, mediated by the NA–serotonin descending pathways, Fig. 20.

The changes in state 2 represent the way hypnosis, placebo, suggestions, distraction, and cognition can affect (suppress) the perception of painful stimuli. Switching from state 1 (normal function) to stage 2 can be affected by administration of opioids, alpha-adrenergic agents, and GABA_A antagonists (bicuculine). The known freedom of pain that often is present during a short period after an accident is probably an example of the changes that represent state 2.

In state 3 function of the neural circuits in the dorsal horn, the excitability of cells is higher than normal, thus almost opposite to that of state 2. In stage 3, the nociceptive receptive fields of neurons in the dorsal horn neurons becomes larger through activation of ineffective (dormant) synapses [32]. This is presumably caused by an increased synaptic efficacy facilitating neural transmission together with reduced inhibition. Stimulation of sensory receptors thereby elicits larger than normal neural activity and sensory activation that normally does not elicit a sensation of pain causes painful sensitivity. This is believed to be one way that light touch can cause a painful sensation known as allodynia. Similar mechanisms may be responsible for the exaggerated prolonged pain experience from moderately strong painful stimuli known as hyperpathia.

Pathologies of nerves may promote a switch of the function of neural circuits in the dorsal horn to stage 3 [33].

State 4 has many similarities with stage 3; one major difference being that the abnormal conditions are caused by an anatomical re-organization, while the changes in function in stage 3 are caused by functional changes (altered synaptic efficacy, etc.). The changes in morphology that is a characteristic of stage 4 include programmed deaths of cells (apoptosis), degeneration or atrophy of synapses, creation of new synapses, and modification of the contacts between cells and synapses. In state 3, Aβ fibers may make synaptic contact with cells that are innervated by C fibers [31]. Instead of normally terminating on cells in layers III–V of the spinal horns of the spinal cord, they may invade the territories of C fibers (lamina II), which can explain why normally innocuous stimulation is perceived as painful (allodynia).

In summary, state 1 is the normal state of the spinal cord’s dorsal horn (and the trigeminal nucleus); state 2 represents decreased response to painful stimulation. State 3 represents the opposite with abnormally high excitability. Finally, state 4 is a (anatomically) permanent state of such increased excitability that causes permanent pain and redirection of information.

Change from stage 1, the normal function occurs when neural plasticity is tuned on first, causing a change in synaptic efficacy and as a further step when state 4 is reached as a result of structural changes. State 4 is a stable stage that is more difficult to reverse than that of 2 and 3. The change in function that occurs in state 2 probably reverses automatically to the normal stage or to state 3.

**Modulation of Pain**

The best known way to modulate pain is by administering common pain relieving medications such as aspirin, ibuprofen, naprosyn, and by opioids that act on several different opioid receptors (mu, kappa, and delta receptors), that are found in many structures, best known are those found in the brainstem in the RVM and PAG. The COX1 and COX2 enzyme systems are involved in pain and analgesics of various kinds are aimed at modulating this enzyme system, either acting on both COX1 and COX2 or specifically on COX2. Selective COX2 inhibitors were introduced some years ago but were in general found to have unacceptable side effects or/and did not offer
the benefits that were expected in the form of less risk for stomach bleedings.

Pain can be naturally modulated by peripheral mechanisms and by central mechanisms. Peripheral mechanisms involve modulation of the sensitivity of pain receptors, and central mechanisms involve complex descending neural pathways that can control the impulse traffic in the ascending pain pathways. The modulation has often been described as sensitization that increases the sensitivity of pain sensations and as de-sensitization that decreases pain sensations. The anatomical bases for different systems that can modulate acute pain are described below.

**Peripheral Modulation of Pain**

Peripheral modulation of pain consists mainly of sensitization of pain receptors and can occur through secretion of norepinephrine from sympatric nerve fibers that terminate close to the receptors. This kind of modulation is caused by the sympathetic nervous system.

**Descending Pathways**

As in sensory systems, pain pathways have extensive descending pathways (Figs. 15.6–15.8) that exert control over impulse traffic in the ascending pathways. Together with the sympathetic nervous system, these descending pathways can sensitize pain receptors (peripheral sensitization) or block pain impulses from reaching the brain in the spinal cord, such as often occurs after trauma and which is a part of “flight or fight” reactions. The results are often freedom of pain in the first short period after a serious trauma.

Three or four separate descending systems, which can modulate the transmission of pain signals in the ascending pathways, have been identified [34]. These are the rostral ventromedial medulla (RVM) (Fig. 15.6), the dorsolateral pontomesencephalic tegmentum (DLTP) (Fig. 15.7) [35], and the NA–Serotonin pathway. In addition, ascending activity from stimulation of nociceptors can be modulated by the norepinephrine (NA)–serotonin pathway that originates in the brainstem reticular formation (Fig. 15.8). In addition, the vagus nerve may also be regarded as a descending pathway that can modulate pain [36, 37].

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![Fig 15.6](image1.png) **Input to the PAG and pathways through which modulation of transmission of pain signals by the PAG can occur through the RVM pathway.** From Møller AR (2006) Neural plasticity and disorders of the nervous system [6]. Reproduced with permission from Cambridge University Press.

![Fig 15.7](image2.png) **Schematic diagram showing the dorsolateral pontomesencephalic tegmentum pathway (DLTP).** From Møller AR (2006) Neural plasticity and disorders of the nervous system [6]. Reproduced with permission from Cambridge University Press.

The RVM and DLTP pathways originate in supraspinal structures. The modulation occurs mainly by influencing neurons in lamina I and II of the dorsal horn.
The PAG is involved in two of these pathways through at least one interneuron in the RVM. The RVM’s target is neurons in lamina I and II of the dorsal horn [38]. The DLPT also targets neurons in the dorsal horn, but it is mainly excitatory (facilitating). There are three types of RVM neurons, on-cells, off-cell and neutral cells. The on-cells that excite dorsal horn pain cells are inhibited by opioids. The off-cells, that are excited by opioids inhibit pain cells in the dorsal horn. The RVM is the main source of serotonin.

It is worth noting that these descending systems have two parts, an inhibitory and an excitatory part, which means that at least the RVM and DLTP can both enhance and suppress impulse traffic in pain circuits in the dorsal horn and the trigeminal nucleus. There are thus parallel inhibitory and excitatory descending pathways. The net effect of activation of these descending systems depends on the balance between the activity in the inhibitory and excitatory paths. This is one of the anatomical substrates for the complex modulation of pain sensations.

**The Role of the Vagus Nerve**

Much less is known about the neural circuits through which the vagus nerve can modulate pain impulses. Approximately 80% of the fibers in the vagus nerve are afferent (ascending) fibers, but so far most studies have concerned the efferent (descending) fibers of the vagus nerve. The afferent fibers terminate in the nucleus of the solitary tract (NST). Recent studies have found that the nucleus can influence many structures of the brain and the spinal cord, including pain circuits (Fig. 15.9). It is not completely understood how vagus stimulation can control central pain, but the effect seems to be related to the fact that the vagus nerve can influence some of the neurons in the dorsal horn neurons that mediate pain [37, 39]. Studies have shown indications that the vagus nerve may be involved in the opioid induced analgesia. Studies in rats have shown that after severing of the vagus nerve, the analgesic effect of morphine decreases [37]. This means that intact function of the vagus nerve is necessary for the analgesic effect of morphine.

Electrical stimulation of the vagus nerve can reduce the pain from controlled stimulation of nociceptors. It also reduces the temporal integration of pain elicited by consecutive impulses (“wind-up”) as well as pain from tonic pressure [36]. Pain sensation such as from electrical stimulation normally show temporal integration and the threshold at high stimulus rates is much higher than the threshold of sensation, which does not show any noticeable temporal integration for stimulus rates between 1 and 100 pps. In an individual with signs of central neuropathic pain, the temporal integration
for pain is abolished and the difference between pain threshold and that of sensation is much smaller than normal [40].

Studies in cats and monkeys have shown that electrical stimulation of the vagus nerve can attenuate the response from neurons in the dorsal horn to many different types of noxious and innocuous stimuli [41].

The results of many studies indicate that the vagus nerve can affect the central processing of pain (central inhibition). The effect of vagal activity on pain may be mediated through the NA–serotonin descending system (Fig. 15.8) [42] as well as endocrine from the adrenal medullae [39].

The complexity of both the descending pathways and modulation of central pain impulses may explain some paradoxical effects. For example, benzodiazepines that are effective in treating pain caused by muscle contractions may enhance other forms of pain see Møller [6].

Animal (rat) experiments have shown that activity in vagal afferents under the diaphragm can modulate somatic pain impulses such as mechanical hyperalgesia [39]. The effect was induced by endocrine signals released from the adrenal medulla. It was concluded that the brain could control the sensitivity of nociceptors all over the body, even when the effect is elicited for an anatomical far distance. This means that nociceptors are under the control of circulating catecholamines in a way that is different from other modulation of the sensitivity of nociceptors, such as that by the sympathetic nervous system.

**Placebo Effect**

The common way of testing treatments is to compare the results of the active treatment verses the inactive (sham) treatment when administered in similar ways.

The expectation of pain relief from what was presumed to be an effective treatment often causes a beneficial effect of the inactive treatment. This is known as the placebo effect.

Placebo effect is the beneficial effect that is obtained from administration of inactive medication or other forms of sham treatment while the participants are told that they are treated for their symptoms. It is now recognized that the placebo effect is real, at least regarding pain [43]. The placebo effect may be regarded as another way of modulating the flow of pain impulses. Many treatments have placebo effects, and it is well known that placebo treatment can reduce pain and has been recognized as a form of treatment for pain [44, 45]. The placebo effect on pain could be caused by endogenous opioids that were liberated because of the expectation of a beneficial effect from the treatment that, unknowingly to the participant in the study, was a sham treatment.

A study of postoperative patients supported that hypothesis and showed that the participants who responded positively to placebo experienced increased pain after administration of naloxone. Those who did not respond to placebo did not respond to naloxone either [45]. Perception of pain is very complex. It is not surprising that emotional factors are involved in many forms of pain, making it understandable that the observed effect of placebo treatment for pain is also complex [39, 44]. Involvement of limbic structures and descending pathways from the prefrontal cortex most likely plays important roles, both in some forms of pain perception and in the beneficial effect of treatment with placebos [9].

**Conclusion**

Pain has many different forms from minor pain that may be regarded as a nuisance to pain that has a wide range of symptoms regarding perception, degree of suffering, and how it can affect the entire life of a person. While the anatomy of acute pain is not more complicated than that of sensory systems, the anatomy of pain that causes serious suffering is complex, and it may involve large parts of the brain. The physiology of acute pain involves similar circuits as the somatosensory systems but severe pain may involve activation of circuits that can modulate pain, and it may re-route information to many different parts of the brain through activation of neural plasticity.

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Naloxone: An antidote to opiates that counteracts the pain relieving effect of opioids.
References

biological basis for mind body interaction. Prog Brain Res 122:271–85.


Keypoints

1. Tinnitus research on humans is difficult, primarily because the pathophysiology of tinnitus is still not well understood.

2. A number of animal models have been developed in order to study conditions that may lead to tinnitus and evaluate treatments for efficacy and safety before being used in human trials.

3. Current tinnitus animal models fall into five general subtypes:
   a. Lick suppression
   b. Operant conditioning
   c. False-positive models
   d. Avoidance conditioning
   e. Startle reflex models

4. Animal models have evaluated tinnitus induced primarily by:
   a. High doses of sodium salicylate
   b. High doses of quinine
   c. High-level noise exposure

5. A number of tinnitus treatments that target specific mechanisms have been proposed and tested in animal models. These include:
   a. Calcium channel antagonists
   b. GABA agonists
   c. NMDA antagonists
   d. Benzodiazepines
   e. Potassium channel modulators
   f. Transcranial magnetic stimulation

6. Tinnitus animal models provide important guidance in the development of new drug therapies.

Keywords
Animal models • Drug therapy • Startle reflex • Tinnitus

Abbreviations

- BW Bandwidth
- GABA γ-Aminobutyric acid
- GPIAS Gap prepulse inhibition of the acoustic startle
- NBN Narrow band noise
- NBPIAS Noise burst prepulse inhibition of the acoustic startle
- NMDA N-methyl-d-aspartic acid
- rTMS Repeated transcranial magnetic stimulation
- SC Scopolamine
- SIPAC Schedule induced polydipsia avoidance conditioning
- SS Sodium Salicylate

Introduction

Behavioral Models of Tinnitus

Over the last 15–20 years, a number of animal models have been developed to facilitate basic research into the biological basis of tinnitus. While the models vary from measuring reflexes to advanced conditioning paradigms, they share a basic feature: animals must discriminate between quiet and the presence of a real sound. When tinnitus is present, the ability to detect quiet becomes compromised and animals behave as if a real sound was present. The following chapter will
introduce a number of animal models and review some of the treatments that have been evaluated using these models. These advances provide the framework to accelerate preclinical and basic research toward the biological mechanisms of tinnitus and the effects of potential treatments. The reader will appreciate the ingenious and creative ways in which researchers have shown that, indeed, animals appear to experience tinnitus.

Animal Models to Assess Tinnitus

Conditioned Lick Suppression: Jastreboff developed the first behavioral model of tinnitus [1] using a conditioned, lick-suppression paradigm. Water-deprived animals were allowed to lick for water when sound was present; however, during randomly presented quiet intervals an unavoidable foot shock was administered at the end of the interval. Delivery of foot shock suppressed licking during quiet intervals, but not during sound intervals. In this conditioned-suppression paradigm, animals learned to lick when sound was present and to suppress licking during quiet. After being conditioned, animals in the experimental group were given a tinnitus-inducing agent (e.g., a high dose of sodium salicylate) while animals in the control group were given a placebo (e.g., saline). During the tinnitus testing phase, the foot shock was turned off and the lick-suppression behavior began to extinguish as foot shock was no longer presented at the end of quiet intervals. The animals in the experimental group that experienced tinnitus during the quiet intervals quickly began to lick during the silent intervals, and the conditioned lick suppression extinguished rapidly. In contrast, the control group did not experience tinnitus during the testing phase and the rate of extinction was much slower as quiet intervals continued to suppress licking, because these intervals were still associated with shock. Tinnitus was assumed to be present if lick suppression extinguished more rapidly in the experimental group (tinnitus) than in the control group. The lick-suppression paradigm was then used to assess the presence, pitch, and loudness of tinnitus induced by high doses of salicylate or quinine [2–5]. While the conditioned lick-suppression model provided useful data, it had some important limitations. First, the onset and offset of tinnitus could not be assessed repeatedly in the same animal. Instead, the technique required two groups, an experimental tinnitus group and a control group. The analysis was based on comparison of group data rather than data from individual animals. Second, the behavior extinguished after 4–5 days. Therefore, tinnitus could only be assessed over a short time interval. With this model, it was not possible to determine if tinnitus was permanent or temporary, or to measure the time course of tinnitus onset and cessation.

Heffner modified the Jastreboff-conditioned lick-suppression paradigm, so that water-deprived hamsters could avoid foot shock if they ceased licking for water during quiet intervals [6]. Hamsters were then exposed to high-intensity noise from 1 to 4 h. Following noise exposure, the foot shock was turned off and rate of extinction of lick suppression in the noise-exposed group was compared to the control group. Tinnitus assessment began 5 days postexposure, as physiological data suggested that tinnitus would begin at this time. Most of the animals exposed for 4 h at the highest intensity (127 dB SPL) extinguished more rapidly than the control animals; this was interpreted as evidence of noise-induced tinnitus. However, only a few animals exposed at lower levels for shorter durations exhibited signs of tinnitus. The results of this study were important for two reasons. First, the results indicated only high-level and long-duration exposures reliably induced tinnitus 5–9 days postexposure, while low-level and short-duration exposures seldom induced tinnitus. Second, only a subset of hamsters developed tinnitus as expected from human reports of noise-induced tinnitus. Heffner’s method, however, has some of the same limitations as the Jastreboff paradigm: (a) a separate control group is needed to infer if tinnitus is present; (b) the behavior extinguishes in 4–5 days making it difficult to determine if tinnitus is permanent or temporary; (c) it is not possible to determine the time course of tinnitus and whether the tinnitus is temporary or permanent.

Operant Conditioning: Bauer and colleagues developed a tinnitus animal model by training food-deprived rats to press a bar for food in the presence of white noise and to stop responding during quiet intervals paired with foot shock [7]. At random intervals, a test tone was substituted for white noise without shock. When the stimuli were test tones, the group of salicylate-treated rats continued to press for food more often than the control group. The explanation for this behavioral difference was that salicylate-treated rats perceived the
tones as “noisy” due to the presence of tinnitus and, therefore, suppressed bar pressing behavior less than controls. This approach allowed for long-term assessment of tinnitus; however, a limitation of this technique was that differences in behavior attributed to tinnitus occurred only at elevated sound levels, often at low frequencies, contrary to tinnitus reported in humans. Thus, results could be more reflective of changes to suprathreshold hearing induced by salicylate, such as changes in sound tolerance, rather than tinnitus. This technique continues to be refined and the authors report that rats with tinnitus have a constant noise floor and require a larger signal-to-noise ratio than normal control in order to hear tones above their tinnitus.

False-Positive Response Models: Guitton used an increase in “false-positive” responses in quiet to infer the presence of tinnitus [8]. Rats were trained to jump onto a pole when sounds were presented in order to avoid foot shock. During quiet intervals, the shock was turned off and animals could safely remain on the cage floor. In the training phase, rats reliably climbed the pole during sound intervals (hits) and seldom jumped on the pole during quiet intervals (false positive). Rats were then treated for 4 days with 300 mg/kg/d of sodium salicylate. Salicylate-treated rats showed a progressive increase in false-positive responses during quiet intervals over the 4 days of salicylate treatment, indicative of tinnitus. After salicylate treatment ended, the false-positive rate began to decline and reached control levels 2 days postsalicylate, indicating cessation of tinnitus. This behavioral paradigm has several appealing features. It does not extinguish, it does not require a separate control group, and it can be used to assess the onset and recovery of tinnitus. There are, however, some potential limitations with this technique. First, once an animal develops tinnitus, there would no longer be any safe periods in the test chamber. In other words, the animal would need to jump on the pole and remain there 100% of the time, i.e., when the noise was present and when tinnitus was present. Consequently, only a few trials could be run per test session. Second, if an animal developed permanent tinnitus, then it would always jump on the pole, making it difficult to distinguish between tinnitus-induced false positives and false positives due to lack of stimulus-controlled behavior. Finally, over time, an animal with permanent tinnitus might learn to distinguish a phantom sound from a real sound and no longer climb on the pole during quiet periods. Data from other animal models, including the authors of this chapter, suggest that over time some animals learn to discriminate their tinnitus from real sound, or that a low-level tinnitus becomes the animals’ “quiet” state.

Ruttiger combined food-reinforced operant conditioning with a false-positive model. His group trained rats to activate a liquid feeder when white noise was presented, and to withhold their response during quiet periods when no food was delivered [9]. After rats were treated with 350 mg/kg of salicylate, they increased their “false-positive” response rate during the quiet intervals, suggesting that they were perceiving tinnitus. The intensity of the phantom sound was estimated to be around 30 dB SPL since the false-positive rate to the 350 mg/kg dose of salicylate was similar to the number of response evoked by a real noise of 30 dB SPL. Lowering the dose of salicylate reduced the false-positive rate in periods of quiet. The main limitation of this technique is the large amount of time needed to train the animals. Other limitations include the ability to detect the frequency of the tinnitus, whether animals can learn to discriminate tinnitus from real sound over time, and extinction of the ability of quiet to reduce responding. Despite these limitations, this method can reliably detect the presence, intensity, and persistence of tinnitus.

Schedule Induced Polydipsia Avoidance Conditioning (SIPAC): The authors of this chapter developed SIPAC to evaluate the onset, offset, and pitch of transient or persistent tinnitus. Under SIPAC, food-restricted animals (85% free feeding weight) are placed under a fixed, 1-min time interval (FT = 1) food reinforcement schedule while water is available in the experimental chamber (animals receive one pellet per minute). Each daily session is 150 min, under which animals receive 150, 45-mg food pellets. Over a few days, animals become polydipsic; they begin to exhibit large bursts of drinking following food pellet delivery. Typically, total session licks for each session range from 2,000 to as high as 10,000 licks. Initially, a 4-kHz narrow band noise (NBN) is played in the background on half of the trials while the other half has no sound (quiet). Once the animals become polydipsic, a brief foot shock is delivered if animals lick in the presence of the 4-kHz NBN. Within 2–3 days animals restrict their licking in the presence of sound and lick predominantly during quiet intervals (<10% licks in the presence of sound). In the final training stage, the sound trials are generalized to 4, 8, 12, 16, and 20 kHz NBN or 16 kHz
tone. Licking in the presence of these sounds also results in brief foot shock. However, no shock is delivered if the animals lick during the quiet intervals. The presence of tinnitus is inferred by a decrease in the licks that occurs during the quiet intervals, while sound intervals are expected to remain unchanged. Recent advancements in the SIPAC technique have allowed the estimation of the pitch of salicylate, quinine, and unilateral noise-induced tinnitus (only one ear is exposed to the noise, the other ear is left unexposed to hear the real sound). The pitch was estimated to be in the 12–16 kHz range for salicylate and quinine-induced tinnitus and 12–20 kHz for noise-induced tinnitus, depending on the frequency of the noise trauma. These ranges are consistent with previously published results using other animal models.

**Gap Prepulse Inhibition Acoustic Startle (GPIAS):** Turner [10] developed an efficient technique to assess tinnitus, which we refer to as GPIAS. The dependent measure in GPIAS is the amplitude of a sound-evoked reflex in response to an acoustic startle stimulus (115 dB noise burst, 20 ms). The acoustic startle reflex is a rapid extension and reflection of a series of muscles resulting in pressure exerted on a platform. The changes in pressure are detected by a piezo transducer attached to the bottom of the platform. Presentation of the startle stimulus reliably induces a robust startle response. However, the amplitude of the response to the startle can be suppressed when a low-level stimulus, or pre-pulse, precedes the startle stimulus. Similarly, a detectable silent gap embedded in an otherwise continuous low-level background noise presented before the startle stimulus can also suppress the startle reflex. Suppression of the startle reflex by a prepulse or gap is referred to as prepulse, or gap prepulse, inhibition as shown on Fig. 16.1. In the Turner study, the gap prepulse stimulus was a 50-ms silent interval embedded in otherwise continuous noise (60 dB SPL, broad band noise, or narrow band noise centered at 10 or 16 kHz); the gap preceded the startle stimulus by 100 ms. In untreated rats, the gap prepulse suppressed the startle response by 50–65%, relative to trials with no gaps. However, in unilaterally noise-exposed rats believed to have tinnitus, prepulse inhibition was normal except for gaps embedded in noise centered at 10 kHz. The authors concluded that the rats had 10 kHz tinnitus that partially filled in the silent gap embedded in the narrow band noise and reduced the ability of the rats to detect gaps in 10 kHz noise. In contrast, gap prepulse inhibition was normal for gaps embedded in broadband noise or narrowband noise centered at 16 kHz, presumably because rats could differentiate the 10 kHz tinnitus from the broadband noise or 16 kHz narrow band noise. More importantly, the same animals that showed impaired GPIAS at 10 kHz also showed evidence of 10 kHz tinnitus measured with an operant bar press discrimination task [11].

**Methods of Inducing Tinnitus**

**Salicylate-Induced Tinnitus:** In our first experiments, we used SIPAC to determine which dose of salicylate would reliably induce tinnitus [12] and to find out if these results were consistent with previous reports. Figure 16.2 shows the typical behavior of a rat with salicylate-induced tinnitus. During baseline testing, the rat made 2,000–4,000 licks in quiet (>90% correct) and almost no licks in sound (40 dB SPL). Following baseline measures, the rat was injected with saline for 2 days. Licks in quiet remained high (correct) while licks in sound remained low, indicating that the injections had no adverse effects on performance. Next, the animal was injected with 150 mg/kg of salicylate for...
two consecutive days. On these days, licks in quiet were far below a 99.9% confidence interval established during baseline/saline conditions, providing a statistical method for detecting the presence of tinnitus in an individual animal. The licks in quiet remained low during the first 2 days of recovery, indicating residual tinnitus. However, the ratio then returned to baseline levels, indicating the absence of tinnitus. A dose of 50 mg/kg of salicylate was also administered to the rat (not shown). This dose failed to suppress licks in quiet, indicating that the treatment was too low to induce tinnitus-like behavior. Finally, the animal was treated with 100 mg/kg of salicylate (not shown), which partially suppressed licks in quiet on the first day, but the effect disappeared by the second day. This result suggested that the 100 mg/kg does not reliably induce tinnitus.

Salicylate Dose–Response: The mean \( n = 5 \) salicylate dose–response data are shown in Fig. 16.3. Licks in sound remained low during the entire experiment, indicating that the response was under stimulus control and real sound remained audible. Licks in quiet during saline treatment and the 50 mg/kg dose of salicylate were high, similar to baseline, indicating an absence of tinnitus. In contrast, licks in quiet were significantly reduced during the 150 mg/kg and 350 mg/kg doses of salicylate, indicating the presence of tinnitus. Licks in quiet were slightly reduced with the 100 mg/kg dose, but the reduction was not statistically significant, indicating an absence of tinnitus. To study the recovery from salicylate-induced tinnitus, we measured licks in quiet after salicylate treatment ended. Licks in quiet were greatly depressed during treatment with 150 mg/kg, remained low 1–2 days posttreatment, and fully recovered to baseline values by the third day.

Quinine-Induced Tinnitus: Jastreboff was the first to report evidence of tinnitus in animals treated with high doses of quinine [4]. Quinine, an antimalarial agent, and its derivatives are still used in sub-Saharan Africa and in military populations serving overseas. High doses of quinine can be quite toxic and have been known to cause birth defects in humans [13]. In addition, high doses of quinine have been reported to induce tinnitus [14, 15]. The mechanism of action by which quinine induces tinnitus is not well understood, but may be related to effects on calcium channel signaling and hyperactivity. We evaluated the effects of quinine using SIPAC and GPIAS to determine the dose, duration, and pitch characteristics...
of quinine-induced tinnitus. Figure 16.4 shows the effects of quinine on SIPAC. At 100–150 mg/kg, quinine significantly reduced licks in quiet, indicating the presence of tinnitus. We repeated the experiment using GPIAS, and again found a dose-dependent reduction in the ability to detect silent gaps. The effects on GPIAS are shown on Fig. 16.5. Note that the pitch of quinine-induced tinnitus was 16–20 kHz.

Salicylate Dose Response Curve and Tinnitus assessed by SIPAC (n=6)

Fig. 16.3 Salicylate dose–response function. Animals drink during quiet intervals with no shock and refrain from drinking during noise intervals that are paired with shock. When treated with saline or a low dose of salicylate (50 mg/kg), animals can still discriminate quiet from sound intervals. When the dose of salicylate exceeds 100 mg/kg, animals behave during quiet intervals as if a sound was there, indicated by a decrease in licks in quiet. These results are consistent with the presence of tinnitus. Note that behavior to the real sound intervals remains unchanged.

Quinine Dose Response Curve and Tinnitus assessed by SIPAC (n=6)

Fig. 16.4 Quinine dose–response function. Animals drink during quiet intervals with no shock and refrain from drinking during noise intervals that are paired with shock. When treated with saline or a low dose of quinine (50 mg/kg), animals can still discriminate quiet from sound intervals. When the dose of quinine is increased to 100–150 mg/kg, animals behave during quiet intervals as if a sound was there, indicated by a decrease in licks in quiet. These results are consistent with the presence of tinnitus. Note that behavior to the real sound intervals remains unchanged.

Effects of 150 mg/kg Quinine (PO) on GPIAS as a Function of Frequency (n=6)

Fig. 16.5 Effects of 150 mg/kg quinine on GPIAS. Compared to baseline measures, there is a progressive decrease in GPIAS at 12–20 kHz. These results indicate that animals cannot reliably detect silent gaps embedded in NBN 12–20 kHz, suggesting the presence of tinnitus. Note that behavior gaps in lower frequency NBN (6 kHz) remains unchanged.
Noise-Induced Tinnitus: We first used SIPAC to determine if individual rats developed tinnitus after unilateral exposure to narrow band noise centered at 11 kHz (120 dB, 2 h). The exposure resulted in a 35–50 dB high-frequency hearing loss (confirmed by auditory brainstem response) immediately after the exposure, but hearing remained nearly normal in the contralateral, unexposed ear. As shown in Fig. 16.6, some rats developed persistent tinnitus (open downward triangles). Note a large decrease in the number of licks in quiet from baseline (B1–5) after noise exposure (N1d–N3d), indicating the presence of tinnitus. Other rats developed transient tinnitus for a day or two (Fig. 16.7, licks in quiet, open downward triangles), and then recovered to baseline by postnoise day 3. Some rats, however, failed to develop noise-induced tinnitus (not shown), consistent with other studies [6]. These results are in agreement with human studies showing that some individuals develop noise-induced tinnitus while others do not (see Chap. 37). These data illustrate the importance of assessing tinnitus in individual rats rather than solely assessing group data.

In addition to SIPAC, we recently evaluated the effect of unilateral noise-induced hearing loss on the development of tinnitus under GPIAS. Like previous studies, animals show a range of variability in their susceptibility to noise-induced hearing loss and noise-induced tinnitus. However, we have found that the maximum hearing loss and tinnitus pitch resulting from the unilateral noise trauma is often ½–1 octave above the noise exposure. For instance, when one group of animals was exposed to 12 kHz bandpass noise (123 dB SPL) unilateral noise trauma, GPIAS was reduced initially across multiple frequencies as shown in Fig. 16.8 (2 h postnoise). However, recovery typically occurred at frequencies both below the noise trauma and greater than one octave above the noise exposure.
trauma and by day 4 postnoise, there was complete recovery (Fig. 16.9). Increasing the frequency of the noise trauma to 16 kHz narrow band noise (120 dB SPL, 1 h) shifted evidence of tinnitus to 20 kHz (Fig. 16.10); results are consistent with the maximum hearing loss in the exposed ear.

In order to ensure that the tinnitus effects observed were not the result of hearing loss, we ran noise burst prepulse inhibition (NBPIAS). Under NBPIAS, a brief 60 dB SPL narrow band prepulse (50–100 ms, 6, 12, 16, 20, and 24 kHz) was presented in a quiet background 100 ms before the startle stimulus. Similar to GPIAS, if the animal detects the prepulse preceding the startle stimulus, the acoustic startle reflex will be greatly reduced compared to conditions in which no prepulse is present, resulting in a high percentage of NBPIAS. We found that unilaterally exposed animals could easily perform this task and had robust NBPIAS, indicating that the animals were able to detect the 60 dB prepulse tone. In contrast, GPIAS was reduced when gaps were inserted into continuous NBN of the same frequencies and intensities as tested by NBPIAS. To further confirm the status of the unexposed ear and exposed ears, distortion product otoacoustic emissions (DPOAE) were obtained from both the traumatized and nontraumatized ears. We found that the nonexposed ear did not differ from baseline measures, while the traumatized ear showed large reductions or absent DPOAE, consistent with the effects of noise on outer hair cell function in the noise-exposed ear. The results from GPIAS, NBPIAS, and DPOAE were consistent with the presence of tinnitus in the 12–20 kHz frequency region, and not the result of hearing loss.

**Tinnitus Treatments**

A number of potential tinnitus suppressing drugs have been evaluated in animal models. The following section presents an overview of drugs from different classes that have been suggested or hypothesized as potential treatments for tinnitus (see also Chaps. 78 and 79).

**Nimodipine**: This calcium channel antagonist was reported to block quinine-induced tinnitus in a dose-dependent manner [4] in rats. A subsequent human trial found evidence that a small subset of patients showed a reduction of tinnitus severity with nimodipine treatment; however, this study lacked an appropriate placebo control [16], and more research is needed on Nimodipine to determine whether it is a viable treatment for some forms of chronic tinnitus.

**Tiagabine and Gabapentin**: The efficacy of tiagabine (GABA agonist that inhibits GABA reuptake) and gabapentin (anticonvulsant, antihyperalgesic, antinoceptive, mechanism of action unknown) was evaluated
in rats with persistent noise-induced tinnitus [11]. Tiagabine was found to have no effect on the behavioral manifestations of tinnitus, whereas gabapentin, at clinically relevant doses (1 and 2.5 mg/kg), significantly reduced tinnitus. However, a recent human clinical trial failed to show that gabapentin was effective in treating tinnitus [17]; while another found that gabapentin reduced the annoyance of tinnitus in a subgroup of patients [18].

*The Excitatory Neurotransmitter Glutamate:* It has been hypothesized that salicylate-induced tinnitus arises because salicylate increases neural excitation by enhancing glutamate-sensitive NMDA (N-methyl-d-aspartic acid) receptor currents in the cochlea [8, 19, 20]. To test this hypothesis, rats were given intraperitoneal injections of salicylate to induce tinnitus. Salicylate-induced tinnitus was suppressed when NMDA receptor antagonists (MK801, 7-chlorokynurenate) were applied to the round window of both cochleas [8, 21]. However, a recent human clinical trial found that the NMDA antagonist flupirtine was ineffective in treating tinnitus.

*Vigabatrin:* Previous studies have suggested that loss of tonic GABA-mediated inhibition could lead to increased neural excitability and tinnitus [22, 23]. To test this hypothesis, unilateral acoustic overstimulation was used to induce tinnitus in rats tested with an operant-conditioned suppression method [22]. The psychophysical discrimination function (suppression ratio vs. intensity) of noise-exposed rats showed evidence of tinnitus-like behavior at suprathreshold levels of the 20 kHz discrimination function compared to the control group. Treatment with low and high doses of vigabatrin (30 or 81 mg/kg) abolished the tinnitus-like behavior. However, tinnitus-like behavior re-emerged during the drug washout period. These results suggested that upregulating GABA-mediated inhibition may suppress noise-induced tinnitus. However, since only group data were presented, it is not clear if vigabatrin can suppress tinnitus in all animals or just some. In addition, since tinnitus assessment was based on discrimination performance obtained at suprathreshold levels, it was not clear whether vigabatrin could abolish tinnitus-like behavior measured in quiet as is done with other techniques (e.g., SIPAC). One concern with using vigabatrin clinically is its serious side effects (e.g., ataxia, psychotic episodes) [24]. These risk factors, unfortunately, reduce vigabatrin’s therapeutic utility for tinnitus treatment.

*Memantine:* Memantine, a spasmylytic drug and NMDA antagonist with antiglutamatergic properties, has been suggested as a possible drug to treat tinnitus [25, 26]. We evaluated the ability of memantine to suppress salicylate-induced tinnitus using SIPAC. In control experiments, we first identified the dose of memantine alone (≤3 mg/kg/d) that did not disrupt behavior. We then induced tinnitus with 200 mg/kg of sodium salicylate (SS), and then administered memantine (M) at 3 mg/kg to see if it would suppress salicylate-induced tinnitus. Treatment with memantine failed to abolish salicylate-induced tinnitus (Fig. 16.11).

*Scopolamine:* Recently, the anti-cholinergic drug scopolamine (SC) has been suggested as a possible drug to treat tinnitus [26]. We first determined the dose of scopolamine alone (≤1 mg/kg/d) that did not disrupt the behavioral response. Then, we induced tinnitus with 150 mg/kg of sodium salicylate (SS). SS alone caused a significant decrease of licks in quiet, indicative of tinnitus. SC alone had no effect on licks in quiet. Afterwards, rats were treated with SS plus SC. SC caused a slight increase in the number of licks, but did not suppress salicylate-induced tinnitus [27].

![SIPAC: Effects of Memantine on Salicylate Induced Tinnitus](image)
Alprazolam and L838417: If tinnitus is the result of loss of central inhibition in response to peripheral hearing loss, it is possible that increasing endogenous inhibition might suppress the aberrant signals of tinnitus. To test this hypothesis, we evaluated the benzodiazipine, alprazolam, and the GABA-A agonist L838417 on salicylate or noise-induced tinnitus using GPIAS. Alprazolam, 0.5 mg/kg, failed to reduce evidence of 16–20 kHz tinnitus in animals treated with 250 mg/kg salicylate. Furthermore, higher doses of alprazolam (>1 mg/kg) exhibited strong sedation effects and attenuated overall startle response amplitudes.

A separate group of animals was exposed to 120 dB SPL unilateral noise trauma with a 16-kHz narrow band noise (BW = 100 Hz). Following noise trauma, animals were tracked for 15 days. Animals showing evidence of 16–20 kHz tinnitus were treated with 10 mg/kg of L838417 at 2 and 15 days posttrauma. L838417 has similar anxiolytic effects as the benzodiazepine, or Chlordiazepoxide, but does not exhibit sedative effects, making it an ideal candidate for the GPIAS paradigm. Initial results showed a partial reversal of noise-induced tinnitus at 48 h and no effect on noise-induced tinnitus at 15 days. These results suggested that early administration of L838417 treatment following noise trauma may reduce the presence of tinnitus temporarily or inhibit the development of permanent tinnitus. However, after 15 days of administration, the drug has little effect on tinnitus. Interestingly, new hypotheses are emerging regarding potential differences between the acute phase and the chronic phase of tinnitus, in regard to their neural locus. Early tinnitus is believed to have a more peripheral component and reside subcortically, while chronic tinnitus may result from more central plasticity changes. These differences could explain some of the results obtained among studies.

Potassium Channels and Tinnitus: Potassium channel dysfunction has been implicated in brain hyperactive disorders such as epilepsy [28, 29]. One of the functions of potassium channels is to regulate the resting state of neurons and the depolarization threshold. The emergence of highly selective potassium channel modulators has brought forth new potential therapies for central hyperactivity. We have begun to evaluate potassium channel modulators and their effect on salicylate or noise-induced tinnitus. Preliminary data show that BK potassium channel agonists reverse the effect of salicylate-induced tinnitus, and may slightly reduce evidence of noise-induced tinnitus. However, further validation is needed to determine if these effects can be replicated and if the effects on salicylate are a drug-on-drug interaction or a reduction of the tinnitus percept.

Repeated Transcranial Magnetic Stimulation (rTMS): Transcranial magnetic stimulation is an approved therapeutic technique for the treatment of depression [30, 31]. It has also been evaluated in humans for the treatment of tinnitus [32–35]. One hypothesis of rTMS for the treatment of tinnitus is that magnetic pulse stimulation may “break up” synchronous activity in the brain associated with the perception of tinnitus.

Fig. 16.12 Animals with evidence of tinnitus at 12–16 kHz were treated to two sessions (one ipsi, one contra) of repeated Transcranial Magnetic Stimulation (rTMS). The animals were awake and restrained and treated with 100 pulses per session. Results suggest a partial reversal of tinnitus, indicated by an increase in GPIAS.

Fig. 16.13 A single animal with evidence of tinnitus at 12–16 kHz was treated to two sessions (one ipsi, one contra) of repeated Transcranial Magnetic Stimulation (rTMS). The animal was awake and restrained and treated with 100 pulses per session. Results suggest no benefit from rTMS in either the ipsilateral or the contralateral condition.
Although the use of rTMS for tinnitus is still in its early stages, there have been some positive results in a select number of tinnitus patients (see Chap. 88). When we evaluated rTMS on rats ($n=4$) with noise-induced tinnitus, we found transient suppression after a single rTMS session (100 pulses) in two of the rats (Fig. 16.12), no change in one rat, and increased tinnitus in one rat (Fig. 16.13). The results suggested that there might be a biological basis for the efficacy of rTMS for at least some tinnitus patients.

**Current Studies**

Advances in our understanding of neural disorders and the development of novel pharmacological agents to manage central disorders provide newfound opportunities for potential tinnitus treatments. Engineered compounds and nanoparticles allow more efficient and effective targeting of specific receptors and channels. A combined preclinical approach among physiological, microbiological, and behavioral studies of tinnitus is likely to accelerate the finding of new potential treatments to manage or eradicate tinnitus. New human clinical trials with drugs that modulate NMDA activity are ongoing and show promising results on noise-induced tinnitus.

**Summary**

While no treatments are available that are effective in all tinnitus patients, a number of significant advancements have been made in tinnitus research. Physiological and functional brain imaging studies have suggested that increased activity along the central auditory pathway is present during tinnitus. Theories of tinnitus have moved away from the inner ear to a central generator for chronic tinnitus. Clinical work is ongoing for better classification of tinnitus subtypes, better patient selection is being implemented for human drug studies, and a number of animal models have been developed. This last point is of critical importance as tinnitus can currently only be assessed from a person’s or an animal’s subjective experience. However, the animal models allow researchers to find biological correlates of behavioral evidence of tinnitus and explore how tinnitus develops from trauma to the inner ear. Additionally, while the locus may remain unclear, preclinical drug trials can be performed to determine if potential treatments reduce behavioral evidence of tinnitus in animals. These advancements have come a long way in establishing tools necessary to reveal the biological markers of tinnitus and to develop effective treatment strategies.

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**References**


Chapter 17
Objective Signs of Tinnitus in Humans

Bertold Langguth and Dirk De Ridder

Keypoints
1. Different methods have successfully been used for detecting tinnitus-related changes in the brain; chief among them are neuroimaging, electroencephalography and magnetoencephalography.
2. These methods make it possible to detect noninvasively neuronal activity in the human brain and determine the anatomical location of the activity.
3. Findings from neuroimaging have already contributed to a better understanding of the pathophysiological changes underlying the different forms of tinnitus.
4. The different neuroimaging methods hold the potential to be further developed as methods for diagnosis, outcome assessment, and outcome prediction.
5. Replication of studies with larger sample sizes and clinically well-characterized individuals with tinnitus is needed.

Keywords Tinnitus • Neuroimaging • Electroencephalography • Magnetoencephalography • Functional magnetic resonance tomography • Positron emission tomography • Diagnosis • Pathophysiology

Abbreviations
EEG Electroencephalography
fMRI Functional magnetic resonance imaging
MEG Magnetoencephalography
MRI Magnetic resonance imaging
PET Positron emission tomography
SPECT Single positron emission computed tomography

Introduction

Many forms of tinnitus are phantom perceptions of sound and therefore related to functional changes in the brain. Identification of the “neuronal correlate” of such forms of tinnitus is of utmost importance for a deeper understanding of the pathophysiology and the development of new effective treatments for these kinds of tinnitus. It should be stressed that tinnitus is most likely a network property and that the “neural correlate” should be understood as such, and should not be viewed as one phrenological “tinnitus hotspot” somewhere in the brain. Several different methods have been increasingly used during the last two decades for the detection of tinnitus-related changes in the brain and in attempts to find where in the brain the physiologically abnormal neural activity is generated and to which extent the pathophysiological changes in humans correspond to those in animal models of tinnitus. In detail, these methods are structural and functional neuroimaging methods and source-localized electroencephalography (EEG) and magnetoencephalography (MEG). These methods make it possible to detect neuronal activity noninvasively in the human brain and determine the anatomical location of the activity. Even the most cautious interpretation of the available data, most of them come from small samples, indicates the potential of neuroimaging, EEG, and MEG as valuable tools in tinnitus research. These methods provide windows to the brain that allow detecting the localization of
tinnitus-related changes in the brain. This knowledge is indispensable for a better understanding of the pathophysiology of tinnitus (see Chap. 21). Very importantly, imaging techniques can be applied both in animals and in humans and can so contribute to bridge the gap between the knowledge coming from clinical data and animal models of tinnitus [1].

Neuroimaging may not only serve as a tool for improved understanding of the pathophysiology but also have an impact on future diagnosis and treatment of tinnitus patients. This can be best illustrated by the recent development of brain stimulation techniques for the treatment of tinnitus (see Chaps. 88 and 90). Neuroimaging findings of increased neural activity in the auditory cortex of tinnitus patients prompted the suggestion to treat tinnitus by focal modulation of this activity with electric or magnetic stimulation. Neuroimaging has the potential to be further developed as an objective diagnostic tool for tinnitus. Most findings from studies of tinnitus-related brain changes come from comparison of groups of individuals with tinnitus with matched groups of individuals who do not have tinnitus. The results of such studies do not automatically mean that each individual with tinnitus has an identical abnormality as the ones detected when groups of individuals are compared. Further studies with larger sample sizes will be needed to estimate sensitivity and specify of different techniques for the diagnosis of tinnitus, since there is not yet enough evidence that any of the presented methods can be recommended for use in routine diagnostic management of tinnitus patients.

A further potential application of neuroimaging is for distinguishing between different forms of tinnitus. It may be assumed that differences in the perceptual characteristics of tinnitus, in the emotions surrounding tinnitus and in the response to specific treatments, would be reflected by specific patterns of neural activity, which could be detected by the use of imaging techniques. By contributing to this differential diagnosis, imaging may in the future also serve as predictor of the efficacy of specific treatments and for the assessment of treatment outcome. This will help to exactly identify the neuronal mechanisms by which specific treatment interventions exert their effects. This knowledge in turn can be useful for improving efficacy of those treatment interventions.

Whereas EEG and MEG measure directly the electrical and magnetic field, which is induced by neuronal activity, “functional imaging” methods such as functional Magnetic Resonance Imaging (fMRI) or Positron Emission Tomography (PET) measure changes in cerebral blood flow, blood oxygenation, and glucose uptake based on the assumption that alterations of neuronal activity are reflected by changes in the hemodynamic or metabolic responses. Results of the use of these methods in the investigation of tinnitus and the results from such studies are summarized in the following chapters.

Tinnitus-related functional changes of neural activity have been investigated with fMRI, PET, and Single Positron Emission Computed Tomography (SPECT). The different methods differ in the correlates of neuronal activity they detect (e.g., cerebral blood flow or glucose uptake) and in their ability to measure resting neuronal activity or stimulus-evoked changes of neuronal activity. Results from the use of these methods for the study of tinnitus will be presented in detail in Chap. 18. High-resolution Magnetic Resonance Imaging (MRI) data have demonstrated changes in the volume of specific brain structures in tinnitus patients. However, it remains to be elucidated, whether these alterations are a consequence of longer lasting changes in functional activity or whether they rather represent a marker for increased vulnerability to develop tinnitus. This will be discussed further in Chap. 19.

Chapter 20 concerns electrophysiologic methods for studies of neural activity. While the EEG records the electrical field, which is produced by neuronal activity, MEG records the magnetic field changes. The use of MEG and EEG is based on the assumption that electrical activity either from electrodes placed on the scalp (EEG) or from measurement of the small changes in the magnetic field that can be measured outside head (MEG) correlates with neural activity in populations of nerve cells. Typically, EEG is recorded by many electrodes placed over the surface of the scalp. Signals recorded by these electrodes can be used to construct a map of the brain’s electrical activity. Both EEG and MEG are characterized by high temporal but low spatial resolution. MEG is more sensitive for currents that are directed tangential to the surface of the skull, whereas EEG detects radial sources best. Both methods...
have the advantage that they do not produce noise that can interfere with auditory recordings as the imaging methods do. EEG and MEG can be used both for measuring resting brain activity and for recording neural activity elicited by sound.

Reference

Keypoints

1. Different functional imaging methods, such as SPECT, PET, and fMRI, have been used for investigating tinnitus.
2. Neuroimaging methods have provided windows to the brain that allow detection of the localization of tinnitus-related changes in the brain.
3. Such studies have shown signs of abnormalities in many parts of the brain, including auditory brain regions but also nonauditory brain areas involved in sensory integration, in attention, or in emotional evaluation.
4. New treatment strategies have evolved from fMRI and PET findings of abnormal neuronal activity in the auditory cortex.

Keywords

Tinnitus • Neuroimaging • Electroencephalography • Magnetoencephalography • Functional magnetic resonance tomography • Positron emission tomography • Diagnosis • Pathophysiology

Abbreviations

FDG-PET Fluor-deoxy-glucose PET
fMRI Functional Magnetic Resonance Imaging
PET Positron emission tomography
rCBF Regional cerebral blood flow
SPECT Single positron emission computed tomography
[15O]-H2O PET Positron emission tomography with radioactively labeled water

Introduction

Tinnitus-related functional changes of neural activity have been investigated with functional imaging techniques such as single photon emission computed tomography (SPECT), positron emission tomography (PET), and functional magnetic resonance imaging (fMRI). The different methods and the results of their use in the investigation of subjects with tinnitus will be presented in detail in the following sections. Finally, we will discuss how the different techniques have contributed to identify the anatomical location of the functional abnormalities that cause some forms of tinnitus.

Single Photon Emission Computed Tomography (SPECT)

SPECT (single photon emission computed tomography) scanning makes use of a radioactive tracer emitting gamma rays to measure blood flow in regions of the brain (regional cerebral blood flow, rCBF). The emission of photons is recorded by a camera that provides a 3D image of the anatomical location of indicators of neural activity. To obtain a SPECT scan, the individual person receives an injection of a small amount of a radio-labeled compound, e.g., Technetium-HMPAO. The distribution of this compound is related to blood flow and is used as a measure for local neural activity.

A study of rCBF using SPECT [1] in 45 depressed individuals, of whom 27 had severe tinnitus, found decreased CBF in the right frontal lobe Brodmann area 45 (Broca, pars triangularis), the left parietal lobe.
area 39 (angular gyrus, part of Wernicka’s area), and the left visual association cortex area 18 (secondary visual cortex, V2) in tinnitus patients compared with nontinnitus patients. In patients with tinnitus, the CBF was increased in the primary, secondary, and auditory association areas of the temporal lobe (Brodmann’s area 41, primary auditory cortex; area 21, middle temporal gyrus; area 22, superior temporal gyrus) as compared to gender-matched controls and depressed patients who did not have tinnitus. The study also showed signs that the superior temporal gyrus bilaterally (primary and secondary auditory cortex) and three further brain areas were more active in depressed patients who had tinnitus than in depressed patients without tinnitus: Brodmann areas 18 (V2), 39 (inferior parietal, angular gyrus), and 45 (Broca’s homologue, VLPFC) [1]. Another study in two individuals who had tinnitus found differences in the temporal, frontal, parietal, hippocampal, and amygdala regions when compared with normative Te-HMPAO SPECT data [2].

**Positron Emission Tomography**

PET has a similarity to SPECT and makes use of a radioactive tracer (a short-lived radioactive isotope) to identify the anatomical location of indicators of neural activity such as blood flow or glucose metabolism.

As the radioactive atoms in the compound decay, they release positively charged positrons. When a positron collides with a negatively charged electron, they are both annihilated and two photons are emitted. The photons move in opposite directions and are detected by the sensor ring of the PET scanner. Reconstruction of the three-dimensional paths of the particles provides information about the maximum accumulation or metabolism of the short-lasting radio-labeled isotope at a higher resolution than obtained with a SPECT scan (1 cm for SPECT).

PET retains unique advantages in studies of auditory processing over fMRI because it is not associated with any noticeable noise, as are fMRI machines, which produce up to 130 dB noise at the location where the person who is scanned is placed. Unlike fMRI, PET can be used to study individuals with cochlear implants or other kinds of implanted electrodes, which do not allow use of fMRI [3]. PET is also much less sensitive to body movements, such as those from arterial pulsations in the brainstem. On the other hand, PET is not widely available, is relatively expensive, and is always associated with exposure to ionized radiation, which precludes repetitive imaging sessions.

The two PET methods have been used in the investigation of tinnitus: one uses a radioactively labeled glucose (FDG PET), which reflects metabolic activity, and the other type uses radioactively labeled water ([¹⁵O]-H₂O PET), which provides a measure for cerebral blood flow.

**Studies Using [¹⁵O]-H₂O PET**

Estimates of changes in rCBF using PET with radioactively labeled water ([¹⁵O]-H₂O PET) have been used as an indicator of changes in neural activity during transient reduction of tinnitus loudness, e.g., by the administration of lidocaine [4, 5].

Several PET studies of rCRB took advantage of the fact that few individuals can modulate their tinnitus by orofacial movements [6, 7] or eye movements, a condition which may occur after surgical operations in the cerebellopontine angle [8]. Thus, Giraud et al. [9] found that such forms of tinnitus are associated with an increase in CBF bilaterally, especially in auditory temporoparietal association areas.

Individuals with unilateral tinnitus who could alter the loudness of their tinnitus by orofacial movements showed indications that neural activity in areas adjacent to the **contralateral** auditory cortex increased and decreased in parallel to the reported change in loudness of tinnitus [10]. In contrast, auditory stimulation in the same individuals resulted in **bilateral** activation of the auditory cortex, suggesting that the abnormal neural activity that caused the sensation of tinnitus originated in the central auditory system rather than the cochlea [10]. When investigating subjects with gaze-evoked tinnitus, Lockwood et al. [11] found signs of CBF alterations in a large part of the frontal, parietal, and temporal cortex, as well as the lateral pontine tegmentum and the primary auditory cortex. Whereas lateral gaze reduced rCBF in the temporal lobe in control subjects, this was not the case in individuals with tinnitus whose condition worsened during lateral
gaze. This finding suggests that gaze-evoked tinnitus may be caused by reduced gaze-evoked inhibition of the auditory cortex [11].

Tinnitus was also associated with more widespread activation of neural structures in the brain not activated by sound stimulation, including activation of limbic structures, which indicated that plastic changes of the auditory nervous system had occurred (see Chap. 12).

Several investigators using PET scans have shown indications that intravenous administration of lidocaine can modulate tinnitus [4, 5, 12, 13]. Most studies of the effect of lidocaine on tinnitus have been done in individuals where lidocaine decreased the loudness of the tinnitus. In such a study, Reyes et al. [5] found that the decrease in the loudness of the tinnitus was associated with changes in the neural activity in the right auditory association cortex. These findings were confirmed and extended by Plewnia et al. [4], who found changes in CBF in a broad region of the auditory cortex (middle temporal gyrus), including areas involved in the integration of sensory stimuli (gyrus angularis) and cognitive processing (posterior cingulated cortex) of sensory stimuli.

In a recent study, Andersson et al. [14] showed evidence that reduction of tinnitus loudness during a cognitive task (silent backward counting) is accompanied by reduced CBF in auditory cortex.

Taken together, measurements of rCBF with [15O]-H2O PET have consistently provided evidence for tinnitus-related increases of neural activity in auditory pathways as well as in some nonauditory neural systems. However, the use of this technique depends on the ability to influence the loudness of the tinnitus by specific interventions, which means that it can only be used in individuals who can modulate their tinnitus.

**Studies Using FDG-PET**

Another, and perhaps more direct method for getting estimates of neural activity uses measurements of regional glucose uptake (FDG-PET) that is related to metabolic activity and, in turn, is a marker for steady-state neuronal activity. This technique has been applied to measure steady-state brain activity in individuals with tinnitus [15–17].

![Fig. 18.1](image) [18F] deoxyglucose (FDG) positron emission tomography (PET) of a patient with right-sided tinnitus. The transversal slice through the temporal brain region shows unilaterally increased metabolic activity in projection to the left auditory cortex.
The unilateral activation pattern resembles findings from Lookwood and colleagues, who observed unilateral auditory cortex activation in individuals with tinnitus using a different method [10]. A major limitation of all published FDG PET studies of tinnitus patients is that data analysis has been restricted to the auditory cortex by using a region of interest approach [19].

**Functional Magnetic Resonance Imaging (fMRI)**

fMRI is a specialized form of MRI that is used for identifying regions of the brain, where neural activity increases in response to neural stimulation, such as sensory stimulation. The use of fMRI is based on the finding that magnetic properties of hemoglobin depend on its oxygenation level and the observation that blood flow and blood oxygenation is closely related to neural activity. Regional changes in hemoglobin oxygenation occur because of local neuronal activation e.g., in response to a stimulus or during a specific task. Blood oxygenation level-dependent (BOLD) contrast is the basis of brain mapping using fMRI. BOLD contrast provides in vivo real-time maps of blood deoxygenation in the brain under normal physiological conditions. Brain regions of increased oxygen consumption are depicted by comparison of two MRI images: one at rest and one with increased oxygen consumption due to a specific task.

The use of fMRI offers several advantages over PET. First, participants are not exposed to ionized radiation; second fMRI is easier to perform, less expensive, and more widely available. Furthermore, fMRI provides high spatial resolution (1 × 1 × 1 mm) [20].

The fMRI obtained in individuals with bilateral tinnitus show symmetrical activation in all investigated areas of the auditory pathways (auditory cortex, thalamus, and inferior colliculus) while all published studies show that in individuals with unilateral tinnitus altered activation patterns only of structures contralateral to where the tinnitus is perceived were observed [21–24]. Lanting et al. found an increased sensitivity of the contralateral inferior colliculus to the loudness of the presented acoustic stimuli, whereas Melcher and Smits observed reduced neuronal activation in the contralateral inferior colliculus in individuals with tinnitus.

For the correct interpretation of these data, it is important to remember that functional MRI activation always represents a comparison between two activation states. Energy usage by the brain depends largely on firing rate [25], and it has been shown that high-frequency activity in the gamma range correlates with the BOLD signal in the auditory cortex [26, 27]. Thus, increased spontaneous neural activity, such as postulated in tinnitus patients, may imply only a limited increase in activity during stimulation with sound due to a ceiling effect. According to this saturation model, hypoactivation in fMRI has been interpreted as a possible indicator of pathologically increased neuronal spontaneous activity.

Functional MRI can also provide information about the tonotopic organization of the auditory cortex (Fig. 18.2). In individuals with pure-tone tinnitus, fMRI during presentation of a tone with the frequency of the tinnitus has made it possible to determine the anatomical localization of the representation of the tinnitus frequency in the primary and secondary auditory cortices. The fMRI technique has been used for finding the best placement of the stimulating electrode for epidural cortical stimulation [28]. However, there

![Fig.18.2 Visualization of auditory cortex activation induced by sounds of different frequencies: activation induced by 4 kHz is displayed in green, activation caused by 0.5 kHz in red](image)
are no fMRI studies available that investigated systematically potential alterations of the tonotopic map in tinnitus patients.

In a recent study, fMRI studies were combined with an emotional paradigm in order to identify tinnitus-related changes in emotional processing (Rosengarth et al., personal communication). The study showed signs of abnormal neural activity in the hippocampus, the parahippocampal gyrus, and the amygdala that were independent from depressive comorbidity. These results provide further experimental evidence for the involvement of limbic brain structures in tinnitus [29, 30].

Summary of the Use of Imaging Methods for Studies of Tinnitus

A summary of the findings of functional imaging studies in tinnitus is given in table 1 and in Fig. 18.3 PET studies have shown which areas of the brain are involved in the tinnitus network: primary [10, 15, 17, 31] and secondary auditory cortex, extending into the temporoparietal junction (=auditory association area) [1, 2, 9, 31]; (para)hippocampus [10]; medial geniculate body [10]; anterior [32] and posterior cingulate cortex [4, 33]; and precuneus and inferior lateral parietal cortex [12]. Voxel-based morphometry adds the subgenual anterior cingulate cortex extending into the nucleus accumbens area [34], and both VBM and fMRI add inferior colliculus [21, 24, 35]. Most of the neural networks activated by tinnitus overlap with brain regions that are involved in attention to and processing of normal sounds include the primary and secondary auditory cortex, parahippocampus, amygdala, as well as the right superior, middle, and inferior dorsolateral prefrontal cortex [33, 36] have the important difference that tinnitus-related activity seems to be predominantly unilateral. More recent studies have shown that other brain areas are activated by aversive sound stimulation.

<table>
<thead>
<tr>
<th>Area</th>
<th>Individuals with tinnitus compared to controls</th>
<th>Individuals with tinnitus: changes in tinnitus, induced by:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Steady-state metabolism</td>
<td>Somatosensory modulation Gaze Lidocaine</td>
</tr>
<tr>
<td>Primary auditory cortex</td>
<td>†1,2,3</td>
<td>†4</td>
</tr>
<tr>
<td>Secondary auditory cortex</td>
<td>†A</td>
<td></td>
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<tr>
<td>Auditory assoc. cortex</td>
<td></td>
<td></td>
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<tr>
<td>Thalamus</td>
<td>†A</td>
<td>†4</td>
</tr>
<tr>
<td>Inferior Colliculus</td>
<td></td>
<td>†5</td>
</tr>
<tr>
<td>Auditory brainstem</td>
<td></td>
<td>†6,7,9,10</td>
</tr>
<tr>
<td>Limbic system</td>
<td>†A</td>
<td></td>
</tr>
<tr>
<td>Frontal lobe</td>
<td></td>
<td>†7,10</td>
</tr>
</tbody>
</table>

Legend:
†: increased asymmetry of FDG uptake
‡: increased response to sound; †: reduced response to sound
†: increased and reduced rCBF corresponding to increased and reduced tinnitus
A: Abnormal assymetry

Studies:
and are related to the reward and emotional system, such as nucleus accumbens and insula [37].

New treatment strategies have evolved from fMRI and PET findings of abnormal neuronal activity in the auditory cortex [38, 39]. The results of several pilot studies using low-frequency transcranial magnetic stimulation (see Chap. 88) and direct electrical stimulation of the auditory cortex (see Chap. 90), which have been based on findings in imaging studies, have shown promising results for treatment of some forms of tinnitus. The application of brain stimulation for treatment of tinnitus patients most likely requires guidance by neuroimaging studies. Areas of the auditory cortex that have shown signs of increased metabolic activity have been identified as targets for stimulation. PET scan is now used for guidance of treatment. Preliminary findings suggest that the results of PET scanning may also serve as a predictor of the outcome of treatment [17, 32, 40].

Even data that come from relatively small sample sizes emphasize the value of fMRI, SPECT, and PET for developing new therapeutic strategies, but also show the potential of these procedures to become clinical tools for the diagnostic differentiation between different forms of tinnitus and for the assessment of treatment outcome.

References

Chapter 19
Findings from Structural Neuroimaging

Berthold Langguth and Michael Landgrebe

Keypoints

1. Structural differences in brain morphology have been detected in individuals with tinnitus compared to control groups.
2. These structural alterations involve auditory and limbic brain structures, suggesting that tinnitus may arise when alterations in both auditory and limbic brain structures occur.
3. It has yet to be clarified whether the observed structural changes represent causes or consequences of tinnitus.

Keywords Tinnitus • Neuroimaging • Magnetic resonance imaging • Voxel based morphometry • Morphological segmentation

Abbreviations

MRI Magnetic resonance imaging
TMS Transcranial magnetic stimulation
VBM Voxel based Morphometry

Introduction

Recent studies, in the search for neural correlates of tinnitus functional neuroimaging, have been complemented by structural imaging research. The search for structural differences between individuals with tinnitus and control groups can in some instances identify brain areas, which are involved in the pathophysiology of tinnitus. Similar to functional imaging, the exact clinical characterization of the pathology of an individual patient is of the utmost importance as is the control for confounding factors such as hearing loss or psychiatric comorbidity. In the interpretation of imaging results, it must be considered that differences in brain structure between individuals with and without tinnitus may be signs of (1) changes that have occurred before the tinnitus became manifest, thus signs of risk factors, (2) the consequences of tinnitus that reflect plastic changes involved in tinnitus pathogenesis, or (3) a combination of both.

Structural imaging studies for research purposes are usually based on high-resolution magnetic resonance imaging (MRI). Different analyses can reveal either changes in gray matter volume of specific brain regions in an individual person (individual morphologic segmentation; [1]) or alterations in the concentration or ratio of volume of gray and white matter at the group level without any restriction to a specific brain region (voxel-based morphometry, VBM; [2]). The major advantage of VBM is that it is a semi-automatic, investigator-independent method, which allows researchers to detect structural alterations in the brain without an a priori hypothesis. This method is therefore ideally suited to give accurate insights into physiological [3] and pathophysiological (e.g., [4]) neuroplastic processes.

Studies Using Voxel-Based Morphometry

Two VBM studies have been published to date [5, 6]. In both studies, individuals with tinnitus and similar clinical characteristics (normal audiogram, no psychiatric...
comorbidity) have been investigated. Both studies revealed structural changes in the auditory and non-auditory system. Muhlau et al. found an increase of gray matter in the medial geniculate nucleus of the thalamus and a decrease of gray matter in the subcallosal region, including the nucleus accumbens. In the study of Landgrebe et al., areas of decreased gray matter were found to be located in the inferior colliculus and the hippocampus. Interestingly, tinnitus has been described after lesions occurring in the inferior colliculus [7] and the hippocampus [8]. The decrease of gray matter in the inferior colliculus suggests that hyperactivity in that area, as demonstrated in animal studies (see Chap. 16) and in functional imaging studies in humans (see Chap. 18), may be a compensatory mechanism. An increase in thalamic gray matter concentration may be the consequence of deprivation of input to the auditory nervous system due to a dysfunction in the auditory periphery. Change in the function of the inferior colliculus may also influence corticothalamic loops.

The Auditory Cortex

Surprisingly, both of the VBM studies mentioned above did not show any changes in the auditory cortex, despite the fact that functional imaging (see Chap. 18) as well as TMS studies (see Chap. 88) indicate that the auditory cortex is involved in the pathophysiology of tinnitus. This could be due to the high inter-individual variability in the morphology of the auditory cortex, which makes it difficult to detect changes of gray matter density with VBM. A study that used a different technique for assessing potential structural changes in the auditory cortex, namely individual morphological segmentation of the medial partition of Heschl’s gyrus [1], found that individuals with tinnitus had significantly smaller auditory cortex volumes than controls. In individuals with unilateral tinnitus, this effect was almost only seen in the hemisphere ipsilateral to the affected ear. In bilateral tinnitus, the volume of the medial Heschl’s gyrus was substantially reduced in both hemispheres. This tinnitus-related volume reduction occurred across the full extent of medial Heschl’s gyrus and was not limited only to the high-frequency part usually most affected by hearing loss-induced deprivation of input to the auditory nervous system.

The Role of Limbic Structures in Tinnitus

In some individuals with tinnitus, the observed change that occurs in non-auditory brain regions, such as the hippocampal [6] and subcallosal regions of the orbitofrontal cortex [5], provides further evidence of the limbic system’s involvement in the generation and maintenance of tinnitus and associated symptoms, as postulated by several models of tinnitus pathophysiology [9, 10]. These findings support the assumption that tinnitus arises if there is (1) increased neuronal activity in specific parts of the central auditory pathways as a consequence of hearing loss (deprivation of input to the auditory nervous system) and (2) a central deficit in cancelation of meaningless signals in the auditory pathways.

Thus, several types of studies seem to agree that tinnitus-related activity in the central auditory pathways is relayed in parallel to limbic structures, which might act to evaluate the importance of the auditory signal. Habituation to meaningless sensory signals, which are mediated by the subcallosal region, the nucleus accumbens, and the hippocampus, is normally assumed to cancel out the tinnitus signal at the thalamic level and prevents the signal from being consciously perceived [5]. However, if this filter fails, increased activity in the central auditory system (e.g., due to altered sensory input) might be consciously perceived as tinnitus and the limbic structures may become involved in forming negative emotional associations with the tinnitus sound.

Conclusion

High-resolution magnet resonance imaging has revealed alterations of the brain structure both in the central auditory pathways and in the limbic system of individuals with tinnitus. Whether the observed structural changes may represent the cause or the effect of tinnitus is yet to be explained. Longitudinal studies may indicate whether the observed structural changes result from tinnitus-related functional changes or whether they constitute a vulnerability factor in the generation of the abnormal neural activity that causes tinnitus.
References

Chapter 20
A Global Brain Model of Tinnitus

Winfried Schlee, Isabel Lorenz, Thomas Hartmann, Nadia Müller, Hannah Schulz, and Nathan Weisz.

Keypoints

1. Subjective tinnitus is characterized by the perception of a phantom sound in the absence of any physical source.
2. While transient tinnitus usually lasts only a couple of seconds to a few hours, chronic tinnitus is an ongoing conscious perception of sound for more than 6 months with low incidence of spontaneous remissions.
3. Empirical studies in animals and humans often show enhancement of cortical excitability in the auditory areas associated with the tinnitus.
4. Theoretical and experimental studies suggest an additional involvement of extra-auditory cortical regions, especially the frontal cortex, the parietal cortex, and the cingulum.
5. Using magnetoencephalographic recordings, we found that these areas are functionally connected with each other and form a global fronto–parietal–cingulate network.
6. The top–down influence of this global network on auditory areas is associated with the distress that is perceived by many individuals with tinnitus.
7. We suggest that both entities – the enhanced excitability of the central auditory system and the integration with a global cortical network – are important to generate and maintain a conscious percept of tinnitus.
8. This chapter will concentrate on how a conscious perception of tinnitus is formed and maintained throughout a lifetime.

Keywords

Chronic tinnitus • Conscious perception • Global network • Cortical connectivity • Top–down • Long-range connectivity

Abbreviations

ACC Anterior cingulate cortex
AM Amplitude modulation
dB Decibel
DPFC Dorsolateral prefrontal cortex
EEG Electroencephalography
ERS Event-related synchronization
Hz Hertz
MEG Magnetoencephalography
OF Orbitofrontal cortex
PCC Posterior cingulate cortex
PDC Partial directed coherence
PET Positron emission tomography
rCBF Regional cerebral blood flow
SLIM Synchronization by loss of inhibition model
SPL Sound pressure level
SSR Steady state response

Introduction

Subjective tinnitus is characterized by a conscious perception of a sound in the absence of any physical source. This sound is typically described as a tone, a hissing or roaring noise, and in some cases as a combination of several sounds. Transient tinnitus, a phenomenon perceived by a large percentage of the population at least once in a lifetime, typically lasts a few seconds to a few hours. A far smaller percentage (5–15%) of people in western societies reports hearing their tinnitus constantly for more than 6 months [1]. Such an
ongoing perception of tinnitus has the potential to impair the ability to concentrate, to disturb sleep, to affect social interactions, and it may even cause psychiatric problems. Indeed, about 1–3% of the general population reports that their tinnitus adversely affects their quality of life [2].

In this chapter, we will concentrate on how a conscious perception of tinnitus is formed and maintained in the brain. In general, our sensory systems constantly receive an overwhelming amount of sensory input, which – although being processed within the nervous system e.g., primary sensory regions – does not enter our consciousness completely. In fact, most of the stimuli remain unconscious and our conscious perception is limited to only a few sensory events. In recent years, several studies have investigated the neural mechanisms for the conscious perception of sensory stimuli [3–6] and with the Global Neural Workspace Hypothesis, Dehaene et al. proposed a model for conscious visual perception that explains several empirical findings and observations [3, 4]. In short, this model points out two requirements for conscious perception: (1) activation of the respective sensory area and (2) the entry of this activity into a global network of long-range cortical couplings. In light of this view, we want to review the current knowledge about chronic tinnitus and suggest a model for the perception of tinnitus.

In the following chapter, we will summarize the findings of altered activity in the central auditory and nonauditory regions as well as the cross-talk between auditory and nonauditory areas in tinnitus. We will give a short overview of the studies on conscious perception of external stimuli. Finally, we suggest a model for tinnitus perception that has the potential to explain several tinnitus phenomena and propose new methods of tinnitus therapy.

Increased Excitability of the Central Auditory System

Many individuals with tinnitus are able to localize their tinnitus to one or both ears. In most cases, the tinnitus sensation is accompanied by an audiometrically measurable damage to the cochlea. Thus, one may think that the tinnitus is generated within the ears; however, this is most likely not the case. If the phantom sound was generated within the ears, a transection of the auditory nerve would reliably eliminate the ongoing perception of the tinnitus sound. To date, there is much evidence refuting this view. There are only a small percentage of patients in whom the auditory nerve section leads to relief from tinnitus. The majority of patients still experience tinnitus after surgical sectioning of the auditory nerve [7, 8]. Furthermore, if tinnitus was generated in the periphery, a systematic enhancement of spontaneous activity in auditory nerve fibers would be present. As summarized by Eggermont and Roberts [9], changes in the spontaneous firing rate of the auditory nerve are rather unsystematic. When tinnitus is induced experimentally in animals, spontaneous auditory nerve activity may be enhanced, reduced, or even remain the same. Thus, the tinnitus perception is elicited irrespective of the utilized technique and the accordant changes in auditory nerve activity. These results suggest that for the majority of individuals, the sensation of tinnitus originates from central rather than peripheral parts of the auditory system. There is a large body of studies demonstrating the importance of central structures in tinnitus. Tinnitus-related changes of spontaneous activity can be found throughout the central auditory system. The spontaneous firing rate is enhanced in the dorsal cochlear nucleus [10], the inferior colliculus, and the primary and the secondary auditory cortex [9].

Neuroimaging studies in humans also suggest a hyperactive auditory cortex in tinnitus. In some individuals, tinnitus can temporarily be suppressed by masking or lidocaine application. Mirz et al. [11] used this effect to investigate changes in regional cerebral blood flow (rCBF) during tinnitus suppression in the positron emission tomograph (PET). They reported a significant reduction of rCBF in the right temporal lobe during tinnitus suppression. However, changes in nonauditory structures were also observed, which will be discussed in the next section. In another PET study, individuals with tinnitus were distracted from their symptoms with the serials seven test (counting silently backwards in steps of seven), which led to a reduction of rCBF in the left and the right auditory cortices [12]. Neuroimaging recordings of tinnitus patients during resting state differ from recordings of individuals who do not have tinnitus inasmuch as the individuals with tinnitus experience an ongoing phantom sound. Using magnetoencephalography (MEG) in resting state recordings, Weisz et al. [13] reported a significant enhancement of delta (1–4 Hz) activity and a concomitant reduction of alpha (8–12 Hz) activity in individuals with tinnitus. These changes were most prominent in
the temporal regions and correlated with the subjective rating of tinnitus distress. A later analysis on an extended dataset also showed a significant increase of gamma frequencies (40–90 Hz) in the left and right temporal lobe of the tinnitus group [14]. These results fit well into a recently proposed framework that explains enhanced synchronization of auditory activity by a reduction of cortical inhibition (“Synchronization by Loss of Inhibition Model,” SLIM, [15]). Synchronized alpha activity is often assumed to be an indicator for active cortical inhibition mechanisms: A decrease in alpha power is associated with an increase in cortical excitability [16–18], while an increase in alpha power (also called Event-Related Synchronization, ERS) reflects inhibition [16]. The alpha desynchronization, as observed in chronic tinnitus, reflects a release of inhibition and thus favors the synchronization of neuronal activity. Altogether, the elevated rCBF (in PET), the enhancement of gamma band synchronization (in MEG), and the augmented spontaneous firing rate (single-unit recordings in animals) all act as an indicator for increased excitability of the auditory cortex in tinnitus.

Integration of Auditory and Nonauditory Brain Activity

Changes in brain activity accompanying tinnitus are not restricted to the auditory cortices. In the study by Mirz et al. referred to above, tinnitus suppression was accompanied by a reduction of rCBF in the temporal lobe, but also in the frontal lobe and posterior brain regions [11]. The MEG study by Weisz et al. [13] demonstrates alpha power decrease and delta power increase mainly located in the temporal lobe, but also extending into frontal and parietal sites.

Furthermore, there are also reports of structural changes in gray and white matter regarding chronic tinnitus. In a voxel-based morphometry study, Mühlaus displayed a decrease of gray matter density in subcortical regions and a gray matter increase in the posterior thalamus, and the medial geniculate body for tinnitus patients compared with healthy controls [19].

These results suggest an involvement of extra-auditory brain regions in the generation and/or perception of the phantom tinnitus sound. As hypothesized earlier by Jastreboff, the neural activity that causes tinnitus is generated within the auditory system, while nonauditory regions are involved in encoding the conscious percept as well as the emotional evaluation of it [20]. This hypothesis is supported by a study conducted in the 1960s, which revealed that a disconnection of the prefrontal cortex resulted in a reduction of tinnitus annoyance in most of the surviving patients [21]. Also, almost all clinicians are aware of anecdotal evidence that chronic tinnitus patients are often not aware or disturbed by their tinnitus (e.g., when distracted), but it can become the focus of attention or brought back into conscious awareness at any time. Based on these results and theoretical considerations, we postulate the existence of a widespread tinnitus network functionally connecting auditory and nonauditory brain regions. If such a network existed, there should be a considerable difference in the long-range cortical networks between participants with tinnitus and control participants who do not report an ongoing perception of tinnitus.

Furthermore, if the connectivity between auditory and nonauditory regions encodes tinnitus distress, a correlation between the functional inter-regional connectivities and tinnitus distress should be revealed. We challenged these suppositions in three studies with MEG recordings in tinnitus and non-tinnitus control participants.

In the first study, we employed auditory steady-state responses (SSR) to entrain the tinnitus network and investigated long-range functional connectivity across various nonauditory brain regions [22]. We presented amplitude-modulated (AM) tones of three different carrier frequencies to 22 participants (12 individuals with tinnitus and 10 controls). One of these stimuli was designed to match the individual tinnitus sound and the two other were control tones that were 1.1 and 2.2 octaves below the frequency of the tinnitus. Cortical connectivity was analyzed by means of phase synchronization in the participants with tinnitus and in healthy controls. We found a deviating pattern of long-range functional connectivity in tinnitus that was strongly correlated with individual ratings of tinnitus intrusiveness. Phase couplings between the anterior cingulum and the right frontal lobe as well as phase couplings between the anterior cingulum and the right parietal lobe demonstrated significant condition times group interactions. They were correlated with individual tinnitus distress ratings in the tinnitus condition. This study provided the first evidence for tinnitus-related alterations in the long-range synchronization between distant brain regions outside auditory areas.
The second study aimed to investigate the cortical networks in the resting state [23]. The analysis was based on a sample of 41 participants: 21 individuals with chronic tinnitus and 20 healthy control participants who did not have tinnitus. Cortical coupling was again analyzed by means of phase-locking analysis between distant brain regions. We found a significant decrease of inter-areal coupling in the alpha (9–12 Hz) band and a significant increase of inter-areal coupling in the 48–54 Hz gamma frequency range for the tinnitus group. Furthermore, an inverse relationship \( (r = -0.71) \) of the alpha and gamma network coupling was observed for all participants. Discrimination analysis revealed a separation of 83% between the tinnitus and the control group based on the alpha and gamma couplings. Post hoc analysis showed an influence of tinnitus manifestation on gamma coupling. In the participants who had a short tinnitus history, the left temporal cortex was predominant in the gamma network, whereas in the participants who had a longer tinnitus duration, the gamma network was more widely distributed across the cortex.

This study demonstrated disturbances in the long-range cortical coupling in individuals with tinnitus under resting conditions. The resting state is of particular interest for tinnitus research since individuals with this condition typically report an enhanced perception of the tinnitus when they are in a quiet surrounding. The results of the second study are in line with several other findings demonstrating the emergence of functional connectivity across widely distributed brain areas, in association with a conscious perception of the stimulus [3, 4, 6, 24, 25]. This connectivity may be an important mechanism of the brain in binding different features of the stimulus to form a comprehensive perception. Additionally, this connectivity might serve as an amplifier that enhances the neuronal activity in sensory areas (e.g., [3]).

In a recently published framework on chronic tinnitus, Weisz et al. proposed a top–down influence of higher order brain areas on the cortical activity in the auditory cortex [15]. With the third study [26] we specifically aimed to assess this top–down influence using partial directed coherence (PDC) – a measure that is based on the concept of Granger causality and allows for investigating the directionality of information flow between distant brain regions in the frequency domain.

Using MEG, we investigated the long-range cortical networks of individuals with chronic tinnitus \( (n = 23) \) and healthy controls \( (n = 24) \) in the resting state. A beam-forming technique was applied to reconstruct the brain activity at source level, and the directed functional coupling between all voxels was analyzed by means of Partial Directed Coherence. Within a cortical network hubs are brain structures that either influence a great number of other brain regions or are influenced by a great number of other brain regions. A strong outflow in this context indicates that this brain area considerably influences the activity of other brain structures. In the tinnitus group, two brain regions were identified with stronger outflow and one site with a weaker outflow. Stronger outflows were located in the prefrontal cortex and in the posterior part (parieto-occipital/occipital) of the brain. The weaker outflow was found in the orbitofrontal cortex (OFC). All these changes in the outflow behavior were found for the gamma frequency band above 30 Hz. A strong inflow means that this brain area is strongly driven by other brain regions.

With respect to the inflow characteristics, we found two sites with significant group differences. The OFC received more inflow in the high-frequency gamma range in the tinnitus group compared to the control group. Posterior parts of the cortex received less inflow from other brain areas in a broad frequency range, including slow waves, alpha, low beta, and gamma frequencies. Furthermore, we found the inflow to the temporal cortices correlated positively with subjective ratings of tinnitus distress: the more the activity in the temporal cortices was driven by other brain regions, the stronger the subjective distress reported by the participants. Additionally, we demonstrated that the inflow to the temporal cortex mainly originates from the prefrontal cortex and the posterior part of the brain; both are structures that we have characterized with a strong outflow within this network.

**A Short Notion on Long-Range Cortical Networks**

Long-range synchronization of distant brain regions has been first reported by Gray et al. [27]. They revealed synchronized oscillatory responses between neighboring columns in the visual cortex of the cat. Based on this finding, they proposed that synchronization combines different features of the visual pattern, which is
A Global Brain Model of Tinnitus

In this chapter, we propose a model for the conscious perception of the tinnitus sound, which is based on the above-mentioned studies on long-distance cortical coupling and extends earlier tinnitus models by Jastreboff [20], Eggermont and Roberts [9], and Weisz et al. [15]. Two levels of tinnitus-related neuronal processing are distinguished in this framework: the local (or sensory) level refers to the activity in the auditory areas. The global level refers to long-range cortical network of functionally connected brain areas.

The Sensory Level

Tinnitus is frequently associated with hyperactivity and enhanced synchronization of neuronal activity in the auditory cortex. Animal studies have shown a systematic...
enhancement of spontaneous neuronal activity of the dorsal cochlear nucleus, the inferior colliculus, the primary auditory cortex, and the secondary auditory cortex (see [9] for a review). Moreover, studies in humans with chronic tinnitus revealed tinnitus-related changes in oscillatory activity of the temporal cortex [13, 14, 37, 38]. In a very recent study, we investigated rock musicians who perceived a transient tinnitus after a loud (~120 dB SPL for ~2 h) band practice. Resting-state activity in the MEG was recorded at two time points: immediately after the practice and at a second day without exposure to loud music. We found a strong enhancement of gamma frequency power (55–85 Hz) in the right temporal cortex during the perception of transient tinnitus, which was also observed on the single participant level in 13 of 14 participants and, importantly, was not correlated with the degree of hearing loss (Ortmann et al., submitted).

The hyperactivity in auditory areas (i.e., enhanced spontaneous firing rate as observed in animal studies) and the stronger synchronization of neuronal activity (as observed through an increase of oscillatory power in the gamma frequency range) both argue for an enhanced excitability of the auditory cortex in tinnitus [9]. The absence of an alpha effect in our transient tinnitus study could imply that a down-regulation of inhibition sets in after synchronization of excitatory neurons and could play a crucial role in the transition to chronic tinnitus. A down-regulation of inhibition would require less excitatory activity to ignite a tinnitus-related cell assembly, putatively evolving into “spontaneous synchronization” (i.e., where spontaneous activity (firing) of neurons suffices for synchronization of excitatory neurons [15]). We suggest that the enhanced spontaneous synchronization of circumscribed tonotopically organized regions of the central auditory system is one necessary prerequisite for the perception of tinnitus.

The Global Level

A second requirement for the conscious perception of tinnitus is the activation of a global network characterized by long-range coupling between distant cortical regions. The brain contains a highly organized pattern of functional connectivity for which we report multiple evidence of disturbance in cases of tinnitus. Based on our studies, we suppose the tinnitus-related global network to spread over the entire cortex. However, four core regions are emphasized particularly: (a) the dorsolateral prefrontal cortex (DPFC), (b) the orbitofrontal cortex (OFC), (c) the anterior cingulate cortex (ACC), and (d) the precuneus/posterior cingulate cortex (PCC). Furthermore, top–down influence of these higher order regions on the auditory cortices modulates the neuronal activity therein. The prefrontal cortex and the precuneus/PCC regions are the main areas for this top–down modulation. This idea of a tinnitus-related global network is an application of the global workspace hypothesis as suggested by Dehaene et al. onto chronic tinnitus [3, 4]. They postulated the existence of global workspace neurons that are distributed over distant areas of the cortex, characterized by a disproportionately large amount of long-range excitatory connections. Information that is processed within this network can easily be accessed by various brain systems; hence, it is hypothesized that this workspace is the basis for conscious perception. People with chronic tinnitus report an ongoing perception of the tinnitus sound. Thus, we propose that the tinnitus sound is constantly kept in the global workspace.

Furthermore, we suppose a top–down influence from the fronto-parietal-cingulate network on the temporal cortices that enhances the neuronal excitability therein. The magnitude of this influence is mediated by the subjectively perceived tinnitus distress. Support for this assumption comes from the above-described study in which we demonstrated significant correlations between the strength of the inflow hubs and the tinnitus distress. As outlined above, we presume that desynchronized alpha activity reflects a state of reduced intracortical inhibition and enhanced neuronal excitability. In a previous MEG-study [13], we demonstrated that the decrease of alpha power in temporal regions correlated strongly with the tinnitus distress as reported by the participants in our study. Two mechanisms are likely to influence alpha power decreases in the resting state: (1) a profound hearing loss that is frequently associated with the occurrence of tinnitus might lead to loss of lateral inhibition in the tonotopically ordered auditory cortex and (2) a top–down influence from higher order brain regions on the temporal cortex might further affect the cortical excitability. Here, we assert that the later mechanism plays the more prominent role in tinnitus of the chronic state. This is largely supported by the fact that temporal alpha desynchronization
correlates well with the tinnitus distress ratings, but not with hearing loss (Fig. 20.1).

In summary, the models state that there are two processes that modulate excitability in the auditory cortices: at the sensory and the global level. The explanation at the sensory level takes into account that chronic tinnitus is usually associated with a profound damage to the hearing system (ear or auditory nerve). The reduced sensory input leads to a decrease of inhibitory mechanisms in the central auditory system and ultimately to an enhancement of cortical excitability therein (and favors the synchronization of spontaneous neuronal activity). We assume that this is the central mechanism in the generation of the phantom sound at tinnitus onset. The second explanation emphasizes a top–down influence of the global tinnitus network on the auditory cortices. We suggest that tinnitus-related information is processed in the globally extended fronto–parieto–cingulate network with influence on the auditory cortex. The magnitude of this influence is positively associated with the strength of the perceived tinnitus distress. Stronger tinnitus distress is characterized by stronger top–down influence leading to a marked alpha desynchronization, which is a neuronal signature of reduced cortical inhibition. We suppose that this mechanism is especially involved in the maintenance of the tinnitus-related enhancement of neuronal excitability in later periods of the tinnitus history. This is supported by the fact that we found significant correlations between tinnitus distress and top–down connectivity, but no results for the bottom–up connectivity.

**Implications of the Model for the Treatment of Tinnitus**

The proposed model explains the partial success of current therapies for tinnitus like Neurofeedback (see Chap. 87), transcranial magnetic stimulation (TMS), and cognitive therapies (see Chap. 73). Repetitive transcranial magnetic stimulation (rTMS) (see Chap. 88) aims to reduce the enhanced excitability in the auditory cortex, which leads to a reduction of tinnitus loudness [39–43]; however, a complete relief of tinnitus is rare. Regarding the global brain model of tinnitus, this is not surprising. Even if rTMS successfully reduces the enhanced excitability in the auditory cortices, the amplification by the global network would constantly fight against it. On the other hand, it has been shown that cognitive therapies also reduce tinnitus symptoms to some extent [20, 44]. In our proposed framework, we speculate that cognitive therapies are able to alter the tinnitus-related global network by changing the conscious elaboration of the tinnitus percept. This can potentially reduce the top–down amplification of the global network on the temporal lobe and thus lower the enhanced excitability therein, though there is still an untreated abnormal pattern of spontaneous activity in the temporal cortex that results from damage to the peripheral hearing system. If this abnormal spontaneous activity reaches a certain threshold, it can enter the global network again by means of the “bottom–up mode” as explained above.
Therefore, we stress the importance of a combination of both branches in tinnitus therapy: reducing the enhanced excitability in the auditory cortex on the one hand (e.g., via rTMS), and changing the global network on the other hand (e.g., via cognitive therapies). We strongly suggest combining both treatment approaches and expect synergy effects that improve the benefit from current tinnitus therapies.

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References

33. Just, MA, Cherkassky, VL, Keller, TA, Kana, RK, Minshew, NJ, Functional and anatomical cortical underconnectivity in autism: evidence from an FMRI study of an executive func-
tion task and corpus callosum morphometry Cereb Cortex, 2007 17:951–961
35. Le Van Quyen, M, Navarro, V, Martinerie, J, Baulac, M, Varela, FJ, Toward a neurodynamical understanding of ictogenesis Epilepsia, 2003 44 Suppl 12:30–43
37. Kahlbrock, N, Weisz, N, Transient reduction of tinnitus intensity is marked by concomitant reductions of delta band power BMC Biol, 2008 6:4
Chapter 21
A Heuristic Pathophysiological Model of Tinnitus

Dirk De Ridder*

Keypoints

1. Tinnitus pathophysiology should explain both tinnitus distress and tinnitus intensity.
2. Distress in tinnitus is most likely generated by an aspecific distress network consisting of the amygdala–anterior cingulate and anterior insula.
3. Tinnitus intensity might be encoded by gamma band activity in the contralateral auditory cortex.
4. This gamma band activity might result from thalamocortical dysrhythmia.
5. Tinnitus distress can be seen as phase-synchronized co-activation of the auditory cortex activity and the aspecific distress network.
6. For tinnitus to be perceived consciously, it requires the auditory cortex activity be embedded in a larger network.
7. This larger network could be the global workspace, the self-perception network.
8. The tinnitus network changes in time, hypothetically via an allostatic mechanism.
9. In chronic tinnitus, the parahippocampus, insula, and dorsolateral prefrontal cortex networks are critical.
10. The parahippocampus is involved via its auditory sensory gating mechanism, suppressing redundant auditory information.

Keywords
Tinnitus • Gamma • Theta • Thalamocortical dysrhythmia • Distress • Deafferentation • Plasticity • Reorganization • Networks

Abbreviations
AC Auditory cortex
ACC Anterior cingulate cortex
BA Brodmann area
BOLD Blood oxygen level dependent
BPS Band pass small
BPW Band pass wide
BRAfN Brain research center antwerp for innovative & interdisciplinary neuromodulation
CAS Complex adaptive systems
DACC Dorsal part of ACC
DLPFC Dorsolateral prefrontal cortex
EEG Electroencephalography
ERP Event related potential
FMRI Functional magnetic resonance imaging
Hz Hertz
IC Inferior colliculus
ICA Independent component analysis
IPS Intraparietal sulcus
IEEG Intracranial EEG
LORETA Low resolution electro tomography
LTP Long term potentiation
MCS Minimally conscious state
MD Mediodorsal
MEG Magnetoencephalography
MGB Medial geniculate body
NB Nucleus basalis
OF Other frequency
PET Positron emission tomography
PCC Posterior cingulate cortex

*With a critical review by the members of the TRI neurostimulation workgroup: Nathan Weisz, Berthold Langguth, Marco Congedo, Winnie Schlee, Arnaud Norena. Other reviewers include Ana Belen Elgoyhen, Elsa van der Loo, Sven Vanmeete, Mark Plazier, Thomas Elbert, Paul van de Heyning, and Aage Møller. Not every idea presented in this heuristic model has full support of all the reviewers.
Introduction

If rational treatments for tinnitus are to be developed, its pathophysiology needs to be understood. However, current knowledge of auditory system physiology is largely insufficient for this purpose. Available data on auditory physiology and the neural correlate of tinnitus can be supplemented by translating physiological data from other systems studied more extensively, such as the visual and somatosensory systems, and by extrapolating from pathophysiological mechanisms known in potentially analogous symptoms such as pain. Being aware of the limitations and the potential risks of such an approach, the proposed model has to be considered as a heuristic approach that results in the generation of testable hypotheses and needs to be corrected and improved accordingly.

Pathophysiology of Tinnitus

The pathophysiological working model of tinnitus has to include the mechanisms involved in the generation of the auditory percept and the intensity of a phantom sound as well as the mechanisms causing the tinnitus-related distress.

Tinnitus Intensity

The auditory system consists of two main parallel pathways supplying auditory information to the cerebral cortex; the same two ascending systems also have a descending counterpart, the tonotopically organized parvalbumin staining lemniscal system and the non-tonotopic calbindin staining extralemniscal system [1–4]. The lemniscal pathways use the ventral part of the medial geniculate body, the neurons of which project to the primary auditory cortex, whereas the extralemniscal pathways use the dorsal part of the medial geniculate body that projects to the secondary auditory cortex and association cortices, thus bypassing the primary cortex [5], Table 21.1. While neurons in the lemniscal pathways only respond to auditory stimulation, many neurons in the extralemniscal pathway are multimodal. Neurons in the ventral thalamus fire in a tonic or semitonic mode while neurons in the dorsal thalamus fire in bursts [6, 7]. Burst firing consists of dense packets of action potentials followed by periods of quiescence [8]. Information theory suggests that, in general, both tonic and burst firing efficiently transmit information about the stimulus. Burst and tonic firing might therefore be parallel computations in the auditory and other sensory systems [8, 9] (Table 21.1).

Based on the differences between the two parallel auditory pathways – the lemniscal being tonotopic and the extralemniscal being less tonotopic – it has been hypothesized that white-noise tinnitus may be caused by synchronous hyperactivity of burst firing in the non-tonotopic extralemniscal system, whereas pure-tone tinnitus may be the result of increased synchronous tonic firing in the tonotopic (lemniscal) system [43]. Narrow band tinnitus could be the result of a co-activation of the lemniscal and extralemniscal pathways.

Tinnitus Distress Matrix

The same subjectively reported tinnitus intensity can be related with severe distress in some people but may well be tolerated in others. The emotional component involved in tinnitus is most likely generated in the emotional circuit imbedded in our brain. Components of the emotional system are the amygdala, the subgenual and dorsal anterior cingulate cortex (ACC), the anterior insula, the ventromedial prefrontal cortex (VMPFC), and the orbitofrontal cortex [44–47]. Some of these areas such as the amygdala [48], the ACC [49], and the orbitofrontal cortex [50] are also involved in the reward system, together with the ventral tegmental area, nucleus accumbens, and mediodorsal nucleus of the thalamus [51].
The lemniscal system – aka the specific system, the tonotopic system, or the non-classical system – has the following characteristics:

- Phyllogenetically old [11, 12]
- Unconscious reflexes [13, 14]
- To secondary cortex [1, 2, 15, 16]
- Less tonotopic [1, 7, 17, 18]
- Slow spontaneous firing rate [19] [20]
- Variable latency response [18, 21, 22]
- Rapid habituation to repetitive stimuli [17, 18, 22]
- Fires predominantly in burst mode [6, 7]
- Stimulus detector [23, 24]
- Non-linear [24–26]
- Overrides tonic mode [24–26]
- Processes changes in auditory environment [24, 27]
- Calbindin positive [1, 16, 28]
- CB increases after deafferentation [29–32]
- Multimodal [17, 33–36]

The extralemniscal system – aka the non-specific system, the non-classical system, or the interoceptive default state – has the following characteristics:

- Phyllogenetically recent [11, 12]
- Conscious perception [13, 14]
- To primary sensory area [1, 2, 15, 16]
- Tonotopic [1, 7, 17, 18]
- Higher spontaneous firing rate [37, 38] [39–41]
- Short latency response [18, 21, 22]
- Slower habituation to repetitive stimuli [17, 18, 22]
- Fires in tonic mode [6, 7]
- Feature detector [23, 24]
- Linear [24–26]
- Weaker than burst mode [24–26]
- Processes the content of change in the auditory environment [24, 27]
- Parvalbumin positive [1, 16, 28]
- PV decreases after deafferentation [42]
- Unimodal [34]

The brain resolves perceptual ambiguity by anticipating the forthcoming sensory environment, generating a template against which to match observed sensory evidence. The ventromedial prefrontal cortex has been implicated as the source of this template [52]. Positive feedback results when sensory evidence is indeed as predicted and raises hemodynamic activity in the ventral striatum (nucleus accumbens) and the posterior cingulated cortex, related to a reward and storing the received information, respectively; negative feedback activates the dACC and the anterior insula, mediated via the habenula [53]. Thus, when the brain has not obtained the information, it needs to guide subsequent behavior, it activates the dACC–insula network to get more information.

Whenever new information is presented, the brain cannot compare this to a template, and therefore activity levels of the dorsal ACC (dACC) might also reflect the salience of the new information for predicting future outcomes [54, 55], guiding optimal decision making in an uncertain world [56].

Functional connectivity studies reveal that the dACC is functionally connected to the anterior insula [57] as well as the thalamus and brainstem [58]. The combined dACC–anterior insula activity possibly subserves intrinsic alertness [58], as the dACC and anterior insula are co-activated during states of arousal [55, 59, 60] and anticipatory arousal [61]. It has been shown that the amount of baseline activity in the dACC and insula predicts how intense a subsequent pain stimulus is being perceived [62]. The combined anterior insula and dACC activation has been suggested to act as a switch from the interoceptive default state to an exteroceptive executive brain state [63].

The human dACC has developed a parallel specialization for motivational drive via a thalamocortical pathway relaying in the mediodorsal thalamus [49]. The direct activation of both the interoceptive cortex and the dACC by the distinct homeostatic modalities corresponds with the simultaneous generation of both a sensation and a motivation [49, 64]. Thus, the function of the dACC might be to integrate motivationally important information with appropriate autonomic and motor responses [61] related to the survival needs of the body [64]. This might be based on the reward learning system, which uses dopamine as one of its major neurotransmitters. Dopamine neurons emit an alerting message about the surprising presence or absence of rewards [65, 66]. Dopamine neurons in the ventral tegmental area (VTA) are activated by rewarding events that are better than predicted, remain uninfluenced by events that are as good as predicted, and are depressed by events that are worse than predicted.

The right anterior insula has been implicated in interoceptive awareness [64, 67] related to the autonomic nervous system, the amygdala could be a relevance detector [68], and the ventromedial prefrontal cortex could be a major link between the autonomic nervous system, regulation of emotion, and stress reactivity [69]. Imaging studies on distress in posttraumatic stress disorder (PTSD) demonstrate activation of the amygdala, insula, medial prefrontal cortex, and anterior cingulate.
cortex [70], which overlaps with the distress network noted in pain and tinnitus. In anxiety disorders (such as social phobia, specific phobia, or PTSD) during emotional processing, the amygdalae and insulae are hyperactive; in PTSD specifically, the dACC and medial prefrontal cortex are hypoactive [71]. This could hypothetically reflect the brain’s suppression of the salience (dACC [54]) of the traumatic template (VMPFC [52]). Thus, even though the same network is active, its composing structures might be differentially activated depending on the task and pathology involved.

In tinnitus, using whole head magnetoencephalography (MEG) phase synchronization analysis has shown that functional connectivity between ACC and the right frontal lobe and ACC and right parietal lobe is correlated to tinnitus intrusiveness, a measure of tinnitus distress. The phase synchronization between ACC and right frontal lobe was inversely correlated with tinnitus intrusiveness, whereas the phase synchronization between ACC and right parietal lobe was positively correlated with tinnitus intrusiveness [72].

Even though no specific studies have looked at the tinnitus distress, Positron Emission Tomography (PET) studies have demonstrated activation of this distress network as well. Tinnitus distress, as measured by the Tinnitus Questionnaire (TQ) [73], is correlated with anterior cingulate activity [74], and the anterior insula is activated in tinnitus [75].

It has been suggested that there is a lateralization of the two components of the autonomic system, with the right insula controlling the sympathetic system and the left insula the parasympathetic system [59, 76, 77]. The same lateralization has been found in the ventromedial prefrontal cortex [78, 79], consistent with earlier data on hemispheric lateralization of parasympathetic and sympathetic control [80]. This could explain why the difference between severe but compensating and severe but decompensating tinnitus distress is related to activation of the right anterior insula (Vanneste submitted), confirmed by heart rate variability data correlated to anterior insula spontaneous activity (van der Loo, unpublished data). Both studies are based on Low Resolution brain Electric Tomography (LORETA) EEGs [81] (Fig. 21.1).

Based on the clinical analogies between tinnitus distress and pain distress and based on neuroimaging data, it is tempting to speculate that the tinnitus distress network and the pain matrix are identical [82]: unpleasantness of pain activates the anterior cingulate [83] and orbitofrontal cortices, amygdala, hypothalamus, posterior insula, primary motor cortex, and frontal pole [84]. One may further speculate that the perception of tinnitus and pain intensity could be related to auditory and somatosensory cortex activation, respectively, but that the distress associated with its perception might be related to activation of a common general non-specific “distress network.” This notion is supported by a recent study that demonstrates activation of this distress network during unpleasant symptoms in a somatoform disorder, even in the absence of a real physical stimulus [85].

Furthermore, the emotional network involved in pain and dyspnea [86] is similar, suggesting that the distress network might be a non-specific system that can be activated by many different kinds of external and internal stimuli.

The conscious perception of tinnitus distress and pain distress could be due to a co-activation of the thalamocortical auditory and somatosensory activity and distress network activity, possibly through synchronization of neuronal activity [72]. This heuristic model can also explain the clinical observation that tinnitus distress is frequently related to the development of tinnitus in stressful periods. Thus, a person in which the distress network is already sensitized, for whatever reason (divorce, work-related problems, etc.), would be more vulnerable to develop distressing tinnitus by increased activation of the auditory system. Once established, the co-activation between the auditory pathways and the distress network might stabilize and become self-sustaining.

Developmental and Adult Plasticity

Plasticity refers to the capacity of the nervous system to modify its organization [87]. The response of the nervous system to environmental changes can involve functional and structural changes. These changes can be induced not only by normal sensory input but also by abnormal sensory input, adaptation to damage of the nervous system, or sensory deprivation [87]. There seems to be a greater potential for plastic changes during development than during adulthood, even though similar mechanisms seem to govern both developmental and adulthood plasticity.
Any alteration of auditory input during the development of the tonotopy will result in reorganization of the tonotopic map according to the altered pattern of incoming neural activity. Thus, the Lamarckian and Darwinian (pangenesis) principle of “use it or lose it” guides both development and subsequent changes in the tonotopy. The auditory system develops in two stages [88, 89]. A first stage of synapse formation or auditory tract formation is genetically determined [90] and requires the release of a chemotropic factor [89, 91]. This is followed by fine-tuning of the synapses, leading to the formation of a tonotopic structure [92]. The development of tonotopy requires electrical activity resulting from auditory input during a critical period [93, 94]. It is the result of self-organization [95] via apoptotic resorption of surplus synapses and neurons [91, 96].

The mature auditory system still demonstrates an important capacity for reorganization, adjusting itself to any change in the auditory environment [97, 98]. The tonotopic maps are not rigid and may alter or reorganize under influence of normal physiological stimuli, as in learning, adjusting the tonotopic map to relevant environmental stimuli [97, 99, 100]. However, the plastic changes also occur in pathological situations such as sound overexposure [101], partial unilateral hearing loss [93, 102], or tinnitus [103].

In addition, the tonotopic map can also reorganize via direct cortical stimulation, as demonstrated in the big brown bat. Electrical auditory cortex stimulation can change the tonotopic map at a cortical [104], thalamic [105], or inferior colliculus level [97, 105], suggesting that the corticofugal pathway is involved in this tonotopical reorganization [98]. This corticofugal system acts as a positive feedback system, which in combination with lateral inhibition sharpens and adjusts tuning of neurons in the thalamus and inferior colliculus [98, 106]. In other words, the corticofugal system acts as a mechanism for reorganization of the thalamus and the inferior colliculus [105], adjusting the tonotopy to auditory experience [97].

Focal electrical stimulation of the cortex activates this corticofugal system resulting in reorganization of the thalamus and inferior colliculus [107], all the way to the cochlea [108], as well as the auditory cortex itself [104]. It induces tonotopic changes by decreasing best frequencies slightly higher than those electrically stimulated, and increasing best frequencies slightly lower than those electrically stimulated [104].
Auditory cortex plasticity is under the influence of the major neuromodulatory systems, such as the cholinergic nucleus basalis [99, 109], the dopaminergic ventral tegmental area [110], the serotoninergic dorsal raphe [111], and the noradrenergic locus coeruleus. The effects on the auditory cortex are understandably not identical for all these neuromodulatory systems. For example, the effect of the nucleus basalis [109] and the VTA [110] can be summarized as follows:

<table>
<thead>
<tr>
<th>Stimulation of</th>
<th>NB</th>
<th>VTA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Size of functional auditory cortex</td>
<td>Increased</td>
<td>Increased</td>
</tr>
<tr>
<td>Size of functional AI</td>
<td>No change</td>
<td>Increased</td>
</tr>
<tr>
<td>Stimulus frequency representation</td>
<td>Increased</td>
<td>Increased</td>
</tr>
<tr>
<td>Adjacent frequency representation</td>
<td>Decreased</td>
<td>Increased</td>
</tr>
<tr>
<td>Spectral selectivity</td>
<td>Increased</td>
<td>No change</td>
</tr>
<tr>
<td>Non-monotonic responses</td>
<td>Increased</td>
<td>No change</td>
</tr>
<tr>
<td>Frequency specificity of the effects</td>
<td>Sharper</td>
<td>Broader</td>
</tr>
<tr>
<td>Tuning of secondary auditory cortex</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Temporal asymmetry of the effects</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Modulation of stimulus-following rate</td>
<td>Undetermined</td>
<td>Yes</td>
</tr>
<tr>
<td>Cross-area synchronization</td>
<td>Yes</td>
<td>Does not apply</td>
</tr>
</tbody>
</table>

The differential effects of these neuromodulatory systems on auditory cortex plasticity might benefit future tinnitus treatments.

### Deafferentation, Tinnitus, and Synchronized Auditory Hyperactivity

In tinnitus, firing rate and synchrony of firing are increased both in the extralemniscal and in the lemniscal systems. In the extralemniscal system, increased firing is observed [120–122] in the dorsal and external inferior colliculus [120], the thalamus [123], and the secondary auditory cortex [121, 122]. Furthermore, quinine, known to generate tinnitus, induces an increased regularity in burst firing, at the level of the auditory cortex, inferior colliculus, and frontal cortex [124]. This fits with the fact that in tinnitus an increased synchrony is found in the cochlear nerve [125–127] and auditory cortex [128, 129]. In tinnitus, an increased tonic firing rate is present in the lemniscal system as demonstrated in the lemniscal dorsal cochlear nucleus [130–135], inferior colliculus [136–139], and primary auditory cortex [140]. Interestingly, in the primary auditory cortex, not only tonic firing is increased, generating the phantom sound, but also the burst firing [129] at a regular basis.

Repetitive stimulus presentation results in decreased neuronal response to that stimulus, known as auditory habituation at the single cell level [141], also known as auditory-mismatch negativity at multiple cell level [141, 142]. Tinnitus is usually constantly present, i.e., there is no auditory habituation to this specific activation at this specific frequency.

### Plasticity and Reorganization in Tinnitus

After noise trauma, tonotopic organization in the cortex is changed such that cortical neurons with characteristic frequencies in the frequency region of the hearing loss no longer respond according to their place in the tonotopic map but reflect instead the frequency tuning of their less affected neighbors [112, 113]. Providing an acoustically enriched environment, spectrally matching the hearing loss prevents this reorganization [114]. Neurons in the reorganized region also demonstrate spontaneous hyperactivity and increased neural synchrony [115–117], which can also be abolished by providing a spectrally matched and enriched acoustic environment. Magnetic source imaging studies [103] confirm this reorganization in humans: the auditory cortex is reorganized such that the frequency area corresponding to the tinnitus pitch is represented adjacent to where magnetic activity is expected on the tonotopic axis. Furthermore, in this study, the amount of reorganization was correlated with the perceived strength of the tinnitus, similarly to what is found in phantom pain [118]. In tinnitus patients, this reorganization is not correlated with the amount of hearing loss [103], which is the primary activator of changes in tonotopic maps [119]. This suggests that reorganization of the cortical tonotopic map, changes in neuron response properties, and tinnitus are correlated.
This corresponds, to some extent, to habituation deficits described in chronic pain.

The Neural Correlate of Tinnitus: Gamma Band Thalamocortical Firing

The EEG power spectrum (of the oscillation rate) and the level of consciousness are correlated [143]. Slow delta frequencies (0.5–4 Hz) are recorded in patients under deep sleep, anaesthesia, and coma. Somewhat higher frequencies, called theta waves (4–7 Hz), are noted in light sleep, and alpha waves (8–13 Hz) are recorded from all sensory areas in a resting state. Frontal beta waves (13–30 Hz) are recorded predominantly when people pay attention to external or internal stimuli. Synchronization of separate gamma band activities (30–80 Hz), present in different thalamocortical columns [144], is proposed to bind [145, 146] distributed neural gamma activity into one coherent auditory percept [147–152]. In general, coherent gamma band activity is present only in locally restricted areas of the cortex for short periods of time [152–156]. Thus, persisting gamma activity localized in one brain area can be considered pathological.

Recent data from the visual system suggest stimuli that reach consciousness and those that do not reach consciousness are characterized by a similar increase of local gamma oscillations in the EEG [157, 158]. Thus, gamma band activity, per se, is not related to conscious perception. Data from the olfactory bulb, as homologue for the thalamus, indicate that percept of odor could be related to amplitude modulation of the gamma band, suggesting that the gamma band is no more than a carrier wave [159, 160]. This idea is based on the fact that a signal (information) must sometimes be attached or superimposed on other voltages at frequencies that move easier in the transmission medium. Attaching signals to other carrier signals is called modulation. Carrier waves are known frequencies that can be readily detected (using a narrow bandwidth receiver tuned to transmitted signal). Retrieving the tinnitus-related information from the gamma carrier wave might therefore be attempted by different methods: by amplitude modulation analysis, frequency modulation analysis, pulse modulation analysis, or by completely different methods such as principal or independent component analysis (ICA) of the spectrally filtered gamma band or raw EEG.

In clinical practice, source analysis of the gamma band activity in tinnitus patients can be performed with LORETA EEGs [81]. If gamma band activity is localized in the auditory cortex, an ICA of the raw EEG filtered for gamma band activity can be performed, and the independent component that co-localizes with the gamma band activity could be considered to contain the tinnitus-related information. Intracranial recordings (iEEG) give a unique way to measure brain activity directly at the site of the electrode, bypassing skin and skull resistance. Comparing these intracranial recordings to simultaneously recorded scalp EEG activity, validation of the independent components measured at scalp level has been given at the site of the intracranial electrode [161]. According to our data, the ICA of scalp EEG could indeed serve as a tool to detect the neural correlate of tinnitus, similarly to what has been suggested for contralateral auditory cortex gamma band activity [162, 163]. Incorporating this concept into the thalamocortical dysrythmia model of Llinas (see below for further information), 40 Hz is a carrier wave, carrying the tinnitus-related information, which could potentially be represented by a co-localized gamma band filtered independent component (Fig. 21.2).

Fig. 21.2 Independent component analysis performed on a 19-channel EEG recording in a patient with right-sided pure-tone tinnitus. The 16th independent component co-localizes with 40 Hz activity. Note that this component is not based on gamma band filtered EEG, which would be essential for if looking for the tinnitus information carried on the gamma wave.
Thalamocortical Dysrhythmia

Tinnitus correlates with gamma band activity, and Llinas has developed this hypothesis further in this thalamocortical dysrhythmia model [163]. This model can be summarized as follows: the thalamus and cortex are interconnected and act in a coherent way. In the sleeping state, the thalamus fires at 4–7 Hz (1–3 Hz during slow wave sleep); in the resting awake state the thalamus fires around 10 Hz, driving the cortex to fire at the same rate [164]. When auditory stimuli are presented, the thalamocortical rhythm becomes activated and increases its firing rate to gamma band activity (>30 Hz). However, in a deafferented state, the thalamocortical columns fire in a burst mode with a frequency of 4–7 Hz. This leads to a decrease of lateral inhibition in the adjacent areas and results in a halo of gamma band activity, called the edge effect. It is hypothesized that this spontaneous and constant gamma band hyperactivity causes tinnitus [156].

Tinnitus is usually constantly present, which suggests that tinnitus-related gamma activity is continuously present, in contrast to normal physiological gamma activity, which waxes and wanes [152–156]. Therefore, it should be possible to retrieve this gamma band activity from the auditory cortex by analyzing short-term recordings of spontaneous electrical activity from the brain. Magnetoencephalography studies demonstrate that indeed gamma band activity is increased in the auditory cortex contralaterally to the side of tinnitus perception [162]. Whether the gamma band activity in the auditory cortex is related to the percept per se or is just an intensity coding mechanism is not clear. The first LORETA EEG data suggest that the spontaneous gamma band activity might be encoding tinnitus intensity [165].

Using data from implanted electrodes overlying the secondary auditory cortex, power versus frequency plots can be made of spontaneous electrical activity. The normal power versus frequency plots demonstrate the typical individual alpha peak of the sensory cortices. In thalamocortical dysrhythmia tinnitus, a theta peak can sometimes be found on iEEG recordings (De Ridder, submitted) similarly to what has been described for MEG. When recording during a period of residual inhibition, after electrical stimulation at the area of the theta peak when no more tinnitus is present, the theta peak disappears, suggesting that the theta peak is causally related to the tinnitus, either the theta itself or, hypothetically, via the decrease of nested gamma [166]. This seems to confirm Llinas’ model, at least at a cortical level.

When analyzing four implanted patients, in whom stimulation results in a decrease of tinnitus intensity, iEEG recordings can be performed with tinnitus at two different tinnitus intensities: one performed while the tinnitus is at rest and another performed during a period of residual inhibition. Theta band activity is higher on all poles of the electrodes when tinnitus intensity is high in comparison with low (Z = −1.826, p = 0.068), a nearly significant result with only four patients.

Using co-registration of the preoperative functional Magnetic Resonance Imaging (fMRI) and the postoperative CT, it can be shown that gamma band activity is highest at the area of Blood Oxygen-Dependent Level (BOLD) activation in all patients. These data give some support at a group level for the idea of thalamocortical dysrhythmia.

Tinnitus is usually constantly present, indicating that no habituation occurs for the tinnitus-related neuronal activity. Using EEG-mismatch negativity, abnormalities have been demonstrated in tinnitus sufferers who are specific to frequencies located at the audiometrically normal lesion edge as compared to normal hearing controls [167], which is compatible with Llinas’ thalamocortical dysrhythmia model [163].

Thalamocortical Dysrhythmia and Reorganization Go Hand in Hand

Increased “synchrony” in theta and gamma band firing in thalamocortical dysrhythmic tinnitus may induce cortical reorganization by simple Hebbian plasticity mechanisms [168]: cells that fire together, wire together. This model would predict that over time the tinnitus-related neuronal changes become more and more stabilized and the tinnitus more difficult to treat. Hebbian learning in the adult requires that the event is behaviorally relevant, i.e., input from nucleus basalis (NB) and VTA in addition to the firing of cortical cells or thalamocortical circuits in parallel. Therefore, the model would emphasize appraisal of the tinnitus, only predicting long-term changes when the tinnitus is given significant attention. The central nucleus of the amygdala and midbrain–striatal dopamine systems are critically involved in the alteration of attentional and
emotional processing of initially neutral stimuli by associative learning [169–171], via its influence on the VTA [169] and nucleus basalis [170]. The insula and anterior cingulate receive the most pronounced innervations from the VTA [172]. It has been demonstrated that 10–50 Hz stimulation at the VTA (in contrast to the MD nucleus of the thalamus) activates the anterior cingulate via a dopaminergic pathway in a frequency-dependent manner [173].

Thus, co-activation of the dorsal ACC with the anterior insulae could result in attaching salience [54, 174, 175] to the tinnitus sound, resulting in reward-based Hebbian long-term plasticity as a (clinically negative) consequence. The dACC exerts a top–down influence on secondary auditory cortex (BA22) gamma band responses [176]. Cortical gamma band activity with associated attentive behavior is under control of the dopaminergic VTA [177]. Stimulating the VTA together with an auditory stimulus of a particular tone increases the cortical area and selectivity of the neural responses to that sound stimulus in AI and via coherent activity in A2 as well [110].

The anterior insula is not only involved in sound detection and in the entry of the sound into awareness but also in allocating auditory attention and in processing of novel versus familiar auditory stimuli [178]. Lesions in the anterior insula lead to contralateral auditory agnosia [179–181].

Under physiological situations, the hippocampus detects new information, which is not already stored in its long-term memory as it arrives. The resulting novelty signal is conveyed through the subiculum, accumbens, and ventral pallidum to the VTA where it contributes (along with salience and goal information) to the novelty-dependent firing of these cells. This results in dopamine release within the hippocampus producing an enhancement of Long-Term Potentiation (LTP) and learning [182]. In the auditory system, the auditory input enters the hippocampus via the parahippocampus [183, 184]. Complex novel sounds in humans activate the left and right superior temporal gyrus and the left inferior and middle frontal gyrus as well as the left parahippocampal gyrus [185]. In a similar fashion, the left superior temporal and left parahippocampal gyrus, along with left inferior frontal regions, are associated with listening to meaningful sounds [186]. The parahippocampal area is involved in sensory gating of irrelevant or redundant auditory information after both 100 ms and 400 ms [183]. This area is activated with the dACC, which peaks at 120 ms and after 240 ms [187]. It is of interest that onset of auditory hallucinations is related to activation of the left anterior insula and right middle temporal gyrus [188, 189], associated with deactivation of the parahippocampal area and anterior cingulate [188].

Thus, in summary, the amygdala might perceive a sound as salient or not [190], which activates the VTA [169] to mobilize the dACC and insulae [173], switching the default state to an executive brain state [63]. The dACC exerts a top–down influence on A2 [191], from where the left parahippocampal area is also activated if the sound is novel [185] or meaningful [186]. The VTA and the (tinnitus) sound result in plastic changes in the primary auditory cortex and from there in the secondary auditory cortex [110]. The posterior parahippocampus is the main node of entry for auditory information from A2 to the medial temporal lobe memory system, where salient information is encoded into long-term memory [184]. The parahippocampus also has an auditory gating function, suppressing irrelevant or redundant auditory information [183], as the dACC does somewhat earlier [187, 192]. Thus, when the dACC and parahippocampus are deactivated, as in the onset of complex auditory phantom percepts (hallucinations), the irrelevant and redundant information is not suppressed anymore, and the activation of the anterior insula and temporal cortex permits the internally generated auditory information to be perceived consciously and attended to [178]. Thus, it can be hypothesized that tinnitus onset could be characterized by deactivation of the dACC and parahippocampus, with activation of the insula and superior temporal gyrus.

**Extending Thalamocortical Dysrhythmia to Darwinian Plasticity: Reverse Thalamocortical Dysrhythmia**

Thalamocortical dysrhythmia predicts that the hyperactive symptoms related to gamma band activity are expressed at the lesion edge, thus adjacent to the missing sensory input. However, both in the auditory system [193] and in the somatosensory system [194], phantom perceptions are those coming from the missing input and not from the edge. This could be explained by including Darwinian plasticity to the thalamocortical
dysrhythmia model. Sensory deafferentation results in expansion of the adjacent non-deafferented region into the vacated area, both in the somatosensory and in the auditory cortex. It has been suggested that a reverse form of plasticity could also exist: deafferented sensory cortex neurons seek information elsewhere in an attempt to survive (hence the name Darwinian plasticity). Neurophysiological and neuroanatomical data, functional imaging, clinical and human electrical brain stimulation data suggest a Darwinian model of brain plasticity. This model is capable of explaining deafferentation-induced symptomatology, which was not well explained by classical plasticity [195]. Whereas the lemniscal thalamocortical dysrhythmia model predicts a reduction of the oscillation frequency in deafferentiated thalamocortical columns, the proposed reverse thalamocortical dysrhythmia model can explain that the deafferented thalamocortical units also oscillate at gamma frequencies and thus can generate phantom percepts that fit the clinical data. Due to increased lateral inhibition related to gamma activity, a halo of low-frequency activity will develop at the lesion edge. This could be called reverse thalamocortical dysrhythmia, which explains that the perceived tinnitus pitch matches the deafferented frequencies (Fig. 21.3).

Cortical reorganization in tinnitus can be visualized using MSI (Magnetic Source Imaging, a fusion of MEG and MRI; Muhlnickel, Elbert et al. [103]). However, MEG is an expensive technique, restricted to a very limited amount of research centers. Therefore, using fMRI as a means of visualizing tinnitus would be advantageous in routine clinical practice, as this technique is available at many clinics and can provide images at high resolution.

fMRI measures a relative difference in oxygen consumption between a resting state and activated state. BOLD contrast takes advantage of the fact that the magnetic properties of haemoglobin depend on its oxygenation. The blood oxygenation in turn reflects changes in neuronal activity. As such, BOLD contrast can be used to provide in vivo real-time maps of blood oxygenation in the brain under normal physiological conditions [196]. Thus, a focal area of increased oxygen consumption can be depicted by subtraction of two MRI images, one at rest and one with increased oxygen consumption due to a specific task. As increased oxygen consumption is correlated to increasing metabolic demands, the BOLD effect is related to event-related synchronization of gamma band activity [197], and BOLD is highly coupled to gamma local field potentials (EEG) in the auditory cortex [198, 199]. This strongly suggests that fMRI can visualize the gamma band-synchronized activity associated with tinnitus.

A scanning paradigm, using music as a stimulus, adequately visualizes the auditory pathways in tinnitus.

Fig. 21.3 Heuristic pathophysiological model of tinnitus intensity generation (Figure by Jan Ost, RN)
patients [200]. fMRI activation is symmetrical in patients with bilateral tinnitus at all investigated areas of the auditory pathways (auditory cortex, thalamus, and inferior colliculus). fMRI activation is significantly decreased in patients with right-sided tinnitus in the left primary auditory cortex (AC) and in the left inferior colliculi (IC). In patients with left-sided tinnitus, fMRI activation is significantly decreased in the right medial geniculate body (MGB). In summary, the contralateral auditory pathways seem to be involved in patients with unilateral tinnitus. fMRI activation always represents a difference in neural activity instead of absolute neural activity. An increase of spontaneous neural activity, such as postulated in tinnitus patients, would mean that the affected brain area during the rest condition is more active than the unaffected side, and that the active condition (sound presentation) will only give rise to a limited increase in activity due to a ceiling effect (known as the saturation model) in comparison to the non-affected side. This can explain the fact that constant pathological neuronal hyperactivity can be correlated to hypoactivation in fMRI [200].

A similar study for tinnitus using tinnitus pitch and character-specific stimuli is currently being conducted. In this study, we compare BOLD activation for tinnitus-specific sound presentation to non-tinnitus sounds presented in the scanner. Only tinnitus-specific sounds induce a significant BOLD change, as demonstrated by a lateralization effect, in contrast to non-tinnitus sounds, which generate a bilateral symmetrical BOLD activation.

Even for tinnitus-specific frequencies, the exact representation might be important. For patients suffering from pure-tone tinnitus, auditory presentation of a pure tone generates a marked asymmetrical BOLD activation, whereas presentation of a narrow band noise creates less BOLD activation, and a white noise generates almost no asymmetry (Kovacs, unpublished data) (Fig. 21.4). However, as mentioned before, auditory tract activation is insufficient to objectively diagnose tinnitus solely based on functional imaging.

A disadvantage of fMRI studies is that a contrast is needed, e.g. by presenting a sound and comparing this to a resting state or other conditions (e.g. other sound). The active condition may include different unspecific components, e.g., different arousal and differences in a patient’s understanding of verbal instruction. Therefore, fMRI studies might suffer from various confounds. The fact that the fMRI-related activation changes are specific for the perceived phantom sound (Fig. 21.4) does, however, suggest that fMRI can indeed be used to study tinnitus.

**Isolated Thalamocortical Dysrhythmia and the Global Workspace Model (Electrophysiologically Explored)**

As tinnitus is a persistent conscious auditory percept, it is important to understand the neural correlates of auditory consciousness, defined as the minimal
neuronal mechanisms jointly sufficient for any auditory conscious percept [201]. This understanding is an essential requirement for defining neurostimulation targets to suppress this auditory phenomenon. It has been suggested by Crick and Koch that in the visual system V1 activation is necessary but insufficient for visual awareness [202, 203]. Thus, isolated thalamocortical dysrhythmia in the primary auditory cortex is most likely not enough to generate the conscious percept of tinnitus. Studies in patients in persistent vegetative state (PVS), who are awake but without awareness – without conscious percepts [204] – demonstrate that these patients have a decreased metabolism in a network of areas consisting of midline areas, such as the anterior cingulate (ACC), which extends into the ventromedial prefrontal cortex (VMPFC), and the posterior cingulate (PCC), which extends into the precuneus. However, the lateral cortical regions also have less metabolic activity, more specifically the parietal and dorsolateral prefrontal cortex areas [205]. Not only is metabolism decreased in these patients, but functional connectivity is also decreased between the intralaminar nuclei of the thalamus and ACC/VMPFC and PCC/precuneus regions, and between ACC/VMPFC and PCC/precuneus [204, 205]. Recovery from PVS is associated with normalization of metabolism and connectivity, suggesting this decreased metabolism and loss of connectivity is critically involved and causally related to the neural correlate of consciousness [206, 207]. Extending these studies to auditory processing of patients in PVS, it was shown that the activation associated with auditory stimuli was restricted to the primary auditory cortex bilaterally in patients in a PVS without functional connectivity between the secondary auditory cortex and temporal and prefrontal association cortices [208], similarly to what has been shown for pain processing [209]. Based on these data, it can be proposed that activity restricted to the primary auditory cortex does not lead to auditory conscious perception, similarly to the somatosensory and visual system, but that this auditory activity becomes conscious when functionally connected to the ACC/VMPFC and prefrontal cortex (BA10) [206].

Baars has proposed the global workspace theory [210], which was extended and electrophysiologically refined for the visual system by Dehaene [211, 212]. The global workspace model, as perfected by Dehaene, can be translated to the auditory system as follows [213]: in (unconscious) preconscious processing, auditory stimulus processing is blocked at the level of the global neuronal workspace, i.e., it remains limited to the primary auditory cortex, while the global workspace is temporarily occupied by another task or is non-active, such as in PVS. A preconscious auditory stimulus may be temporarily buffered within the primary auditory cortex (discussed below) and later accessed by the frontoparietal system, once it is released by its present distracting task. In this case, information switches from unconscious to conscious. Conscious processing occurs when the accumulated stimulus-evoked activation exceeds a threshold and evokes a dynamic state of global reverberation [214] ("ignition") across multiple high-level cortical areas forming a “global neuronal workspace,” particularly involving prefrontal, cingulate, and parietal cortices, the same areas that are decreased in metabolism and functional connectivity in PVS. These areas can maintain the information online and broadcast it to a variety of other processors, thus serving as a central hub for global access to information – a key property of conscious states.

Subliminal processing corresponds to a data-limited situation where the auditory stimulus reaches only specialized cerebral sensory networks (i.e., secondary and auditory association areas), without reaching a threshold for global ignition and, thus, without global reportability. The orientation and depth of subliminal processing may nevertheless depend on the top–down state of attention.

So, when an auditory stimulus is presented, it will activate the primary auditory cortex after about 17 to 30 ms [215, 216] and the primary auditory cortex (A1) remains activated up to 300 ms generating a Pa (P50), Nb, Na, P1 en N100 ERP [217]. This persistent A1 activation is characterized by an early (85 ms) posterior and a late (115 ms) anterior N1 component [218, 219]. In other words, the primary auditory cortex neurons synchronize multiple times to generate positive and negative ERP peaks. At 50 ms, the information is not only processed in the primary, secondary, and association auditory cortex [220] but also in the frontal cortex [183, 221], more specifically, in Brodmann’s areas 6 and 24 [192, 221, 222]. There might be a parallel signal transmission to the ACC and auditory cortex analogous to what has been shown for the somatosensory system. Somatosensory stimuli arrive at the ACC and somatosensory cortex simultaneously, as evidenced by intracranial recordings of evoked potentials [223]. This might reflect
1. The dorsal attention system, which is associated with externally directed cognition, includes regions in the frontal eye fields, ventral premotor cortex, superior parietal lobule, intraparietal sulcus, and motion-sensitive middle temporal area [236–238].

2. The hippocampal-cortical memory system, a network of regions that are active during passive mental states linked to internally directed cognition (the default network) [239, 240], includes regions in ventral medial prefrontal cortex, posterior inferior parietal lobule, retrosplenial cortex, posterior cingulate, and the lateral temporal lobe [235, 236, 239, 241, 242].

3. The frontoparietal control system is an executive control system guiding decision making by integrating information from the external environment with stored internal representations [243]. It includes many regions identified as supporting cognitive control and decision-making processes including lateral prefrontal cortex, anterior cingulate cortex, and inferior parietal lobule [235].

4. The emotional system is a network based on functional connectivity with the amygdala and includes subgenual and dorsal anterior cingulate, orbitofrontal, insular, and dorsolateral prefrontal cortex, as well as strong interactions between amygdala and parahippocampal gyrus [244].

The global workspace has not been delineated anatomically. It can be hypothesized that the areas involved in the global workspace overlap with regions of these four networks.

However, that is still more than the minimal requirement for conscious perception [201].

Sleep studies have shown that the inferior and midfrontal gyrus, inferior parietal area, and medial parietal area are less active in Rapid Eye Movement (REM) sleep in comparison to wakefulness [245], suggesting that these areas are important for wakefulness and processing of external input but less important for awareness. The superior frontal and superior parietal areas with the intraparietal sulcus are equally active during wakefulness and REM sleep, as well as the VMPFC [245], suggesting that these areas are important for awareness/consciousness and could potentially be the minimal network required for awareness. It is striking that the dorsal attentional network, which selects and links stimuli and responses and hereby influences subsequent processing of stimuli in sensory cortex, is located in exactly the same areas: intraparietal sulcus (IPS) and superior parietal lobule (SPL), and dorsal frontal cortex along the precentral sulcus [237, 246], except for the VMPFC. The ventral attentional network, which interrupts and resets ongoing activity,
consists of the temporoparietal junction (including the STS and gyrus), the inferior parietal lobule, and the mid and inferior frontal gyrus as well as the frontal operculum, and anterior insula [237, 246]. Thus, the inferior parietal and mid- and inferior frontal area, which are less active in REM compared to wakefulness [245] and are part of a resetting network [246], might be critically involved in updating current conscious information processing with novel external input.

Thus, based on both PVS and sleep studies, it can be proposed that the network consisting of the superior frontal–superior parietal–VMPFC–intralaminar nuclei has to be functionally connected for internally or externally generated auditory stimuli to be consciously perceived. These areas are activated after 200–300 ms and are involved in the generation of the P300, which is one of the requirements for stimuli to be perceived consciously (in the visual system).

Subliminal stimuli can be deeply processed and activate similar brain areas as consciously perceived stimuli [158]. Both perceived and non-perceived visual stimuli cause a similar increase of local (gamma) oscillations in the EEG, but only perceived words induce a transient long-distance synchronization of gamma oscillations across widely separated regions of the brain [157, 158], compatible with the global workspace model. Furthermore, only visual stimuli that are consciously perceived induce enhanced theta oscillations over frontal regions and demonstrate an increase of the P300 component of the event-related potential and an increase in power and phase synchrony of gamma oscillations [158].

As previously mentioned, the neural generators of the auditory P300 are the inferior parietal lobe/temporoparietal junction (TPJ), the supplementary motor cortex (SMA), the dorsal anterior cingulate cortex (dACC), the superior temporal gyrus (STG), the insula, and the dorsolateral prefrontal cortex [231] (in other words, the ventral attentional network plus dorsolateral prefrontal cortex). Thus, the P300 seems to interrupt and reset ongoing activity to what is being processed in the DLPFC, or in working memory [247]. This is very similar to the frontoparietal control system [235].

It has been suggested that the P300 is the electrophysiological correlate of global workspace activation, implying that the global workspace consists of the dorsolateral prefrontal cortex, dACC, SMA, and inferior parietal area extending into the STS [248].

If auditory cortex activation is essential but not sufficient for auditory conscious perception, where is the percept being transformed into a conscious percept? Data from monkey studies in the somatosensory system suggest it could be the prefrontal cortex [249]. Activity of primary somatosensory cortex neurons covaries with the stimulus strength but not with the animal’s perceptual reports. This is similar in tinnitus: tinnitus intensity correlates with gamma band activity in the contralateral auditory cortex [165]. In contrast, the activity of the medial premotor cortex (MPC) neurons does not co-vary with the stimulus strength but does so with the animal’s perceptual reports [249]. In further agreement with the global workspace model, it has been demonstrated in the somatosensory system that the neural correlate of subjective sensory experience gradually builds up across cortical areas starting at the somatosensory cortex and ending in the premotor areas of the frontal lobe [250], which might have a hidden sensory function [251]. This idea of premotor cortex activity related to conscious sensory perception fits with the sensorimotor contingency philosophy of consciousness [252] described in the book *Action in Perception* [253], which suggests that seeing is a way of acting, a way of exploring the environment. This intentional drive sensation dates back to Aristotle and Thomas Aquinas [254] and has been proposed to be a working mechanism in olfaction as well [255].

Thus, neural activity alone is not sufficient to produce vision, but neural activity contributes to experience only as enabling mastery and exercise of laws of sensorimotor contingency [252].

It is of interest that it was shown that N1, P2, and P3 are attenuated in chronic tinnitus patients [256, 257]. However, no source analysis was performed, and N1 attenuation is not found all the time [258]. One explanation can be that N1 is only attenuated in patients with low distress [259]. Another study found a difference in N1-P2 in unilateral tinnitus sufferers on the basis of N1-P2 intensity dependence and N1-P2 amplitude. A bilateral tinnitus group differed from controls by greater intensity dependence of the N1-P2 component and shorter N1 latency [260]. Using MEG, it was also shown that amplitude ratio M200/M100 represents a clear-cut criterion to distinguish between tinnitus patients and individuals without tinnitus [261], and the abnormal M200/M100 normalized when the tinnitus disappeared [262].
However, this M200/M100 abnormality in tinnitus patients could not be confirmed by another study [263].

Based on the above-mentioned heuristic model, it can be hypothesized that the ERPs should be performed with tinnitus-matched sound and non-tinnitus-matched sound. Obtaining a LORETA ICA of N100 should correlate to two aspects of tinnitus: one component relating to tinnitus distress (the ACC component) and one component to tinnitus intensity (auditory cortex component). In a similar way, the P/N 200 should be analyzed by ICA to make the distinction between distress and intensity. Similarly, the P300 should be analyzed for the presence of tinnitus, with P3a gamma band activity examined for the presence of distress and P3b for the presence of the sound.

It can be further hypothesized that the P50 (and N400) might be abnormal in tinnitus, as there is no sensory gating involved for the tinnitus-matched sound, whereas the P50 and N400 could be normal for non-tinnitus-matched sound.

PET studies have shown which areas of the brain are involved in the tinnitus global workspace network (Fig. 21.5): primary [75, 264–267] and secondary auditory cortex, extending into the temporoparietal junction (the auditory association area) [265, 268], (para)hippocampus [75], medial geniculate body, [75], anterior [74] and posterior cingulate cortex [269, 270], and precuneus and inferior lateral parietal cortex [271]. Voxel-based morphometry adds the subgenual ACC extending into the nucleus accumbens area [272], the hippocampus, and the inferior colliculus [273], which is confirmed by fMRI [274, 275]. Magnetoencephalography also finds abnormal spontaneous activity as well in the prefrontal cortex (BA10) [276]. Most of the tinnitus network overlaps with an aversive sound-processing network consisting of the primary and secondary auditory cortex, parahippocampus, amygdala, and right superior, middle, and inferior dorsolateral prefrontal cortex [277]. Later studies extended the aversive sound network to the auditory association, nucleus accumbens, and insula area [278].

The Tinnitus Network Changes in Time

Clinical data suggest that the longer tinnitus lasts the more difficult it becomes to treat. This has been shown for microvascular decompressions [279–285] and transcranial magnetic stimulations [269, 286–288]. Even though it is most likely a gradual continuous change, tinnitus duration of 4 years might be a practical point for clinicians to differentiate acute from chronic tinnitus (De Ridder, in press, Neurosurgery). This was first noted in microvascular decompressions by Möller, later by others performing the same surgery [279–285], and most recently was extended to rTMS investigations [286, 287]. A MEG study looking at phase-locked connectivity in the tinnitus network found that in patients with a tinnitus history of less than 4 years, the left temporal cortex was predominant in the gamma network, whereas in patients with tinnitus duration of more than 4 years, the gamma network was more widely distributed including more frontal and parietal regions [289]. Thus, even though the areas involved might still be the same, the functional connectivity and weight of the hubs between the involved areas might change.

In a recent EEG study, these network changes were also analyzed spectrally. Results indicate that the generators involved in tinnitus of recent onset (<4 years) seem to change in time with increased synchronized activity contralaterally in the auditory cortex, DLPFC/premotor cortex, dACC, and insula. This is associated with an increase in gamma band connectivity between

Fig. 21.5  The tinnitus
global workspace network, as summarized from functional neuroimaging studies. Red:
anteor distress network.
Blue: posterior tinnitus
intensity network
the parahippocampal cortex, auditory cortex, and the insula ipsilaterally to the tinnitus side and DLPFC contralaterally to the tinnitus side. All other connections seem to decrease in time (vanneste, submitted).

It is interesting to note that in chronic tinnitus, the degree of response to auditory cortex rTMS on TQ distress was correlated with tinnitus-associated activation of the anterior cingulate cortex [74].

Recently, the idea of allostasis, defined as the adaptive process for actively maintaining stability (homeostasis) through change [290], has been introduced in medicine [290]. It has been shown that allostasis is controlled by the brain [291, 292]. Homeostasis relates to the mechanisms that maintain stability within the physiological systems and hold all the parameters of the organisms internal milieu within limits that allow an organism to survive [290, 293, 294]. Allostasis, on the other hand, relates to the maintenance of stability outside of the normal homeostatic range, where an organism must vary all the parameters of its physiological systems to match them appropriately to chronic demands, for example, by resetting the system parameters at a new set point [290, 295, 296]. An allostatic state has been defined as a state of chronic deviation of the regulatory systems from their normal state of operation with establishment of a new set point [296]. It has been especially investigated with regard to the Darwinian [297] adaptive nature of stress and its possible maladaptive consequences, called allostatic load. The allostatic load then leads to pathology [291, 292, 298]. Drug addiction is hypothesized to involve a change in drug reward set point and reflects an allostatic, rather than a homeostatic, adaptation (i.e., outside the normal set point) [295, 296].

The brain areas controlling allostatic stress in an organism are suggested to be the amygdala and the prefrontal cortex [291, 292, 297], as well as the ACC and insula [175]. Based on parallels between addiction and pain, it has been suggested that in chronic pain the concomitant tolerance (adaptive decreases of the drug’s efficacy) and hyperalgesia might be the result of the development of a new allostatic equilibrium [299]. Conceptually, in chronic tinnitus, a new allostatic equilibrium could develop, resulting in hyperacusis and persistence of the phantom sound. The dorsal ACC is involved in adaptive decision making and value evaluation [300] by adapting its activity when a new piece of information is witnessed, reflecting its salience for predicting future outcomes [54] by utilizing dopamine reward prediction error signals, but only when something can be learned [301]. Thus, the dorsal ACC might be involved in resetting this equilibrium. Metaphorically speaking, the dorsal ACC attributes salience to the phantom sound and resets its equilibrium allostatically, so that the sound remains consciously perceived via resetting the parahippocampal auditory gating.

The allostatic equilibrium resetting can be located in the dACC and parahippocampus, as both regions are involved in auditory sensory gating [183, 192], i.e. suppression of irrelevant or redundant auditory information. Thus, if there is an allostatic reset of what auditory information is important or not, the dACC will be important as well as the parahippocampal area.

The parahippocampus is functionally connected to the inferior lateral parietal cortex regions along the midline including posterior cingulate and retrosplenial cortex extending into the precuneus, and subgenual ACC extending into the ventral medial prefrontal cortex [241].

The posterior parahippocampus is the main node of entry for auditory information to the medial temporal lobe memory system, where salient information is encoded into long-term memory [184]. The left parahippocampal gyrus along with left inferior frontal and left superior temporal regions are specifically associated with listening to meaningful sounds [186]. The parahippocampal area has also been linked to the unpleasantness of the auditory information [302], in contrast to the left amygdala, which is related to the salience of the aversive auditory (verbal) information [190].

Based on visual system data, it has been suggested that the parahippocampal cortex may play a broad role in contextual association [303, 304]. If complex auditory phantom phenomena (such as auditory hallucinations) and simple auditory phantom phenomena (such as tinnitus) share common pathophysiological mechanisms, it is of interest to note that at onset of auditory hallucinations, the parahippocampus becomes deactivated as well as the anterior cingulate [188]. Furthermore, when analyzing the difference between responders and non-responders to auditory cortex stimulation by means of LORETA EEG, non-responders demonstrate increased theta activity in the left parahippocampus, whereas responders have increased gamma band (30–40 Hz) activity in the (left) parahippocampal area $t(9)=1.98; p<0.05$ (van der Loo, unpublished data). Perception involves the processing of sensory stimuli and their translation into conscious experience. A novel percept can, once synthesized, be
maintained or discarded from awareness. Visual perception is associated with distributed bilateral activation in the posterior thalamus and regions in the occipito-temporal, parietal, and frontal cortices. In contrast, sustained perception is associated with activation of the left prefrontal cortex and left (para) hippocampus [305]. Thus, if tinnitus is considered a sustained auditory perception, it could explain why amytal tests of the amygdalohippocampal area are capable of suppressing tinnitus in chronic unilateral tinnitus [306].

**The Tinnitus Network: A Summary**

A stimulus only makes sense if it is related to and incorporated the person’s self-percept. Therefore, the self-perception network, consisting of the ACC-vmPFC, PCC–precuneus, superior frontal-parietal, and STS, has to be activated for the tinnitus to be consciously perceived (Fig. 21.6). This is supported by the data from PVS patients.

The tinnitus intensity is related to auditory cortex activity, which might be controlled by dACC–insula baseline activity, expressing that the tinnitus is salient. The tinnitus percept, per se, might not be encoded in the auditory cortex but be represented by DLPFC–premotor activity, connected to the self-perception network via the PCC–precuneus activity. This could be analogous to the somatosensory processing, where stimulus intensity is encoded by somatosensory cortex activity and the conscious perpect in the frontal cortex. The parahippocampus might serve as an entry to auditory memory, pulling the missing information due to deafferentation from memory (Fig. 21.6).

**The Tinnitus Network: Future Perspectives**

Since the recent development of network science [307–311] to study complex adaptive systems (CAS), these analyses have been introduced in brain science [312–318] as well. The underlying idea is that CAS, whether it is the internet, ant societies, social interactions, the weather, or economy, are structured by similar universal rules [319].

Network topology describes how different nodes in a network are connected or linked. It was initially assumed that networks predominantly form randomly, in which each node is connected to another node randomly, characterized by a Poisson distribution of its connectivity [307]. All nodes are equal in this network. More recently, scale-free networks have been described [311], in which some nodes are more connected and more clustered (i.e., have a shorter path length, turning them into hubs). This suggests that some nodes are clearly more critical with regard to the robustness of the network. Both random and scale-free

![Fig. 21.6 Heuristic tinnitus network interactions](image-url)
networks are very robust to random errors, but scale-free networks are more sensitive to attacks on hubs. Eighty percent of nodes can be removed in scale-free networks without failure, but if some critically important hubs are removed, the network system fails. Most likely, these scale-free networks become incorporated into hierarchical networks [320], permitting incorporation of modularity and scale-free behavior of the network.

The approach to studying complex adaptive systems has recently been extended to the human brain, as the brain clearly fulfils the criteria of a complex adaptive system [312, 313, 315]. The topological network approach can be applied to brain anatomy [318, 321], electrical [317, 322] and magnetic brain activity [314], and blood oxygenation changes (fMRI) [312, 313].

The entire brain is not ruled by one network, but most likely different topologies exist depending on the brain area and functional state of the brain. The brainstem is organized like a small world, but is not scale free [323]. The cerebellum seems to be structured like a regular or strictly local network [324], the hippocampus more like a random [324] and small world network, and the cortex has both small world [312, 325] and scale-free [326, 327] properties. These different network systems might be integrated in a hierarchical system of functional modules [320].

This network approach to studying the brain of patients with tinnitus could benefit the future neuromodulation management for individuals with this condition. Based on this short introduction to network analysis, it becomes clear that if tinnitus is related to scale-free hub disability, neuromodulation makes sense, as with limited targeting the persistent tinnitus network might be normalized again. This will, however, be impossible in random networks and will not be useful in regular networks. A recent study demonstrates that the hubs in tinnitus might consist of the PCC, dACC, and sgACC, extending into the OFC and parahippocampal area [328]. More similar studies with higher resolution will permit future pathophysiologically-based hub targeting in tinnitus.

**Conclusion**

There is insufficient literature to develop an evidence-based neuropathophysiological model of tinnitus, but a heuristic model can be conceived when available tinnitus data are supplemented by knowledge from other sensory systems, as well as the limbic, autonomic, and motor systems. Since it has been suggested that plasticity uses similar mechanisms in all sensory areas, extrapolating information from other sensory systems seems acceptable.

Tinnitus intensity is correlated with increased gamma activity in the contralateral auditory cortex, possibly as a reaction on reduced auditory input via thalamocortical or reverse thalamocortical dysrhythmia, resulting in lack of inhibition and increased synchrony, which in turn may lead to topographic map reorganization in the auditory cortex.

The tinnitus percept, per se, is almost certainly not related to isolated synchronous gamma band activity in the auditory cortex, but requires co-activation of the ill-defined global workspace or a self-perception network.

The distress some tinnitus patients perceive seems to be correlated to increased activity in the amygdala, anterior cingulate, and right anterior insula. Tinnitus distress might also be the result of synchronization of auditory thalamocortical dysrhythmia and distress network activation.

In time, the neural generators of tinnitus might change, possibly only by spectral modifications within the tinnitus global space network, hypothetically based on an allostatic mechanism.

Future studies, applying techniques from network science might demonstrate which hubs are critical for maintaining the tinnitus percept and therefore could be good targets for tinnitus neuromodulation treatments.

**References**


Keypoints

1. There is no established methodology for clinical trials of tinnitus treatment.
2. Inter-study comparability is difficult due to insufficient characterization of investigated samples and variation of the used assessment and outcome measures.
3. Clinical trials in tinnitus should follow standards set by the guidelines of Good Clinical Practice, by the Consort statements, and should be registered in a clinical trial registry.
4. The design of the clinical trial depends on the clinical question, which should be answered by the study.
5. Placebo-controlled randomized trials represent the gold standard for testing efficacy of treatment approaches. However, in order to save resources, a stepwise approach seems reasonable, which involves pilot open trials as a first step to screen for potentially promising treatments and which is followed in case of positive outcome by randomized controlled trials.
6. Due to the heterogeneity of tinnitus, the best possible characterization of the investigated study sample, with respect to clinical or neurobiological characteristics, is highly desirable.
7. Outcome criteria for therapeutic trials have to be reliable, valid, specific, and relevant.
8. Trial design should be based on statistical estimation of sample sizes and power in order to minimize the risk of type I and type II errors.
9. To enhance inter-study comparability, international accepted standards for patient assessment and outcome measurement should be followed.
10. Standardization of clinical trial methodology will enhance clinical research in tinnitus by facilitating data comparison across trials and allowing pooled data analyses of multicenter study results in international databases.

Keywords Tinnitus • Clinical trials • Placebo • Inter-study comparability

Introduction

Clinical trials are conducted in order to answer questions regarding the safety and efficacy of new treatment options. Ideally, clinical trials will answer these questions as accurately as possible. However, there are several constraints for the design and conduct of a clinical trial, such as recruitment of patients (especially when large sample sizes are required or selective inclusion and exclusion criteria are being imposed), adherence of patients, or financial resources. Thus, the optimal trial design has to find a balance between adequacy to address the clinical question and feasibility as defined by the research infrastructure.

Randomized Controlled Trials: Bias Reduction and Ethical Issues

The gold standard for evaluating treatment efficacy is the randomized controlled clinical trial (RCT) [1].
Randomization is the key element and refers to the fact that patients are assigned to a treatment or a control group by chance. That is, a patient who has been found eligible to participate in the trial and has also provided informed consent to do so, has an equal chance to be assigned to either the treatment or the control condition. Neither the patient nor the physician is involved in this decision. In practice, treatment assignment is laid down in a randomization code that is compiled before the start of the trial by using tables of random numbers or generating a list of random numbers by computerized tools. The important conceptual and methodological consequence of randomization is – given that the sample size is sufficiently high – that any known and unknown factors related to the patient (patient history, co-morbidity, gender, age, expectations) are balanced out across treatment and control condition. Thus, the process of randomization establishes structural equivalence and eliminates biases (i.e., systematic errors that jeopardize the interpretation of study results) that are due to confounding factors.

A second major technique to reduce bias is blinding. The idea is to keep the treatment assignment of patients confidential throughout the course of the trial and ensure that all patients are treated in a consistent way. Ideally, neither patients nor doctors know which treatment or control group a patient is assigned to; this is called a double-blind study. An even more strict design is a triple-blind study, in which neither the patient and the doctor nor the researcher assessing treatment outcome or being involved in data analysis have information regarding treatment assignment. There are cases, however, in which only the patient can be blinded; this is a single-blind study. A study in which treatment assignment is known to all participants is called an open study.

The fact that randomized trials impose the element of chance regarding treatment assignment and require blinding as an additional methodological feature gives rise to substantial ethical concerns. The two most popular, and conflictive, points of criticisms read as follows: (1) patients in the treatment arm may be exposed to a dangerous regimen whose safety is not yet proven; (2) patients in the control arm may be withheld from a potential beneficial new treatment option. There is broad and undisputed social and scientific consensus that randomized controlled trials have to comply with the highest ethical standards. The two most important ethical requirements are sufficient evidence regarding the safety of the new treatment option, and, at the same time, uncertainty regarding the reliability and magnitude of its potential benefit. This state of uncertainty is called equipoise: only then a randomized trial may be conducted as a decision experiment to clarify whether there is a reliable and clinical meaningful difference between treatment and control condition.

Further Considerations When Planning a RCT

Clinical trials can be performed as monocentre or multicentre trials. Performance of RCT as multicentre trials makes it possible to include a larger sample in shorter time and also allows controlling for specific center effects, but the organization also becomes more demanding. It should also be mentioned that not in all situations are RCTs the best solution. First of all, RCTs are methodologically demanding, expensive, and time consuming. Therefore, there should be pilot data available that identify the treatment approach under the study as promising and allow estimation of the effect size of the intervention. Knowledge of the sample size, in turn, is a necessary prerequisite for sample size and power calculations. However, identifying an intervention as “promising” is a difficult task. One practical approach may be a stepwise process starting with a smaller, open pilot trial. If the results of the pilot trial have been promising, the next step can be to test this intervention in a RCT. Furthermore, data from the pilot trial may help to identify inclusion or stratification criteria. Alternatively, RCTs may be started based on promising results of preclinical experiments (e.g., animal studies). However, the lack of in vitro bioassays or validated animal models is a major problem in the development of new drugs for tinnitus therapy.

Recent promising advances in this field, however, may facilitate drug development in the near future (see Chap. 16), but it remains to be determined to what extent data from drug trials in the available animal models can be extrapolated to efficacy in humans. Another problem of RCTs is that the efficacy of a treatment is, in general, evaluated by assessing the mean change of the outcome measure for each group. Using this approach, individual patients with a very good treatment response may be missed. Hence, in addition to the evaluation of group averages, study
results should be analyzed for predictors of positive treatment response or subgroups of responders. The search for predictors of treatment response may be further facilitated by pooling the information from different trials in meta-analyses or databases, assuming that the studies have been performed according to specific standards [2]. It is important to note that such post hoc data-driven analyses have to be interpreted carefully and require further prospective confirmatory studies because they are never corrected for multiple comparisons.

Given the complexity of tinnitus and the many treatment options and possible combinations available, tinnitus may be a target for complex interventions. The UK’s Medical Research Council (MRC) has developed a conceptual and methodological framework for complex interventions [3]. The continuum of increasing evidence comprises five consecutive phases [4, 5]:

1. **Theory**: the theoretical basis is laid down suggesting a specific intervention will have the expected effect (preclinical phase).
2. **Modeling**: all components of the interventions are described and their interrelations and expected outcomes are specified.
3. **Exploratory trial**: preliminary evidence is obtained that this treatment has the intended effect. This phase helps to improve the final study design, intervention, and control groups, as well as assessment strategies.
4. **Confirmatory randomized controlled clinical trial (RCT)**: this crucial phase is designed to answer whether the complex intervention really works.
5. **Long-term implementation**: this final step includes a subsequent study that evaluates the validity of the complex intervention under real-life circumstances.

Promising examples applying this model have been published [4, 5], which may also serve as a general model for clinical tinnitus research.

**Control Groups**

Placebo-controlled trials can be performed in cross-over designs or parallel group designs. In cross-over designs, each patient represents his own control, which minimizes the influence of confounding factors. Furthermore, a much smaller sample size is sufficient to reach the same statistical power. However, the main shortcomings are that a cross-over design limits follow-up periods and requires a wash-out period before switching to the other kind of intervention (placebo or active treatment) in order to avoid the effects of the first intervention influencing the second intervention. The estimation of a sufficient duration of the wash-out time is sometimes difficult, since both known factors (e.g., long half-life times of pharmaceutical agents) and unknown factors (e.g., induced neuroplastic changes in the central nervous system, which may occur with a delay after treatment) may play a role. Thus, the design of a cross-over trial always involves a trade-off between study duration and the risk of potential carry-over effects, which may confound study results. Moreover, cross-over designs cannot be used when the timing of the intervention is critical, e.g., for the investigation of acute treatment intervention in tinnitus with recent onset.

Parallel group designs, in contrast, allow long observation periods and do not have to deal with potential carry-over effects. However, in addition to the already mentioned requirement of larger sample sizes, differences between groups, with respect to potential confounding variables, may become relevant. Such factors include, but are not limited to, age, gender, tinnitus cause and duration, tinnitus severity at baseline, level of hearing loss, co-morbid psychiatric symptoms (depression or anxiety), etc. Despite randomization, these factors may be unequally distributed in different treatment groups, especially when relatively small sample sizes are investigated. In such cases, these confounding factors have to be considered in the interpretation of the results.

As described earlier, blinding of the study is an important issue [1] to keep treatment regimens consistent and to control for effects of associated anticipation and expectation. Whenever possible, clinical trials should therefore use double blinding, which means both patient and therapist are blinded. In pharmacological trials, the use of placebo medication allows effective double blinding. However, in nonpharmacological treatments, blinding represents a substantial problem (e.g., repetitive transcranial magnetic stimulation [6, 7]) or may almost not be possible (e.g., cognitive behavioral therapy). For such interventions, waiting list controls or “treatment as usual” controls are sometimes used. However, these controls are vulnerable to expectations and unspecific effects of the
interaction between patient and therapist and tend to overestimate the effects of the active intervention. Quite generally, the choice of the optimal control condition becomes more difficult when nonpharmacologic interventions such as psychotherapy, physiotherapy, or brain stimulation are tested [8].

**Trial Duration and Follow-Up**

One limitation of current clinical trials in tinnitus is their short duration typically ending after 3 months. Since tinnitus represents in many cases a chronic condition often characterized by significant spontaneous changes in intensity, assessment of treatment effects beyond 3 months would be desirable. This is even more important in treatments, which exert their effect with delay (e.g., behavioral focused treatments like cognitive behavioral therapy or tinnitus retraining therapy). Furthermore, longer follow-up periods enable researchers to determine adherence rates to treatments, which have been reported to be quite low for some kind of interventions (e.g., the use of white-noise generators [9]).

**Sample Size and Statistical Testing**

Clinical trials may have different goals. They can be performed in order to find the optimal dose of a treatment or to demonstrate noninferiority of a specific intervention in comparison to a standard one. In most cases, the aim of a clinical trial is to determine whether or not a treatment is effective for a given indication. To achieve this, it is important that the trial has been designed in a way that false-positive and false-negative errors are minimized. False-positive means that the trial suggests a treatment is more effective than placebo, when this is not actually true; the results of the trial are due to other influences. This kind of error can be minimized by using good blinding conditions, matching for potential confounding variables, and use of randomization procedures. False-negative results mean that the trial suggests a treatment is not more effective than placebo; the reason for this result can also be that the sample size of the trial was too small in order to detect the difference between both study arms. To avoid this problem of insufficient power, definition of clinically meaningful changes (e.g., a reduction of a given number of points in a tinnitus questionnaire), power analysis, and determination of sample size is essential. Whenever possible, power calculations should be based on the results of existing studies or the preliminary findings of pilot studies [10]. After completion of the trial, statistical testing should also address the issue of potential false-negative errors. Treatment efficacy will be determined by analysis of the treatment effect on the primary outcome variable. To avoid the statistical problem of multiple testing, the primary outcome measure should be restricted to one variable. Other variables may be included as secondary outcome measures. Finally, drop-outs from the study may represent a substantial problem in the statistical analyses of the trial. Data from patients who dropped out of the trial should not be disregarded; one option is the so-called “last observation carried forward” (LOCF) approach. This kind of analysis is also called “intention-to-treat” analysis, which is the most widely used approach in analyses of clinical trials. However, if the drop-out rate gets too high, analyses and interpretation of results become difficult. Drop-out rates may increase in study populations with high spontaneous remission rates (i.e., the patients have no need to further stay in the trial), long study durations, or mild forms of tinnitus. These problems have to be kept in mind during the design of the trial. In general, it is advisable to seek statistical consultation by an expert in clinical trials in advance of planning the trial, in order to avoid such problems.

**Statistical Significance and Clinical Relevance**

There is a crucial distinction between statistical significance and clinical relevance. Statistical significance refers to the fact that a finding is statistically reliable. It is important to note that empirical research, in general, can never prove a finding is 100% correct. All solid research can do is minimize the error that false conclusions are drawn from data. In the behavioral and medical sciences, there is consensus that an error rate of less than 5% is tolerable, often expressed as \( p < 0.05 \) (sometimes stricter levels of significance are applied, such as \( p < 0.01 \) or \( p < 0.001 \)). Obtaining a difference between the treatment and the control group accepting \( p < 0.05 \) means that if the experiment is repeated 100 times, in roughly 95% of the
cases the difference will show up again and 5% of the experiments will fail to observe the effect.

Whether a result reaches statistical significance generally depends on the mean difference between the two groups, the variability of the results (standard deviation), and on the sample size. As the number of participants increases, the difference between groups required to reach statistical significance becomes smaller. As an example, with a sample size of several hundred patients, mean improvements of one or two points on a tinnitus scale may reach statistical significance. Such results are frequently criticized with the argument that such a difference may not be clinically relevant.

Clinical relevance relates to the magnitude of an effect. Thus, clinical relevance is not a statistical or methodological issue; it refers to the fact that clinicians and/or patients regard a difference between a new treatment and a control treatment as “large enough” or “important.” For instance, a difference in success rates between treatment and control of 10% may be considered clinically relevant. The definition of clinically important differences is particularly challenging with regard to questionnaire data. Questionnaires are used for detecting meaningful changes of clinical symptoms. Since the currently available questionnaires have not been developed and validated for detecting treatment-induced changes, empirical data about the minimal change required for clinical relevance is limited. For the tinnitus questionnaire, a reduction of five points has been proposed as a minimum change for an individual patient in order to be of clinical relevance [11].

**Reporting Clinical Trials: Consolidated Standards of Reporting Trials**

Consolidated standards of reporting trials (CONSORT) encompass various initiatives developed by the CONSORT group to alleviate the problems arising from inadequate reporting of randomized controlled trials. The main product of the CONSORT group is the CONSORT Statement [12, 13], which represents an evidence-based minimum set of recommendations for reporting randomized trials. It offers a standard way for authors to prepare reports of trial findings, facilitating their complete and transparent reporting, reducing the influence of bias on their results, and aiding their critical appraisal and interpretation. The CONSORT Statement comprises a 22-item checklist (Table 22.1) and a flow diagram (Fig. 22.1), along with some brief descriptive text. The checklist items focus on reporting how the trial was designed, analyzed, and interpreted. The flow diagram displays the progress of all participants throughout the trial.

**Table 22.1** The CONSORT Statement 2001 checklist (items to include when reporting a randomized trial) is intended to be accompanied with the explanatory document that facilitates its use

<table>
<thead>
<tr>
<th>Paper section and topic</th>
<th>Item</th>
<th>Descriptor</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Title and abstract</strong></td>
<td>1</td>
<td>How participants were allocated to interventions (e.g., “random allocation”, “randomized”, or “randomly assigned”).</td>
</tr>
<tr>
<td><strong>Introduction</strong></td>
<td></td>
<td>Scientific background and explanation of rationale.</td>
</tr>
<tr>
<td>Background</td>
<td>2</td>
<td>Eligibility criteria for participants and the settings and locations where the data were collected.</td>
</tr>
<tr>
<td><strong>Methods</strong></td>
<td>3</td>
<td>Precise details of the interventions intended for each group and how and when they were actually administered.</td>
</tr>
<tr>
<td>Participants</td>
<td>4</td>
<td>Specific objectives and hypotheses.</td>
</tr>
<tr>
<td>Interventions</td>
<td>5</td>
<td>Clearly defined primary and secondary outcome measures and, when applicable, any methods used to enhance the quality of measurements (e.g., multiple observations, training of assessors).</td>
</tr>
<tr>
<td>Outcomes</td>
<td>6</td>
<td>How sample size was determined and, when applicable, explanation of any interim analyses and stopping rules.</td>
</tr>
<tr>
<td>Sample size</td>
<td>7</td>
<td></td>
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</table>

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<tr>
<th>Paper section and topic</th>
<th>Item</th>
<th>Descriptor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Randomization – sequence generation</td>
<td>8</td>
<td>Method used to generate the random allocation sequence, including details of any restrictions (e.g., blocking, stratification)</td>
</tr>
<tr>
<td>Randomization – allocation concealment</td>
<td>9</td>
<td>Method used to implement the random allocation sequence (e.g., numbered containers or central telephone), clarifying whether the sequence was concealed until interventions were assigned.</td>
</tr>
<tr>
<td>Randomization – implementation</td>
<td>10</td>
<td>Who generated the allocation sequence, who enrolled participants, and who assigned participants to their groups.</td>
</tr>
<tr>
<td>Blinding (masking)</td>
<td>11</td>
<td>Whether or not participants, those administering the interventions, and those assessing the outcomes were blinded to group assignment. If done, how the success of blinding was evaluated.</td>
</tr>
<tr>
<td>Statistical methods</td>
<td>12</td>
<td>Statistical methods used to compare groups for primary outcome(s); Methods for additional analyses, such as subgroup analyses and adjusted analyses.</td>
</tr>
<tr>
<td>Results</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Participant flow</td>
<td>13</td>
<td>Flow of participants through each stage (a diagram is strongly recommended). Specifically, for each group report the numbers of participants randomly assigned, receiving intended treatment, completing the study protocol, and analyzed for the primary outcome. Describe protocol deviations from study as planned, together with reasons.</td>
</tr>
<tr>
<td>Recruitment</td>
<td>14</td>
<td>Dates defining the periods of recruitment and follow-up.</td>
</tr>
<tr>
<td>Baseline data</td>
<td>15</td>
<td>Baseline demographic and clinical characteristics of each group.</td>
</tr>
<tr>
<td>Numbers analyzed</td>
<td>16</td>
<td>Number of participants (denominator) in each group included in each analysis and whether the analysis was by “intention-to-treat”. State the results in absolute numbers when feasible (e.g., 10/20, not 50%).</td>
</tr>
<tr>
<td>Outcomes and estimation</td>
<td>17</td>
<td>For each primary and secondary outcome, a summary of results for each group, and the estimated effect size and its precision (e.g., 95% confidence interval).</td>
</tr>
<tr>
<td>Ancillary analyses</td>
<td>18</td>
<td>Address multiplicity by reporting any other analyses performed, including subgroup analyses and adjusted analyses, indicating those prespecified and those exploratory.</td>
</tr>
<tr>
<td>Adverse events</td>
<td>19</td>
<td>All important adverse events or side effects in each intervention group.</td>
</tr>
<tr>
<td>Discussion</td>
<td>Interpretation</td>
<td>20</td>
</tr>
<tr>
<td>Generalizability</td>
<td>21</td>
<td>Generalizability (external validity) of the trial findings.</td>
</tr>
<tr>
<td>Overall evidence</td>
<td>22</td>
<td>General interpretation of the results in the context of current evidence.</td>
</tr>
</tbody>
</table>

From ref. [13]. For more information, visit www.consort-statement.org
Good Clinical Practice

Good clinical practice (GCP) is an international quality standard for clinical trials involving human subjects that is provided by the International Conference on Harmonization [14]. GCP guidelines include protection of human rights for subjects in clinical trials, but they also include standards on how clinical trials should be conducted and define the roles and responsibilities of clinical trial sponsors, clinical research investigators, and monitors.
There is international consensus that all studies that involve a medical or therapeutic intervention on patients must be approved by a supervising ethics committee. In some countries, this body is called institutional review board (IRB). Most ethics committees or IRBs are located at the local investigator’s hospital or institution, but there also exists a central IRB for investigators who work at smaller institutions. The main function of ethics committees or IRBs is to ensure that clinical trials are performed according to ethical standards. Major issues include full and informed consent of participating human subjects, qualification of staff, performance of clinical trials according to GCP, as well as the design of the trial. Informed consent is clearly a necessary condition for, but does not ensure, ethical conduct. The final objective is to serve the community of patients, or future patients, in a best possible and most responsible way.

Ethical difficulties can arise when treatments have been established over time, even if their evidence has never been clearly documented in a well-conducted trial. This is, for example, the situation in steroid treatment for acute hearing loss with tinnitus [15]. Placebo-controlled randomized clinical trials are required in order to investigate whether steroids have a beneficial effect or not in this group of patients. However, this would imply that patients, who are randomized to placebo treatment in such trials, would be deprived of an established and potentially effective treatment. On the other hand, as long as no placebo-controlled RCTs are available, a large number of patients all over the world receive a treatment with unknown efficacy.

Specifics of Tinnitus Trials

In the design of clinical trials for tinnitus, several aspects are of specific importance and are therefore discussed in detail. These include heterogeneity of diagnoses, duration of illness, and outcome assessment.

Study Population – Tinnitus Subtypes

An important aspect in the design of clinical trials is the establishment of criteria to stratify or classify tinnitus patients. Tinnitus can occur as a result of insults to the ear, such as from noise exposure or administration of specific pharmacologic agents. It can also be caused by ear or head injuries, some diseases of the ear, and ear infections. In some cases, the causative agent remains unknown. Moreover, the manifestation of tinnitus can vary, ranging from intermittent tinnitus perception with little impact on daily life to a devastating roar that occurs 24 h a day preventing sleep and the ability to do intellectual work, leading to social isolation. Tinnitus is also often associated with other symptoms, such as hyperacusis and distortion of sounds. Affective disorders such as anxiety, phonophobia, and depression often accompany severe tinnitus and that can lead to suicide. With such differences in etiology and symptoms, heterogeneity within tinnitus patients is expected. If a treatment would exist that suppresses all forms of tinnitus irrespective of its etiology or its clinical characteristics, defining statistical analysis methods are performed according to a properly planned study design. Registration of clinical trials further allows controlling for publication bias (negative studies have a higher risk to be never published) and facilitates coordination between efforts of different research groups. Several public clinical trial registries exist; the largest of them is www.ClinicalTrials.gov that is run by the United States National Library of Medicine (NLM) at the National Institutes of Health, currently holding registrations from over 60,000 trials from more than 150 countries in the world. In addition, it has become good research practice to publish short versions of study protocols before starting a clinical trial [16, 17].

Trial Registration and Publication of Study Protocols

Public registration of clinical trials has been introduced with the primary purpose of improving public access to clinical trials where individuals with serious diseases and conditions might find experimental treatments. Moreover, registration of clinical trials became an important methodological requirement since it ensures sample sizes, primary outcome measures, and
subgroups would not be necessary. However, analysis of available clinical trials reveals a high variability in treatment outcomes, which is most probably due to the fact that patients included in a clinical trial suffer from different forms of tinnitus [18]. The fact that a subgroup of patients who have intermittent tinnitus that sounds like a typewriter, popcorn, or ear clicking receive significant benefit from carbamazepine [19, 20] indicates that “subtyping” tinnitus is highly recommended.

Two different strategies can be used to account for the inhomogeneity of tinnitus. Either a more homogeneous group is created by selective inclusion criteria or different groups within a large sample are stratified according to specific criteria. Criteria for selection or stratification have to be identified in both strategies. Among the criteria that have been suggested for delineating different forms of tinnitus are objective versus subjective tinnitus, perceptual characteristics such as pulsatile versus nonpulsatile tinnitus, the perceived localization, the duration, the frequency composition, the response to specific interventions (lidocaine, sound stimulation, somatic maneuvers), or co-morbidities such as hearing loss or concurrent psychiatric symptoms. However, there is no generally established classification system for the different forms of tinnitus to date, and there is an urgent need for further research in order to identify classification criteria. In addition to clinical and audiologic aspects, new neuroimaging methods may provide further information (see Chap. 17). Also, the identification of subgroups by cluster analysis of different clinical variables has been proposed [21].

However, all efforts of stratifying tinnitus patients into subgroups is self-limited by the amount of available patients and the time needed to recruit them. The use of very selective inclusion criteria has the consequence that results cannot be generalized to other forms of tinnitus. Therefore, defining subgroups is always a compromise between homogeneity on one side and sample size and generalizability on the other side. However, the chances for significant results in clinical treatment trials increase with homogeneous samples, e.g., when known predictor variables for a specific treatment are considered in the design of a clinical trial. Frequently, at the time of the design of a clinical trial, it is not yet known which subgroup of tinnitus patients may benefit best from treatment. A practical approach in this situation is a standardized assessment of potentially relevant clinical characteristics and a post hoc analysis of responder groups. As is later described, such analyses have to be interpreted carefully. They always have to be considered as exploratory and require confirmation in further studies.

Patients who undergo any additional tinnitus-specific treatment should be excluded from a clinical trial in order to minimize potential confounding factors that may influence treatment outcome. This may be difficult for long-term treatments such as hearing aids or white-noise generators. At least the use of these devices should be documented. An additional strategy to address this issue is the documentation of baseline stability by repeated tinnitus assessments before initiation of the intervention under study.

Also, relevant comorbid disorders like depression or anxiety may be excluded from study participation, except when the treatment to be tested is specifically focusing on this given comorbidity in tinnitus (e.g., antidepressants in tinnitus patients with comorbid depressive symptoms). As an example for the significance of defining subgroups for clinical trials, we will discuss the relevance of tinnitus duration in more detail.

**Duration of Tinnitus**

Acute and chronic forms of tinnitus differ in many respects. First of all, pathological mechanisms are likely to be different. Acute tinnitus is frequently accompanied by acute hearing loss. It is expected that an intervention that can improve hearing function will also have a beneficial effect on tinnitus. Examples of such interventions are steroid administration or treatment with hyperbaric oxygen. However, it is expected that an intervention that improves tinnitus by restoring hearing after acute hearing loss only has a beneficial effect in acute tinnitus with acute hearing loss and not in chronic tinnitus. For such an intervention, it is important to establish the therapeutic window for successful administration.

In general, tinnitus that persists more than 6 months is considered chronic [16]. However, this distinction is arbitrary and is based on national definitions, not on pathophysiologic knowledge. For example, in a current study, subacute tinnitus is defined as having between 3 and 12 months duration (see clinical trials...
Recent neuroimaging data indicate that there may be different stages of chronification over the course of several years, which differ in their pathophysiology [22], further underlining the relevance of tinnitus duration in addition to the distinction between acute and chronic.

Whereas initiation of tinnitus may be triggered by pathologies in the inner ear, the relevant pathological changes of tinnitus chronification take place within the central nervous system. Alone, this makes it obvious that the success of specific treatment interventions will depend on tinnitus duration. Therefore, acute and chronic forms of tinnitus should not be included in the same treatment trial. Otherwise, the chance of a nonresponding subgroup is artificially increased. Another problem in the acute phase of tinnitus is the high spontaneous recovery rate. Until now, there were no reliable predictors of spontaneous remission. Therefore, large sample sizes were needed for detecting significant differences between an intervention and placebo. Furthermore, if tinnitus is only of transient and mild character, the drop-out rate may be increased leading to statistical difficulties. On the other hand, for some interventions, there may be a short therapeutic window and treatment in general may become more difficult with increasing tinnitus duration [11]. Hence, there is a need for clinical trials that investigate treatment interventions both at early stages of tinnitus and in chronic tinnitus. However, in the design of a clinical trial, the duration of tinnitus should be considered as an important criterion. How this is reflected in the design of the clinical trial, e.g., by selection or stratification of the study population, depends on the intervention and its assumed mechanism.

**Outcome Measures**

Since tinnitus is a purely subjective phenomenon, assessment of treatment effects is not trivial. At the same time, the use of an adequate outcome is probably the single most important factor in the design of clinical trials. One possibility for tinnitus measurement is the assessment of tinnitus loudness, either by a visual analogue scales (VAS) or by matching or masking methods. However, psychoacoustic methods like loudness matching or minimal masking level are subjective methods and can give only indirect approximations of tinnitus intensity. Furthermore, assessments of tinnitus loudness have shown only limited reliability, and there is only a poor correlation between the intensity of the tinnitus as qualified by matching techniques and the degree of annoyance the tinnitus creates [23]. Hence, evaluating treatment effects on tinnitus should rather focus on tinnitus associated with suffering than on tinnitus loudness alone. For the assessment of tinnitus severity, there are several validated questionnaires available (see Chap. 47). However, most of the questionnaires for assessment of tinnitus severity have been designed and validated for diagnostic purposes in order to discriminate subgroups (e.g., to separate mild from severely affected tinnitus patients [24]) but not for evaluating treatment-induced changes. Thus, the available questionnaires are not specifically sensitive for the assessment of treatment-related changes in tinnitus severity. Furthermore, it is not clear which change in these questionnaires is of clinical relevance [25]. It should also be noted that most of the questionnaires have been validated by using the Beck Depression Inventory (BDI), and therefore their scores correlate highly with the BDI scores [26]. Hence, in a sample of tinnitus patients with comorbid depression, an intervention that has an antidepressant effect, but no effect on tinnitus, would probably result in reduced tinnitus scores, just by reducing depressive symptoms.

Efforts are underway in order to design specific questionnaires to evaluate treatment-induced changes in tinnitus [24]. Currently, there is consensus that until such an evaluative questionnaire is validated in different languages and internationally established, the use of one of the available validated questionnaires for the assessment of tinnitus severity is the most appropriate outcome measurement. There is widespread recognition that consistency between research centres, in how intervention outcomes are measured, would allow better comparability of different trials. At the first Tinnitus Research Initiative meeting held in Regensburg in July 2006, which gathered worldwide tinnitus experts, an attempt was made to establish a consensus both for patient assessments and for outcome measurements [2]. There was an agreement that the questionnaire most widely used and validated in most languages is the tinnitus handicap inventory [27], which should for the sake of comparability be included in every trial. It should also be
noted that the first Phase III trials currently performed for a tinnitus drug use the TBF-12 as main outcome criterion, which is essentially a short version of the THI with selection of 12 sensitive items out of the 25 items of the THI [28]. If the trials are successful and the drug is approved by the U.S. Food and Drug Association (FDA) and the European Medicines Agency (EMEA), the TBF-12 might become a reference for further pharmacologic trials.

In this context, it should also be emphasized that each clinical trial should have only one primary outcome measure – a validated questionnaire. The primary outcome measure has to be defined a priori (i.e., before the trial starts) and is the main criterion for statistical determination of the efficacy of the treatment under trial. Additional measures may be included as secondary outcome measures (e.g., assessing change in depressive symptoms using the Beck Depression Inventory [29]).

**Outlook and New Challenges**

Taken together, although efforts in finding effective treatment strategies for tinnitus are increasing, the situation in the therapeutic daily routine is disappointing. Hence, there is a big need to facilitate clinical research in order to find new effective treatment options. Major tasks will be defining and better characterization of tinnitus subgroups based on clinical symptomatology and treatment response as well as detecting predictors for therapy response. To achieve these goals, a large number of tinnitus patients have to be investigated and systematically included in clinical trials. If these investigations were performed in a standardized way, data could be pooled for analysis, which would significantly increase statistical power and enable detection of potential predictors. Unfortunately, studies currently conducted are very heterogeneous, with respect to quality of the design and outcome measures used, thereby jeopardizing the comparability of the results. Hence, future research may be very much facilitated, if there is an overall consensus about key diagnostic and outcome measures and the will to share these results for analyses via an international tinnitus database. Promising attempts have been made to find such agreements [2], which would be the basis for internationally acting research networks.

**References**

8. Tyler, RS, Noble, W, Coelho, C Considerations for the design of clinical trials for tinnitus Acta Otolaryngol Suppl, 2006; 556:44–9
16. Landgrebe, M, Binder, H, Koller, M, Eberl, Y, Kleing, T, Eichhammer, P, et al Design of a placebo-controlled,


Part II

Tinnitus Seen by Different Specialties
Chapter 23
The Otolaryngologist

Tobias Kleinjung

Keypoints

1. This chapter describes the role of the otolaryngologist (ENT specialist) in the diagnosis and treatment of tinnitus.
2. Apart from the general practitioner, the otolaryngologist is the first point of contact for many tinnitus patients.
3. Otological diagnosis must be performed in patients with acute tinnitus and basic audiological screening must be arranged.
4. The acute treatment of new-onset tinnitus is also the domain of the otolaryngologist.
5. In patients with chronic tinnitus, the role of the otolaryngologist – ideally as part of a multidisciplinary team – is to coordinate further diagnostic and therapeutic measures.

Keywords Otolaryngology • Tinnitus • Hearing loss • Counseling

Introduction and History

Advances in science in recent decades have completely redefined the role of the otolaryngologist in terms of the diagnosis and treatment of tinnitus. In the past, the otolaryngologist was often working alone in managing patients with tinnitus. The spectrum of therapeutic options available to the otolaryngologist was soon exhausted, particularly when treating patients with chronic tinnitus with no underlying otological cause. The great suffering experienced by these patients prompted committed otolaryngologists to repeatedly undertake heroic, but usually frustrated curative efforts, even extending as far as sectioning the eighth cranial nerve [1]. A wide variety of therapeutic approaches, all of which had the labyrinth as their focus, also failed to yield successful outcomes [2]. The justifiable demands of patients for further help led in many cases to a profound disturbance of the doctor–patient relationship. “You must learn to live with your tinnitus” – a comment frequently heard from the lips of doctors – was tantamount to admitting further treatment attempts would be futile. The frustrated patient often looked for a new doctor or sought refuge in paramedical treatments. The recognition that mechanisms unfolding outside the ear are key factors in the etiology and perception of tinnitus brought with it a change in the management of tinnitus patients. It became clear that to concentrate solely on the labyrinth cannot do justice to the problem of tinnitus. The development of the neurophysiological model of tinnitus [3] showed how successful treatment strategies can be designed that are outside the core competency area of the otolaryngologist. In patients with chronic tinnitus, the brain has now become the focus of treatment attempts. Aside from diagnostic and exploratory measures, the role of the otolaryngologist – as part of a team – is to coordinate the treatment of patients with tinnitus in conjunction with specialists from other disciplines.
The Role of the Otolaryngologist in a Modern Tinnitus Clinic

Patients with new-onset tinnitus generally seek an appointment first with an otolaryngologist. Thus, it is an important part of the otolaryngologist’s role to be the primary point of contact for the tinnitus patient. In Germany, tinnitus patients make up a large proportion (20–25%) of people seeking medical help from an otolaryngology clinic [4]. For this reason, every otolaryngologist in clinical practice should acquire competence in the management of tinnitus patients; as a rule, such competence should extend beyond the knowledge gained in the course of specialist training, which tends to focus on surgical treatments. The otolaryngologist should acquire the understanding of the pathophysiology of tinnitus and the capacity to empathize with the patient. During the initial consultation, it is important to collect information about circumstances relating to the onset of tinnitus, the nature of the tinnitus, any possible concomitant hearing loss, as well as the patient’s psychosocial background (see Chap. 47). The importance of this first contact with the patient is immeasurable. The needs of an otherwise healthy patient with the symptom “tinnitus” must be taken seriously, but every effort must be made to keep the patient from being overly focused on this symptom. Catastrophic statements such as “Your tinnitus might be a sign of a brain tumor” may have devastating consequences for the patient’s further clinical course.

Otolological diagnosis is a core competency area for the otolaryngologist (see Chap. 48). The purpose of otological diagnosis is to identify potential diseases of the external ear, middle ear, or inner ear that might be a possible cause of tinnitus. In many patients, this may result in a straightforward therapy, such as the removal of wax from the ear canal. For other diseases, such as otosclerosis, surgery (see Chap. 83) may lead to the abolition of tinnitus. In most tinnitus patients, however, an otological examination will reveal no abnormal findings.

Otolological diagnosis should always be followed by audiological testing, which can discriminate between different forms of hearing loss. Such information should enable the otolaryngologist to explain to the patient how the tinnitus might have developed. This can help many tinnitus patients cope with their tinnitus and perhaps require no further therapy.

For patients who need further therapy, the otolaryngologist should direct the patient to an appropriate specialist in areas, such as the temporomandibular joint, cervical spine, etc., for further diagnostic work up.

Once the results of all investigations have been completed by all the specialists involved, a team conference should be held to draw up a treatment strategy for the individual patient. The interdisciplinary approach permits personalized treatment strategies that can be tailored to the requirements of the individual patient. Even though many forms of tinnitus remain “incurable” in the classic sense, this modus operandi brings a higher degree of satisfaction both for patients and for the physicians treating them.

Tinnitus that occurs as a part of sudden hearing loss should be treated with appropriate medications, such as intravenous steroid therapy (see Chap. 56) and rheologically active medication [5]. The usefulness of such vasoactive infusion therapies is still under debate in the English-speaking world [2].

The otolaryngologist can also make important contributions on a scientific level. As the first point of contact, the otolaryngologist will be familiar with many different patients and can therefore be useful in the recruitment, follow-up, and the assessment of patients in clinical studies.

References

Chapter 24
The Role of the Audiologist in Tinnitus Practice

Grant D. Searchfield and David M. Baguley

Keypoints

1. Audiologists play a significant role in most models of tinnitus health care provision, including both the assessment and management of tinnitus as reported by Henry et al. (Am J Audiol 14:49–70, 2005; Am J Audiol 14:21–48, 2005).

2. Audiologists have expertise assessing auditory function; training in auditory physiology and psychology prepares them to provide tinnitus counseling; and they are able to fit hearing aids and other instruments for tinnitus therapies.

3. In some situations, the Audiologist will be part of a multidisciplinary team (e.g., in a large metropolitan hospital) that may potentially include Otologists, Neurologists, Hearing Therapists, and Psychologists.

4. In other circumstances, the Audiologist may work in comparative isolation and be responsible for the majority of tinnitus care.

5. In this chapter, the authors consider the Audiologists’ perspectives of tinnitus.

6. We describe Audiologists’ skills and attributes in their role in tinnitus management; present models for tinnitus practice; and introduce a Matrix framework from within which clinicians can choose strategies for patients with varying needs.

Keywords Tinnitus • Audiology • Assessment • Rehabilitation

What is an Audiologist?

Audiologists are professionals trained in the clinical application of hearing science. Audiology exists as a health care profession in English speaking countries, the Americas, and the Pacific Rim; elsewhere similar roles are undertaken by medically qualified professionals or technicians [1]. Henceforth, in this chapter, we use the name “Audiologist” to identify professionals with nonmedical University qualifications in audiology. However, much of what we discuss could equally apply to Audiological Physicians, Hearing aid Acousticians, and others who provide nonmedical assessment and management of tinnitus.

Audiologists have a broad scope of practice encompassing most aspects of hearing assessment and management. This practice includes behavioral and electrophysiological evaluations of hearing, rehabilitation of hearing loss through technology (hearing aids, cochlear implants), hearing loss prevention, and assessment and management of balance and tinnitus. Due to the strong association between hearing loss and tinnitus (see Chaps. 34 and 35), it is not surprising that most Audiological associations or registration bodies recognize that tinnitus is at the core of audiology practice.

“Audiologists are qualified to evaluate, diagnose, develop management strategies, and provide treatment and rehabilitation for tinnitus patients” [2].

Audiology began to develop as a distinct profession after World War II [3]. Shortly after this, audiological methodology (such as hearing aids) began to be applied for treating tinnitus [4, 5]. The use of sound as a treatment medium became more common with the development of ear-level maskers in the 1970s and the use of this technology in newly developed tinnitus clinics [4, 6]. The profile of audiology in tinnitus management rose
again in the 1990s with the widespread adoption of Tinnitus Retraining Therapy (TRT, [7]). The current decade has seen the continued development and diversification of tinnitus management methods available to audiologists. Some notable additions to the audiologists’ armory being Tinnitus Activities Treatment [8], Audiologic Tinnitus Management [9, 10], Neuromonics [11], and modified versions of TRT [12]. As the scale of tinnitus worldwide has become apparent, Hearing aid manufacturers and technology companies have also become more involved in improving management tools. A comprehensive survey of hearing health care practice internationally identified that audiologists are responsible for tinnitus management in most countries [1] .

**Audiology Skills Applied to Tinnitus**

The skills that audiologists acquire during their training and practice of “general” audiology are directly applicable to the more specialized area of tinnitus (Table 24.1). For example, audiologists need to counsel anxious hearing aid candidates and provide support to emotional parents on diagnosis of hearing loss in children. This counseling is not too dissimilar to counseling the distressed tinnitus sufferer.

Most educational programs in audiology offer limited training specific to tinnitus [10], but there are many regular opportunities for audiologists to gain further tinnitus knowledge. Scientific meetings, such as the International Tinnitus Seminars and Tinnitus Research Initiative meetings, are excellent opportunities to learn of the latest scientific developments in the field. Annual training workshops, such as one hosted annually by Iowa University as well as the European Tinnitus Course in Cambridge, build on existing knowledge to provide additional skills toward tinnitus practice. Audiologists should also have sufficient training to implement practice models described in this book and in other publications [13, 14]. Many established tinnitus clinics (including those at Addenbrooke’s Hospital and The University of Auckland) are willing to share experiences and clinical protocols with clinicians new to the field.

**Diagnosis and Assessment**

Given that patients with hearing and balance issues are internationally being referred directly to an audiologist, care should be taken to ensure effective and efficient diagnosis. This will usually involve four themes:

**Table 24.1** Consideration of audiological skills in general use and as applied to tinnitus. American Speech and Hearing Association audiology scope of practice guidelines [56] were used to formulate the categories of audiology practice

<table>
<thead>
<tr>
<th>General audiology</th>
<th>Tinnitus audiology</th>
</tr>
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<tbody>
<tr>
<td>Hearing loss prevention and promotion</td>
<td>Tinnitus prevention and promotion</td>
</tr>
<tr>
<td>Identification of auditory and balance disorders</td>
<td>Identification of tinnitus and tinnitus related pathology</td>
</tr>
<tr>
<td>Behavioral and electrophysiological assessment of auditory function</td>
<td>Behavioral and electrophysiological assessment of tinnitus</td>
</tr>
<tr>
<td>Intraoperative monitoring of audiology function</td>
<td>Intraoperative monitoring of audiology function during tinnitus-related neurosurgery</td>
</tr>
<tr>
<td>Assessment and management of Auditory Processing Disorders</td>
<td>Evaluation and intervention of tinnitus (and Hyperacusis) in recovery from head and neck trauma</td>
</tr>
<tr>
<td>Otoscopy and middle ear function tests examining for the obstruction of external auditory meatus and middle ear pathology</td>
<td>Evaluation of potential contribution of external and middle ears to tinnitus symptoms.</td>
</tr>
<tr>
<td>Assessing the “Hearing needs” of patients</td>
<td>Tinnitus needs assessment</td>
</tr>
<tr>
<td>Referral and consultation with other professionals</td>
<td>Referral and consultation with other professionals</td>
</tr>
<tr>
<td>Development of intervention and rehabilitation plans</td>
<td>Tinnitus management plan</td>
</tr>
<tr>
<td>Select and fit hearing aids and/or assistive devices to improve hearing</td>
<td>Select and fit hearing aids and/or sound generating devices for tinnitus management</td>
</tr>
<tr>
<td>Assessment and management of severe-profound hearing loss with Cochlear Implants</td>
<td>Assessment and management of tinnitus accompanying severe-profound hearing loss with Cochlear Implants</td>
</tr>
<tr>
<td>Use counseling to address psychosocial aspects of hearing loss and provide communication skills</td>
<td>Tinnitus counseling</td>
</tr>
<tr>
<td>Assessment of hearing intervention outcomes</td>
<td>Tinnitus outcomes measurement</td>
</tr>
</tbody>
</table>
Identification of treatable otological pathology
Assessment of hearing and tinnitus testing
Assessment of tinnitus handicap
Identification of treatable psychological symptoms, such as anxiety and depression

Each of these themes are now discussed in turn. Given appropriate teaching and support, there is no reason why an audiologist should not diagnose otological pathology. A protocol approach is advocated, wherein investigations and onward referral are indicated by the presence of certain symptoms or test findings. An example is in centers located in the UK (Cambridge and Liverpool for example) where patients with unilateral tinnitus and/or asymmetric hearing thresholds are referred for Magnetic Resonance Scanning by the audiologist leading the Tinnitus Clinic. Similarly, when an autoimmune hearing loss is suspected, appropriate serological tests can be requested. When abnormalities are found an otological opinion is then necessary. This extension to the traditional audiologist scope of practice allows a direct-access model of service provision that is both efficient and cost effective.

Audiometric testing is clearly an essential element of the assessment of the tinnitus patient, and in many patients tympanometry will also be routinely undertaken. The issue of testing for loudness tolerance in the tinnitus population, many of whom may have hyperacusis, is somewhat controversial and discussed in Chap. 3. There are well-established audiometric methods of tinnitus pitch and intensity matching (see Chap. 24). The clinical utility of such measures is not high, however. Electrophysiological measures of tinnitus are being developed and opportunities may exist for audiologists to implement these in clinical practice.

The assessment of tinnitus handicap is another essential element in patient management. Many questionnaire instruments are available to determine the impact of tinnitus. These questionnaires typically inquire as to the effects of tinnitus on work or leisure activities, sleep emotion, and in some cases hearing [15, 16]. The clinician must choose one or two questionnaires that are reliable, sensitive to change with treatment, and of low impact to the patient. The need for a universal outcome measure for tinnitus treatments has been recognized [17]. Whether clinicians are willing to move from existing questionnaires to a standard index will be tested in the next few years.

The assessment of anxiety and depression is also strongly indicated in tinnitus patient management. The Beck Depression Inventory [18] and State-Trait Anxiety Inventory [19] are the gold standard in this regard, but the questionnaires may alarm some individuals, promoting greater distress. The Hospital Anxiety and Depression Scale [20] is in widespread use in tinnitus clinics in the UK, and is a low impact screen for these symptoms. As psychologists developed the scale, it is credible to that community which helps onward referral.

Management

Audiology-based methods of tinnitus intervention demonstrate their benefits across psychological and audiological/neurophysiological domains, including but not limited to attention, habituation, and learning [21]. Most strategies used by audiologists incorporate the use of sound making devices along with counseling [22] (see Chap. 74). Although it has been argued that the use of sound stimulation has limited benefit over counseling [23] and even that it is counterproductive [24], there is increasing evidence that placed in an appropriate counseling framework sound does provide additional assistance [11, 25]. How sound should be used and with what counseling approach is most appropriate has been a source of considerable debate [26, 27]. Although treatment strategies used by audiologists differ, fundamentally they are actually very similar – sound therapy and counseling. The most appropriate intervention used by audiologists should be governed by the needs of the individual seeking help. At a bare minimum, an audiologist should be able to offer positive advice and refer to other clinicians involved in tinnitus management. The major elements of tinnitus management from an audiology perspective are described below.

Tinnitus Needs Assessment

A key stage in preparing an audiological management plan is determining the needs of the individual. This tinnitus needs assessment attempts to identify
how the tinnitus affects an individual. Using tinnitus questionnaires can help identify general emotional and lifestyle needs of patients. The Client-Orientated Scale of Improvement (COSI, [28]) is a widely used needs assessment tool in hearing aid selection. The COSI-Tinnitus [29] can be used to identify an individual’s specific needs and goals for tinnitus management. Through dialog and acknowledgment of the tinnitus sufferer’s complaints the groundwork for counseling can be laid. The patient’s needs can be addressed through variations of counseling and sound therapy.

**Audiological Interventions**

Specific counseling (see Chap. 70), masking (see Chap. 74), and habitation (see Chap. 75) based treatments are mentioned elsewhere in the book. We describe them briefly here considering the role of Audiologists.

**Counseling**

To help patients understand tinnitus and facilitate their coping with the condition, clinical approaches to the management of tinnitus include the use of counseling (see Chap. 70). In this context, audiology counseling interventions can range from simply providing advice, to bibliotherapy [30], directive counseling [31], and psychoeducation [32]. In some cases, referral for formal psychological assessment and treatment will be indicated, e.g., CBT [33] (see Chap. 54), though some patients may be resistant to that [34]. Audiologists should be able to address patient concerns and misplaced beliefs due to broad knowledge of auditory physiology, psychology, and aural rehabilitation. Counseling accompanying the fitting of sound devices would be very similar to the counseling of an audiologist should provide individuals with hearing aids in a comprehensive aural rehabilitation program. Some audiologists use CBT-based techniques, which address a patient’s reaction to tinnitus, and provide relaxation training and cognitive restructuring as part of their scope of practice (e.g., [36]).

**Masking**

Tinnitus masking uses sound to cover tinnitus to some degree and should be used with counseling [6]. Masking could be considered the core audiological-based treatment for tinnitus [37]. During the 1990s, masking became somewhat maligned as a treatment method, but the principles and clinical application still remains a useful tool for audiologists. Masking is commonly associated with the use of ear-level devices produced by hearing aid manufacturers with whom audiologists should have strong working relationships. Audiologists familiar with tonal masking in audiometry should understand that tinnitus masking does not obey normal peripheral masking rules [38, 39]. Tinnitus masking is likely due to central processing mechanisms, similar perhaps to informational masking [40]. Complex sounds may be more useful in this treatment than constant broadband noise commonly in use [41].

**Habitation**

Habitation is the decline in responses to a signal that is not important [42]. The most well-known clinical models of tinnitus habitation are the model of Hallam et al. [43] and Tinnitus Retraining Therapy [7]. Audiologists have tended to gravitate toward the TRT model due to familiarity with; the auditory system as expressed in the underlying model [44]; directive counseling and the use of instruments for sound therapy [45]. TRT has been simplified to suit different clinical settings [12] and other published management protocols exploit the aspects of habitation [8].

**Sound Therapy Technology**

Audiology has a strong technology focus; this is also the case in its tinnitus role.

Although tinnitus patients often crave silence, this silence can “feed” the tinnitus by increasing the tinnitus signal relative to background noise. With little competition the auditory system will naturally divert attention resources to the remaining signal – tinnitus [48]. While avoiding silence is simple advice, it does
help patients who fail to make the connection between tinnitus perception and background noise levels. Desktop sound generating devices produce a variety of different sounds (for example, ocean waves, rain, and running water), and these have been found useful to reduce tinnitus effects at night [49]. Nighttime is often when tinnitus sufferers experience heightened tinnitus awareness due to low sound levels and absence of other competing sensory input. Desktop devices are available through hearing aid distributors or electronics retailers.

The value of digital music players (e.g., MP3 players) as tinnitus aids have been recognized by both patients and audiologists [9]. Sounds can be produced by computer programs in the clinic or downloaded from the Internet. Use of prerecorded sounds is an easy way of obtaining treatment sounds that patients find comfortable and easy to listen to. Music can be used in an informal way to promote positive emotional effects to reduce tinnitus [50]. The Neuromonics treatment [11] uses music in a customized form as part of its audiology-focused treatment protocol. Although each year the size of digital players decreases and their battery capacity increases, they are still less convenient to wear on a regular basis than hearing aid style sound generators [52]. Personal music players also do not address any accompanying hearing handicap the way hearing aids or combination devices do.

Hearing aid style in-the-ear and behind-the-ear sound generators produce noise stimuli of variable intensity and frequency. The selection and manufacture of these devices is much the same as for hearing aids, but without the sophistication of signal processing necessary for amplification. Hearing aids themselves are often an underrated tinnitus management technology. Detailed protocols for the fitting of hearing aids in tinnitus treatment are available [29] (Chap. 74). The development of more advanced signal processing appears to have increased success rates [29, 53]. Audiologists are intimately familiar with hearing aids, including their selection, electroacoustic, and subjective evaluation of performance. Hearing aids have the benefit of addressing hearing as well as tinnitus needs. Combination instruments combine a hearing aid with a built-in tinnitus masker. They are available from a limited number of hearing aid manufacturers, but they attempt to combine the benefits of amplification with generated sound [54]. The technology, in these devices, has in the past lagged behind that of the best hearing aids. However, this technology gap appears to being addressed by some manufacturers. A potential advantage of combination devices is their independence from environmental sound levels for effect. In persons with severe to profound hearing loss, cochlear implants become a management option [55].

Matrix Approach to Therapy Selection

Several authors have suggested a progressive management approach based on tinnitus severity and how it manifests itself [14, 56]. One approach to therapy selection is to use a Matrix model in which the audiologist selects intervention depending on the needs of the individual. Selecting the most appropriate elements from both psychological and technological axes can target individual needs assessed during an interview. The key to successfully implementing such an approach is to understand the problems the patient reports and their reaction to each treatment element. Individuals with high emotional needs are provided with more in depth counseling. Those with greater complexity of auditory injury may require more complex technological solutions. The potential strengths of audiologists in this management role are their ability to work across both technology and counseling strategies (Fig. 24.1).

Referring on

Within the context of an audiologist led Tinnitus Service, there will be patients who need to access professionals of other disciplines when issues arise that are beyond the scope of the audiologist’s practice. The majority of these will be referrals to Otology and Psychology services, but there may also be occasions when referral to disciplines, such as Hearing Therapy (for addition counseling), Neurology (e.g., Head injury), Maxillofacial surgery (e.g., temporo-mandibular joint assessment), or Physiotherapy (for one-to-one instruction in relaxation techniques) may be indicated. Building relationships with professionals of such disciplines is an essential part of developing a Tinnitus Clinic. The ease with which these relationships are formed will vary with context and may be most straightforward in a University Acute Hospital
setting. Outside the hospital setting, audiologists need to seek out professionals to network within a useful way (Fig. 24.2).

A clear framework for which issues lie within, and without, the boundaries of clinical practice for an audiologist has been developed by Flasher and Fogle. The areas that are considered to lie within and outside boundaries are detailed in Table 24.2. This framework deserves some reflection. The issues said to be within need to understand the cause and neurophysiological basis of tinnitus — directive counseling is one way of providing this. (c) Patients with significant anxiety, depression, and hearing loss are likely to be best served by hearing aids (or combination instruments) along with referral to psychology or psychiatry services.

Table 24.2  Audiology counseling scope of practice and referral guidelines, based on Flasher and Fogle

<table>
<thead>
<tr>
<th>Within scope of practice</th>
<th>Referral</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interviewing the patient/family</td>
<td>Chemical dependence</td>
</tr>
<tr>
<td>Presenting the diagnosis</td>
<td>Child or elder abuse</td>
</tr>
<tr>
<td>Providing information about the diagnosis</td>
<td>Chronic depression</td>
</tr>
<tr>
<td>Discussing interventions for the diagnosis</td>
<td>Legal conflicts</td>
</tr>
<tr>
<td>Dealing with the patient’s reaction to the diagnosis</td>
<td>Marital problems</td>
</tr>
<tr>
<td>Supporting the strengths of the person and their efforts to regain function</td>
<td>Personality disorders</td>
</tr>
<tr>
<td>Supporting the strengths of the family to help them interact optimally with the patient</td>
<td>Sexual abuse and sexual problems</td>
</tr>
<tr>
<td>Creating supportive empowerment for the patient and family to develop the ability to manage their own problems and be independent of the clinician</td>
<td>Suicidal ideation</td>
</tr>
</tbody>
</table>

Fig. 24.2  Hub and spoke model of key relationships from an audiologist led service
Summary and conclusions

Audiologists should feel that they have the skills to implement many of the tinnitus treatments in this book. Useful protocols for managing tinnitus have also been published elsewhere [32]. Increasingly, audiologists are adopting evidence-based practice models [55]. Tinnitus practice models should also reflect current evidence and audiologists should adapt their methods based on the evidence available. Audiologists should have the training and flexibility to incorporate changes in the understanding of physiology, psychology, assessment, and management technologies as they occur. In many respects, audiology practice is one of human-technology interaction. Rapid advancement in sound technology applications and hearing instruments should see audiologists among those at the forefront of tinnitus treatment innovation for the foreseeable future.

References

Keypoints

1. Tinnitus is always both a medical and a psychological phenomenon.
2. A medical condition might be responsible for the emergence of tinnitus, but psychological factors play an important role in individual processing of inner noises.
3. Characteristics of tinnitus-like loudness do not determine the tinnitus-related distress.
4. The primary goal of psychological interventions is to promote habituation and to improve the patient’s ability to reduce the impact of tinnitus on the quality of life.
5. Psychological approaches offer:
   (a) Diagnostic assessment
   (b) Management of tinnitus
6. Psychological interventions should be an integral part of tinnitus management and not be made dependent on existence of a mental disorder.
7. Early referrals to a psychologist are desirable to conduct a thorough assessment of tinnitus-related complaints and to undertake a comprehensive functional analysis of the problem.

Keywords Tinnitus • Psychological assessment • Psychoeducation • Psychological treatment • Multiprofessional team

The Psychological Perspective on Tinnitus

Tinnitus always has to be regarded as being both a medical and a psychological phenomenon. Even if there is a medical reason for the emergence of tinnitus (e.g., hair cell damage), it is the brain that generates the inner noise when interpreting an altered pattern of nerve signals. This “abnormal” perception is further processed by the brain, and then psychological factors come in to play an important role regarding how the tinnitus is evaluated and coped with. Nevertheless, when proposing a “psychological dimension” of tinnitus, it does not mean that tinnitus is a mental disorder. To classify patients with tinnitus on the basis of hypothesized underlying medical conditions as “organic” or “nonorganic” (respectively “psychogenic”) is not reasonable either. Likewise, it is not at all advisable to attempt to modify the patient’s personality. Instead, the consequences of tinnitus (i.e., behavior and cognition regarding tinnitus) must be made the central issue in psychological assessment and intervention.

Due to the fact that tinnitus is not verifiable by any objective measurement, patients easily get the feeling that their sensations are not taken seriously. They are afraid that their symptoms are considered to be imagined, not real or feigned. For that reason, it is important to emphasize the validity of tinnitus as a sensory experience to the patients.
**Psychological Approaches: Assessment, Psychoeducation/Counseling, Psychological Treatment**

The primary goal of psychological interventions is to improve the patient’s ability to reduce the impact of tinnitus on the quality of life, i.e., to teach and improve coping strategies. Psychological approaches can offer assessment and management of tinnitus.

**Psychological Assessment**

A comprehensive assessment is essential before the implementation of therapy. Apart from medical and audiological parameters, perceptual, attentional, emotional, and behavioral aspects have to be equally considered. Topics of psychological assessment include characteristics of tinnitus (loudness, localization, pitch of sound) and the progression of tinnitus (onset, duration, intensity, increasing and decreasing factors). Beyond that, cognitive-emotional evaluation and coping (e.g., catastrophic thinking, helplessness, anger, sadness, etc.), psychological impairments related to tinnitus (depression, irritability, sleeping problems, and so on), effects of tinnitus on life (e.g., work, social interactions), sources of stress apart from tinnitus (e.g., live events, daily hassles), operant factors (e.g., avoidance behavior), comorbidity (e.g., mental disorders, hearing loss), treatment history, and treatment expectations have to be evaluated (see Table 25.1).

In addition, sometimes it might be important to disentangle connections between tinnitus and other afflictions, preventing tinnitus from becoming a scapegoat for all other problems. Assessment of tinnitus also includes the patient’s view of his/her problem: Although from a psychological perspective, the etiology of tinnitus can be neglected most of the time, the way in which patients interpret the cause for tinnitus can be essential for coping efforts. Patients may indicate that the intensity or loudness of their tinnitus causes difficulties in the areas of sleep, concentration, hearing, social relationships, or work and is therefore made responsible for their increasing anxiety and depression. However, as research shows, there appears to be little correlation between the subjective loudness of the tinnitus and the degree to which a person is impaired by it [1, 2]. Tinnitus distress cannot be regarded as dependent on the severity of the tinnitus sensation or as a function of “loudness.” A variety of features of tinnitus, together with characteristics of the individual, have to be considered in assessing tinnitus-related distress. It is the patient’s reaction to tinnitus rather than the symptom itself that separates the individual who simply “experiences” tinnitus from the individual who seeks medical or psychological help because of tinnitus [3].

Psychological assessment is accomplished through interviews, questionnaires, severity rating, and so on. Sometimes, diaries are used for documenting frequency, duration, intensity, and other parameters of tinnitus impairment. The introduction of diaries arouses fear in some patients that it might cause exacerbation of tinnitus because attention is focused on it. However, this is not harmful. On the contrary, it might rather be an opportunity to make patients aware of the alliance between attention direction and perception.

**Psychoeducation/Psychological Counseling**

An understanding of the assumed neurobiological basis of tinnitus, and its cognitive, emotional, and behavioral factors is essential for successful coping. Knowledge about tinnitus ought to be enhanced, and patients are asked to take psychological aspects of tinnitus into account. Educational programs cover topics like assumptions about the causes of tinnitus, give information about exacerbating factors and the prognosis, and give an overview of treatment possibilities, etc.

Many patients are afraid that tinnitus might become worse over time, that they will go deaf, and they consider
the disorder to be a severe illness. Such beliefs and concerns shift attention to tinnitus and increase its awareness. Patients need to be taught about the relationship between selective attention on tinnitus and its cognitive-emotional and behavioral consequences.

It is also important to inform patients that a large number of individuals are not impaired at all and are able to cope effectively with tinnitus. Poor coping has been found to be associated with the lack of control over sound and failure to habituate [4]. Besides the perception of noises, various other problems may contribute to a negative emotional status and patients may incorrectly attribute it entirely to tinnitus. In addition, patients often are worried that tinnitus becomes worse over time. However, research has shown quite the opposite: the number of complaints tends to decline the longer the tinnitus has been present [5, 6].

Instead of focusing on the unrealistic goal of tinnitus elimination, modifying the ways of coping is a more realistic goal. However, this can be challenging and involves assisting patients in the identification of aggravating factors and dealing with negative emotions (e.g., anxiety, anger, depression, etc.), sleeping problems, interference with social and recreational activities, concentration dysfunctions, and deterioration of performance.

Sometimes, patients also need help avoiding countless ineffective treatments and considerable costs.

**Psychological Treatment**

Treatment has to be tailored to the specific needs of patients, and patients should become active participants in all assessment and treatment procedures. Psychological approaches depend on the working alliance between therapist and patient rather than on so-called compliance. Psychological treatment is a collaborative effort rather than directive approach [7].

Before beginning any psychological treatment, a therapeutic rationale has to be developed and offered to the patient. A general hypothetical model of tinnitus tolerance was first developed by Hallam [8] and enhanced by Kröner-Herwig [9].

This model of tinnitus tolerance described by Hallam [10] suggests that tinnitus can be equated with any other auditory stimulus to which a person may or may not attend and that habituation to tinnitus noises and development of tolerance is the normal response, even though this process may take time. Habituation takes place when an originally new stimulus becomes “well known” and has no relevance for taking any action. Habituation fails if the stimulus is endowed with a negative evaluation (threat, impairment, anxiety). Attention to the inner noise is correlated with distress since it is associated with negative thoughts yes, but it might be German-English; and may also interfere with other activities (e.g., falling asleep or reading a book).

In this model, suffering from tinnitus is explained as a failure of habituation or adaptation. At least three classes of variables are considered influential to the process. These variables can be divided into:

- **Sensory factors**: The characteristics of the stimulus (i.e., intensity and quality).
  It is assumed that noises which are more salient and show a more variable and irregular pattern require a longer period of habituation.

- **Perceptual factors**: Environmental conditions (e.g., intensity of other stimuli and the competing demands on attention).
  For some patients, masking by natural sounds will frequently occur. Different activities and competing sensory perceptions ought to divert attention from tinnitus.

- **Psychological factors**: It is assumed that the more meaningful, especially the more threatening, a stimulus is the more attention it will receive, which creates a positive feedback loop: the more tinnitus is attended to the more the person is involved in negative cognitive emotional processing. High levels of cortical arousal are supposed to delay habituation. A patient’s style of information processing and general distractibility may influence habituation as well. Furthermore, CNS pathology affecting the neural pathways involved in attention, habituation, and appraisal has to be considered as well (see Fig 25.1).

Psychological therapies aim to assist patients in controlling attention by learning to direct attention away from tinnitus (attention-control techniques) and in bringing negative cognitive processes under self-control (cognitive restructuring techniques). Behavior modification techniques aim at reducing avoidance behavior motivated by tinnitus and increasing adaptive problem solving. In addition, different forms of relaxation
training, including biofeedback, are offered to find a way of coping with tension related to tinnitus, sleeping difficulties or other sources of stress (see Chap. 71).

**General Recommendations Regarding Treatment Protocols**

Psychological interventions should be an integral part of tinnitus management and not be based on the existence of a mental disorder, despite the fact that in some cases anxiety or depressive disorders can accompany tinnitus attributed distress. Early referrals to a psychologist are desirable to undertake an assessment of tinnitus-related complaints, identification of psychiatric comorbidity, and to undertake a comprehensive functional analysis of the problem.

Figures 25.2 and 25.3 show two different models of cooperation between medicine and psychology. Referrals to psychologists after various medical and audiological treatments have failed in removing or diminishing tinnitus are counterproductive. And simply telling the patient to accept and ignore the ringing in their ear is not enough – if it were they would have already done so. Giving the information that “nothing more can be done” and that the tinnitus might be “psychogenic” is often interpreted by patients as uncaring and insensitive. Such sentiments hamper the search for and acceptance of psychotherapeutic help (see Fig. 25.2).

McFadden [11] stated that “treatment of psychological factors without adequate preparation of the patient often results in confusion and alienation.” It is vital to inform the patient that while the tinnitus is “real,” the maladaptive response is creating the distress, and this is where patients themselves can intervene. Moreover, cognitive-behavioral therapy (CBT), albeit a primarily psychological approach, may have...
significant neurophysiological consequences, as suggested by the principles of neuroplasticity and cortical reorganization [12–14]. Resistance is often minimized when the patient recognizes that this approach works on the biological level as well.

When tinnitus is experienced, a patient will very likely consult his or her doctor first. From the very beginning physicians, audiologists, and psychologists should work together as partners (see Fig. 25.3). Medical assessments and interventions prevail at the beginning (see yellow triangle in Fig. 25.3). Psychological assessment, counseling, and treatment should become more significant over the course of time (see blue triangle in Fig. 25.3).

While medical interventions attempt to remove (“to cure”) tinnitus, psychological intervention rather supports patients in learning to tolerate the noises and handle tinnitus-related impairments (“to manage”). The patient role is very different in medical and psychological settings as well. While doctors “treat" a disease and patients are more or less passive recipients of treatment, patients have to actively participate in psychological approaches. The collaborative style maximizes patient involvement, encourages the patient to take responsibility, and minimizes the feeling that the therapist is imposing his or her own view.

The time of referral to a psychologist and psychotherapist has to be planned carefully as well. The arrows in Fig. 25.3 show different time points. The treatment management symbolized by arrow 1 shows an example where tinnitus exclusively is seen as a medical condition. Psychological factors are neglected. Patients expect that tinnitus is being removed by medical interventions. When this goal fails (which is the case in chronic tinnitus most of the time), patients increasingly get disappointed and frustrated. In response, they might visit other doctors (“doctor-shopping”) or become desperate and hopeless. The medical expert should be helpful in deciding when is the right moment for seeking the cooperation of a psychologist by assessing the cognitive-emotional and behavioral impact of tinnitus on the patient early and repeatedly. Arrow 3 characterizes quite the opposite: The patient is referred to a psychologist, without (or insufficiently) performing medical and audiological assessments. In this case, tinnitus is considered exclusively “psychogenic.” Medical and audiological factors are ignored. Both ways have their shortcomings in the long run. In any medical condition which is predisposed to become chronic, medical as well as behavioral variables have to be considered equally. It is mainly the cognitive, emotional, and behavioral response to tinnitus that separates patients experiencing the symptoms from patients who are suffering from tinnitus, and behavior can be changed at any time. Due to that, psychologists play an important role in tinnitus management regardless of the existence of psychiatric disorders.

**References**

des chronischen Tinnitus. Beltz: Weinheim
and management of tinnitus distress. In: Tinnitus 156–175
Treatments. National Academy: Washington, DC
information processing in the primary auditory cortex. Nat
Neurosci 1, 727–731
son with other sensory systems. Trends Neurosci 22, 74–80
Clin North Am 36, 249–266
Keypoints

1. Subjective tinnitus is not a disease, but a symptom and the many forms of tinnitus probably have different pathophysiology.
2. For a long time, it was believed that tinnitus arose from the ear and that the anatomical location of the physiological abnormalities which cause tinnitus was the ear. However, it was later understood that most forms of tinnitus were caused by the expression of neural plasticity.
3. The fact that most forms of tinnitus are disorders of the nervous system puts emphasis on neuroscience for treatment of tinnitus.
4. This chapter is focused on the treatment of tinnitus and how neurologists can be involved in the evaluation and diagnosis of patients with tinnitus.

Keywords

Tinnitus • Phantom perception • Neural plasticity • Auditory pathway • Neurotology

Abbreviation

TRI Tinnitus research initiative
GABA Gamma amino butyric acid
TMS Transcranial magnetic stimulation

Introduction

There are two main types of tinnitus – objective and subjective tinnitus (see Chap. 2). Objective tinnitus is caused by sounds generated in the body and then transmitted to the ear, whereas subjective tinnitus is caused by an abnormal neural activity. Objective tinnitus is rare, but subjective tinnitus is a frequent disorder that occurs with different severity; it can be just noticeable, an annoyance, or it can cause suffering that severely reduces the quality of life. There are no objective tests that can measure subjective tinnitus, and the only person who can evaluate a person’s tinnitus is the person who has the tinnitus. This is one of the aspects of subjective tinnitus that is similar to central neuropathic pain [1] (see Chap. 14).

It is generally agreed that subjective tinnitus is not a disease, but a symptom, and the many forms of tinnitus probably have different pathophysiology. For a long time, it was believed that tinnitus arose from the ear and that the anatomical location of the physiological abnormalities that cause tinnitus was the ear. However, it was later understood that most forms of tinnitus were phantom sounds [2] caused by the expression of neural plasticity [1] (Chap. 12). Realizing the complexity of tinnitus has highlighted the importance of interdisciplinary research. The fact that most forms of tinnitus are disorders of the nervous system has placed emphasis on neuroscience in the studies of tinnitus (see Chap. 10).

Objective tinnitus is caused by sound generated in the body, reaching the ear through conduction in body tissues [1] (see Chap. 10). The source can be turbulent flow of blood in an artery, where there is a constriction, or it can be caused by muscle contractions. Unlike subjective tinnitus, an observer can often hear objective tinnitus using a stethoscope.
Subjective tinnitus consists of meaningless sounds that are not associated with a physical sound, and only the person who has the tinnitus can hear it. This chapter discusses subjective tinnitus and focuses on neurologic evaluation of patients with tinnitus, describing the methodology for obtaining a clinical history to classify the tinnitus and make suggestions to the patient regarding the treatment to relieve tinnitus.

Tinnitus is a Neurological Entity

Tinnitus is considered a phantom perception similar to neuropathic pain, sharing a similar pathophysiology and clinical symptoms. Both neuropathic pain and tinnitus are perceptions that occur without the physical stimulation of receptors, which are considered to be the result of maladaptive neural plasticity (see Chaps. 12 and 34).

The expression of neural plasticity can change the balance between excitation and inhibition, promote hyperactivity, and the activation of specific parts of the nervous system not normally involved in processing sounds, such as the nonclassical auditory pathways (extralemniscal pathways) (see Chap. 8). The strongest promoter of expression of neural plasticity is the deprivation of input, which explains why tinnitus often occurs together with hearing loss [1] (see Chap. 11).

Different treatments, such as training and sound exposure to magnetic stimulation, including the use of neuromodulation, are based on the assumption that hyperactivity and reorganization of the nervous system are the causes of many of the different forms of tinnitus.

How the Clinical Neurologist is Involved in Tinnitus Diagnosis?

Individuals with mild tinnitus often do not need any treatment, but some need an assurance that their tinnitus is not a sign, therefore, of a severe disease. A detailed clinical history is essential in all patients with severe tinnitus with an aim to find an etiology if possible. A consensus at a first Tinnitus Research Initiative (TRI) meeting (www.tinnitusresearch.org) agreed upon items that should be assessed, and it has created a tinnitus questionnaire designed for assessing the severity of tinnitus and comorbidity [3].

In parallel to the otologist’s examination necessary in all patients with tinnitus, the neurologist should focus on patients who are suspected of having neurological disorders that may be involved in causing tinnitus or which occur as comorbidity to tinnitus. Examples are vascular malformations and brain tumors. Intermittent types of tinnitus may occur in individuals with migraine and epilepsy (see Chap. 61). Participation of neurologists is also important in the management of patients with somatic tinnitus that occurs without an apparent cause. A multidisciplinary approach should be employed for treating patients with severe tinnitus, selecting the most suitable therapies for each patient [3].

Treatment of Tinnitus with a Neurological View

While the pathophysiology of the different forms of tinnitus remains poorly understood, electrophysiologic and functional neuroimaging studies have recently shown the evidence of an association between severe chronic tinnitus and abnormal functioning of the central nervous system (CNS) [1, 2] (see Chap. 10). Abnormal neuronal firing within the auditory pathway may account for the perception of sound when there is no physical sound present (tinnitus). In detail, neuroimaging studies demonstrated that tinnitus is associated with increased activity of the inferior colliculus [4, 5], the thalamus [6, 7], the auditory cortex [8, 9], and the limbic structure (amygdala) [10, 11]. Evidence has been presented that the activation of neural plasticity is involved in chronic tinnitus as well as failing homeostatic mechanisms, thus resembling the pathologies of chronic pain syndromes [12–14]. Animal models of tinnitus suggest that \( \text{Ca}^{2+} \) signaling pathways as well as imbalance between the GABAergic and glutamatergic system are among known involved mechanisms [15].

Effective therapies of tinnitus have been based on known pathophysiology of tinnitus. Modulation of somatic inputs by transcranial magnetic stimulation (TMS) (see Chap. 88) and retraining therapy, alone or in combination, have been used (see Chap. 73).

TMS is a noninvasive method that can relieve tinnitus by modulating the excitability of neurons in the
auditory cortex to decrease the hyperexcitability believed to cause some forms of tinnitus (see Chap. 88) by direct electrical stimulation of the cerebral cortex (see Chap. 90). All these treatments are aimed at reversing hyperexcitability of the auditory pathways in the central and peripheral nervous system.

Medications used for reducing the increased neuronal excitability, anticonvulsants, have been repeatedly used for the treatment of tinnitus as well as antidepressants, benzodiazepines, acamprosate, and melatonin [15] (see Chaps. 78 and 79). Other methods that are in use for the treatment of tinnitus act to modulate or stimulate the somatic sensory system similar to what is used to treat central neuropathic pain (see Chaps. 91 and 94).

**References**

Chapter 27
The Psychiatrist

Berthold Langguth

Keypoints

1. Tinnitus is not a psychiatric disorder, but shares some relevant aspects with psychiatric disorders.
2. Tinnitus is frequently accompanied by psychiatric comorbidities.
3. Tinnitus research can benefit from recent advances in psychiatric research, e.g., in neuroimaging, genetics or clinical trials methodology.
4. Diagnosis of psychiatric comorbidity and psychopharmacologic treatment should be performed by psychiatrists, and psychotherapy by psychotherapists.
5. A multidisciplinary collaborative approach seems to be the most promising strategy both for the management of the tinnitus patients and for tinnitus research.

Keywords Tinnitus • Brain disorder • Psychiatry • Psychosomatics • Psychotherapy • Multidisciplinarity

Introduction

Traditionally, tinnitus is treated by otologists and audiologists. This can reasonably be explained by the fact that tinnitus is a sound and thus subjectively located to in the ears. It is common to seek help for hearing problems from otologists and hearing specialists (audiologists). Recent advances in neuroimaging and biomedical neuroscience, however, have initiated a paradigmatic shift by demonstrating that tinnitus is generated by an alteration of neural activity in the brain.

Tinnitus as a Brain Disorder

The central auditory system is involved in most forms of tinnitus, but also nonauditory brain areas, such as the frontal cortex or the limbic system. The changes in the central auditory system that cause tinnitus not only arise from auditory deprivation, but can also be caused or modulated by somatosensory input. Whether changes that occur in neural activity in the central auditory pathways are perceived as tinnitus depend on the degree of coherence of neural activity in populations of neurons and synchronous coactivation of a global neural network. The extent to which limbic brain structures are involved may determine the emotional burden (see Chaps. 9, 10, 13, 17, and 21 for more details).

Thus, many studies have demonstrated that the location of the pathology underlying tinnitus and tinnitus distress is in the brain. Hence, there is no doubt that the specific experience of neuroscientists, neurologists, and psychiatrists in the investigation of brain disorders can provide important contributions to tinnitus research.

Should Tinnitus be Considered to be a Psychiatric Disorder?

That the anatomical localization of tinnitus pathology is the brain does not automatically mean that tinnitus should be classified as a neurological or psychiatric
B. Langguth
disorder and treated by these disciplines. Other factors have to be considered, such as which discipline is best equipped to provide diagnostic and therapeutic management of patients with tinnitus. Also, historical aspects, the organization of the different national health care systems or the currently used diagnostic classification systems, may play a role in the choice of specialist best able to care for patients with tinnitus.

Taking into account all these aspects, one may find several arguments for classifying tinnitus as a psychiatric disorder in addition to the fact that the pathology of tinnitus is localized in the brain: (1) Tinnitus as a perceptual disorder shares phenomenological similarities with auditory hallucinations; (2) Emotional and cognitive impairment are core symptoms of tinnitus, and there are high rates of psychiatric comorbidity (see Chap. 62); (3) Psychoeducation (counseling) is probably the most widely used treatment for tinnitus, and psychotherapy is the only intervention for which efficacy has been shown in a Cochrane meta analysis [1]; (4) Also, most promising results of pharmacologic treatment have been shown for drugs used to treat psychiatric disorders, such as antidepressants or anticonvulsants [2] (see Chap. 78).

On the other hand, the importance of hearing disorders as the most important risk factor for tinnitus has been recognized, highlighting the relevance of detailed otologic and audiologic assessment. Treatments, which compensate for hearing loss, such as hearing aids or cochlear implants, have been clearly shown to improve tinnitus (see Chaps. 74 and 77) [3]. These arguments alone already justify that tinnitus patients are treated primarily by otologists and audiologists. There is probably general agreement that otologists and audiologists are most competent to provide information about hearing, but also to give recommendations on how to deal with hearing disorders and tinnitus. This is similar to other diseases, where one would expect, for example, the diabetologist to give recommendations for physical activity and eating behavior and not the behavioral therapist. Nevertheless, many ear specialists may benefit from some psychological training about how to best convey information or behavioral recommendations in their management of patients with tinnitus.

**The Role of the Psychiatrist**

Similar like audiological diagnosis and treatment require competent professionals, specific training is needed for psychiatric diagnosis and psychotherapeutic treatment. In this context, a clear distinction is also necessary between counseling, which comprises general information delivery and behavioral recommendation for tinnitus patients and psychotherapy. While counseling may be primarily the task of otologists or audiologists, cognitive behavioral therapy can best be performed by a specifically trained psychotherapist. Involvement of a specialist is also needed for the diagnosis and treatment of the frequently occurring psychiatric comorbidities.

But who is this specialist? Even among medical doctors there is confusion about the different “psycho-disciplines.” Which discipline can provide the best benefit to patients with tinnitus: psychology or psychiatry, psychotherapy or psychosomatics, psychoanalysis or behavioral therapy? There is no general answer to this question because competences of the different disciplines vary across countries and health systems but an orientation can be given. Diagnosis of psychiatric disorders can be made by psychiatrists and specifically trained psychologists. The different forms of psychotherapy all require specific training. Both psychologists and psychiatrists can be trained to become psychotherapists. Legally, pharmacologic treatment can only be prescribed by psychiatrists or other medical doctors.

**Tinnitus: A Psychosomatic Disorder?**

The term psychosomatics highlights the interaction between psychological and somatic aspects of health and diseases. Psychosomatics can be considered as a reaction to the body–soul dualism, which influenced our thinking since Descartes. However, recent advances in neuroscience clearly demonstrate that psychological factors, such as motivation, emotions, beliefs, or expectations, are related to neural activity in specific brain circuits and that successful psychotherapeutic treatment induces changes in these brain networks. Thus, neither the distinction between “somatic” and “psychological” etiology for a
symptom or a disease, nor the distinction between somatic and psychological treatment can be justified anymore. This also has important consequences for the still widespread assumption that symptoms without a detectable somatic correlate have a nonsomatic, ergo psychological, etiology. There is no reason to believe that there should be more unresolved psychological conflicts in individuals with tinnitus than in individuals who do not have tinnitus. There is also no evidence for any etiological relationship between psychological conflicts and the emergence of tinnitus. On the contrary, we know that tinnitus has always a neuronal correlate. If it is not possible to detect such changes in every patient, then this only reflects the present lack of methods and perhaps lack of knowledge. Any form of successful intervention, whether sensorial (hearing aids), physical (physiotherapy), pharmacologic, or psychological, exert their effect through the modulation of neural activity.

**Conclusion**

It is not helpful to distinguish between somatic and psychological causes for tinnitus. Instead, research should be directed toward identifying the neuronal correlates of the different forms of tinnitus and their comorbidities. It would be a mistake to ignore knowledge and methodological experience from biological psychiatric research. For example, neuroimaging is a valuable tool for targeting neuromodulation, and genetic research can contribute to the identification of molecular structures that can be manipulated by pharmacologic treatment.

Psychiatrists have an important role in clinical management of the patient with tinnitus. When there is suspicion of psychiatric comorbidity, a referral to a psychiatrist is critical and necessary when the patient has suicidal tendencies. (Criteria on how to identify these situations are given in Chap. 54 Tinnitus with psychiatric comorbidity). Prescription of psychopharmacologic drugs is best done by psychiatrists and whenever psychotherapy is indicated for the treatment of tinnitus, it should be performed by a trained psychotherapist.

It becomes clear that no single clinical specialty will be able to cover all relevant aspects of diagnosis and therapy for the different forms of tinnitus. Instead, close collaboration between different disciplines seems to be the most promising strategy, both for the clinical management of tinnitus patients and for tinnitus research. Our own experience from close interdisciplinary collaboration at tinnitus centers has taught us that such a concept that can be realized successfully to the benefit of patients with tinnitus.

**References**

Chapter 28
The Neurosurgeon

Dirk De Ridder

Keypoints

1. Neurosurgeons can contribute in a similar fashion to treatments of tinnitus as they currently do in pain treatment.
2. Neurosurgeons should collaborate with other clinicians and basic neuroscientists to help elucidate the pathophysiology of tinnitus.
3. Invasive neuromodulation can be helpful in selected forms of intractable tinnitus.
4. Different intracranial pathologies exist that can cause tinnitus amenable to surgical treatment, both of the non-pulsatile and of the pulsatile type.
5. Non-pulsatile tinnitus can be considered analogous to pain and results from changes in neural networks of the brain.
6. Pulsatile tinnitus is mostly related to anomalies of blood vessels in and around the brain.

Keywords  Tinnitus • Neurosurgeon • Tinnitus • Pulsatile • Non-pulsatile • Neurosurgery • Neuromodulation

Abbreviations

CPA  Cerebellopontine angle
CSF  Cerebrospinal fluid
ENT  Ear nose and throat
Gy  Gray (unit of absorbed radiation)

MRI  Magnetic resonance imaging
TMS  Transcranial magnetic stimulation

Introduction

Tinnitus has traditionally been a field belonging to ear nose and throat (ENT) surgeons, audiologists, and psychiatrists, except for some forms of pulsatile tinnitus, such as anomalies of the cerebral blood vessels, which have usually been treated by neurosurgeons.

Recently, both basic research [1] and clinical research [2, 3] have focused on the brain’s involvement in the generation of tinnitus, opening the tinnitus field up to neurologists and neurosurgeons specialized in the field of tinnitus (see also Chap. 26).

Neurosurgeons treat patients with pain in an invasive way, and based on the analogy between some forms of pain and tinnitus [4–7], both of which can be considered deafferentation or phantom phenomena [8], the step to treating tinnitus for neurosurgeons is not as big as it looks at first sight (see Chap. 94).

A patient’s referral to a neurosurgeon for pain relief was once considered bad news, because the choice of procedures was limited to the creation of lesions, offering significant risk and only modest success [9]. Neurosurgery used to be considered the “pursuit of the impossible by the irrepressible” [10]. In a similar way, the tinnitus field still considers the neurosurgeon a last resort, when everything else fails and the patient is suicidal or distressed by the tinnitus. Neurosurgical approaches to tinnitus are still too often described as “the half mad being operated upon by the mad” [10].

Advances in technology and an improved understanding of pain have helped to develop more effective procedures to such an extent that a recent textbook [11]
D. De Ridder discusses more than 30 types of procedures used in more than 18 major categories of pain [9]. However, as stated in the textbook, this does not mean that the neurosurgical procedures should always be the first line of treatment, as chances for pain relief are greatest when neurosurgery is but one piece of a comprehensive plan incorporating all possible treatment modalities [9].

Neurosurgeons treat the cause of pain (for example, disk surgery and microvascular decompressions) or use invasive neuromodulation when the cause of the pain is unknown or cannot be treated. In a similar way, the neurosurgeon should be involved in tinnitus treatment, dealing with the cause of the tinnitus, and by using neuromodulation to treat the symptoms.

There are indeed a series of pathologies that can cause tinnitus, either as their principal symptom or as one in a constellation of symptoms. Knowing the clinical course of the tinnitus in these pathologies is needed in order to be able to prognostically address these pathologies surgically. Some examples of specific diseases that often have tinnitus among their symptoms and that can be treated surgically are vestibular schwannoma, Arnold-Chiari malformations, arachnoid cysts, and others. Treatment of such diseases belongs to the classical repertoire of neurosurgery. However, neuromodulation through electrical (or magnetic) stimulation or lesioning, which are effective methods in treating disorders such as tinnitus and some forms of pain, also belong to the armamentarium of modern neurosurgery. Although neurosurgical procedures are traditionally the last resort in the battle against tinnitus, it is of interest for the tinnitus field to learn from neurosurgical pain management, which has brought relief to many patients with pain where other treatments have been ineffective. There are reasons to believe that neurosurgical treatment of tinnitus may evolve to become as widely used for treatment of tinnitus as it is now for treatment of pain.

However, brain surgeons should not limit themselves to developing new treatments for tinnitus based on analogy with pain. As a brain surgeon, one has a unique and unparalleled access to the brain, permitting recordings directly from the brain. It is important that they team up with basic neuroscientists to collaborate in order to gain as much valuable information as possible during the short window of direct brain access [12]. The power of intraoperative studies of brain function has a long history beginning with Penfield in the 1930s [13], extending to modern times where large parts of our understanding of the function of many systems of the human brain is based on intraoperative studies in patients undergoing neurosurgical operations [14, 15].

The neurosurgeon treating tinnitus should ideally work in a multidisciplinary team consisting of not only clinicians but also basic neuroscientists. Therefore, as long as no standardized neurosurgical treatments become available for tinnitus suppression, the neurosurgeon should not limit himself/herself to be a “sophisticated manual laborer” but should also be a “researcher” attempting to better understand the pathophysiology of tinnitus in order to develop new treatments for this elusive symptom (see Chaps. 21, 90, and 94).

**Neurosurgical Approaches to Tinnitus**

Tinnitus can be divided into two entirely different entities: pulsatile and non-pulsatile tinnitus [16–18]. Pulsatile tinnitus is usually related to vascular anomalies or intracranial hyper- or hypotension and is not related to an abnormal function of the auditory system. Non-pulsatile tinnitus, on the other hand, is critically related to an abnormal function of the auditory system.

**Non-Pulsatile Tinnitus**

Non-pulsatile tinnitus can be considered an auditory phantom phenomenon [8], resulting from auditory deprivation or deafferentation [1]. Any lesion along the auditory tract altering its normal function can cause non-pulsatile tinnitus. Ménière’s disease, vestibular schwannoma, cerebellopontine angle (CPA) lesions, arachnoid cysts, microvascular compressions, Chiari malformation, and brain tumors are causes of non-pulsatile tinnitus that can be treated surgically.

If no cause for a patient’s tinnitus can be found and thus no causal treatment can be offered, attempts to provide permanent relief from treatments such as electrical stimulation should be tried. First, non-invasive stimulations at different targets of the auditory system (promontory stimulation, transcranial magnetic stimulation...
[TMS], transcranial direct current stimulation, transcranial electrical nerve stimulation) to test if a permanent implant could be beneficial should be performed. However, no prognostic relation has yet been shown between the effect of TMS aimed at the auditory cortex or cortical electrical stimulation at the level of the auditory cortex.

In Vestibular schwannoma, a high-pitch tinnitus (described as ringing or steam from a kettle) is present in 60–85% of the participants in a recent study [19]. Since the advent of stereotactic irradiation, vestibular schwannomas are often treated by radiosurgery, especially gamma knife radiosurgery. This seems to have a similar effect on tinnitus as microsurgery, although it seems to induce less tinnitus in the short term after treatment. Studies have shown that the tinnitus in 12–46% of such patients improves after the treatment [20] and tinnitus develops in only 4% of the patients after radiosurgery [20, 21]. The tinnitus experienced by patients who underwent microscopic surgery for removal of vestibular schwannoma disappeared in 16–50% of the participants in a study [22, 23]. Other studies have shown that after surgery to remove vestibular schwannoma, the tinnitus is reduced in 16% of patients, in 55% it does not change, and in 29% it becomes worse [22, 24, 25], especially when hearing is saved in surgery [24]. While the prevalence of tinnitus before and after operations, where hearing preservation is not attempted, is not significantly different, there are significant differences in tinnitus before and after operations where hearing is saved. When tinnitus is absent preoperatively, 85% of the hearing preservation group develops tinnitus after the operation while only 31% of patients in whom hearing preservation was not attempted developed tinnitus [24]. The results of other studies are, however, more optimistic, showing that only 8% of patients developed tinnitus after hearing preservation operations for vestibular schwannoma [26].

Gamma knife treatment has advantages over surgery as well as disadvantages. Gamma knife radiosurgery is less invasive and requires shorter hospitalization and convalescence periods [27]. The development of facial palsy or paresis is extremely rare (1% if irradiation dose is <14 Gy), and hearing can be saved in almost 80% of patients if 13 Gy as the maximum dose is respected [28]. The technique is, however, limited to lesions less than 3 cm in size and carries a greater risk for the development of post-treatment hydrocephalus and a certain, though small, risk of dedifferentiation into a neoplasia (malignancy). In small lesions (<1.5 cm) without serviceable hearing microsurgery and gamma knife treatment have comparable rates of tinnitus, tumor control, facial nerve function, and trigeminal function. However, stereotactic radiosurgery has a greater risk of long-term balance problems compared to microsurgery [29]. In general, gamma knife surgery might be better for vestibular schwannoma treatment in the short term [30], with similar effects on improvement and worsening of tinnitus as surgery. Surgery after failed gamma knife treatment has an increased risk for facial palsy due to strong adhesions [31] (see Chap. 85).

Other CPA lesions [32] such as meningo, epidermoid tumors, lipoma, choroid plexus papilloma, epithelial cysts, teratoma, cavernoma, and hemangioma are sometimes associated with non-pulsatile tinnitus, usually together with other symptoms depending on the location of the lesion and the degree of brainstem, cerebellar, or cranial nerve compression.

Arachnoid cysts are a rare cause of non-pulsatile tinnitus. It is a congenital or posttraumatic/post-inflammatory disorder [33, 34], leading to vague symptoms [35]. However, infratentorial [36] arachnoid cysts can sometimes mimic Ménière’s disease as well. Arachnoid cysts producing tinnitus can occur in the CPA [35, 37, 38], but also retroclival, retrocerebellar, and lateral of the cerebellum [39], with postoperative improvement of the tinnitus [39]. Usually, symptoms of intracranial hypertension are associated with non-pulsatile tinnitus [35, 40]. Surgical treatment consists of marsupialization1 or excision of the cyst [40]. Also supratentorial Sylvian fissure arachnoid cysts can generate isolated tinnitus, and tinnitus suppression can be the result of marsupialization of the cysts if they act as a mass lesion [41]. Supratentorial cysts can also mimic Mérié’s disease [42]. Imaging studies using intrathecal contrast to verify if an arachnoid cyst-like lesion communicates with normal cerebrospinal fluid (CSF) flow can help to ascertain whether an arachnoid cyst could act as a mass lesion and thus be symptomatic or not. Magnetic resonance imaging (MRI) sequences looking for a flow void within the cyst can be helpful as well [43].

1 Marsupialization: Surgical alteration of a cyst or similar enclosed cavity by making an incision and suturing the flaps to the adjacent tissue, creating a pouch. (From: The American Heritage® Stedman’s Medical Dictionary.)
Ménière’s syndrome is a clinical entity consisting of episodic vertigo, fluctuating sensory hearing loss, tinnitus, and aural fullness (see Chap. 38). This syndrome is caused histopathologically by endolymphatic hydrops that can be caused by many pathologies – traumatic (acoustic, iatrogenic, or temporal bone trauma and labyrinthine concussion), infectious/inflammatory (autoimmune inner ear, see Chap. 60), Cogan’s syndrome, chronic otitis media, viral or serous labyrinthitis, syphilis, tumoral (leukemia), congenital (deafness, Mondini dysplasia), or in the setting of connective tissue or bone disease (Letterer-Siwe disease, Paget’s disease, otosclerosis), and others [44]. Tinnitus worsens both in intensity and as a function of duration and bilateral disease [45]. It is perceived as worse than in a comparable group of tinnitus sufferers due to acoustic trauma or otosclerosis [45].

In Ménière’s disease, any kind of surgery, whether vestibular nerve section, cochlear nerve section, endolymphatic sac surgery [46], or gentamicin injections [47], never seems to produce greater than 50% tinnitus control – a small improvement upon the 30% spontaneous disappearance in its natural history [48]. Endolymphatic sac surgery, independent on whether decompression, exclusion, or shunting is done, improves or cures tinnitus in 40% of patients with Ménière’s disease [49]. This is similar to intratympanic gentamycin application, a less invasive technique with 27–69% tinnitus improvement [50–52].

In a recent review paper on vestibular nerve section performed for tinnitus [53], the proportion of patients in whom tinnitus was exacerbated postoperatively ranged from 0 to 60%, with a mean of 16.4%. The proportion of patients in whom tinnitus was unchanged was 17–72% (mean 38.5%), and in whom tinnitus was improved was 6–61% (mean 37.2%). These results are similar to gentamycin and endolymphatic sac surgery. In the majority of patients undergoing vestibular nerve section, ablation of auditory efferent input (and thus total efferent dysfunction) to the cochlea was not associated with an exacerbation of tinnitus [53].

In otosclerosis, tinnitus is very common; up to 91% of individuals with otosclerosis have tinnitus and 38% is severely affected by it [54]. Successful stapedectomy causes disappearance of non-pulsatile tinnitus in up to 40–73% tinnitus [55–60] with another 32–37% improving. In individuals who did not have tinnitus before stapedectomy, the risk of developing it after the surgery is almost non-existent. Only in 10%, the operation does not improve the tinnitus [60] and in another 8% it worsens [57]. Rarely, otosclerosis also produces arterial pulsatile tinnitus, due to a neovascularization at the site of stapes fusion. Stapedectomy can sometimes cure this rare form of pulsatile tinnitus [17].

A tumor in the auditory cortex, compressing the auditory cortex, can cause ipsilateral fluctuating non-pulsatile tinnitus as the sole symptom, probably due to a direct influence on normal cortical sound processing. Removal of the lesion resulted in abolishing the tinnitus in 4 out of 5 patients who had the operation [41]. Tumors elsewhere along the auditory tract (for example, the brainstem) rarely present with tinnitus only but usually give rise to additional symptoms related to the tumor’s closeness of other neural structures in the brainstem.

For intractable non-pulsatile tinnitus, auditory brainstem implants [61] (see Chap. 77) and auditory cortex stimulations can give relief in intractable non-pulsatile tinnitus [62–64]. These treatments are based on a recently developed pathophysiological model for non-pulsatile tinnitus, based on auditory deprivation or deafferentation as the initial trigger for tinnitus generation. Studies have shown that a decrease of auditory input induces a slowing of auditory information processed in the thalamocortical loop generating slow wave activity (delta en theta oscillations) [7, 65], with a decrease in lateral inhibition [66] and a halo or edge of increased activity [7, 67]. This is also called thalamocortical dysrhythmia [7] associated with cortical reorganization [68, 69]. The most likely mechanism that links hyperactivity and reorganization is synchrony [1]. Synchronization of the gamma band activity could possibly induce topographical reorganization via simple Hebbian mechanisms (cells that fire together wire together) [1]. Therefore, it seems logical to try and modify this tinnitus-related auditory cortex reorganization/hyperactivity in an attempt to suppress the tinnitus. This can be achieved using neuronavigation-guided TMS, a technique that is capable of modulating cortical activity. If TMS is capable of suppressing tinnitus, the effect could be maintained by implantation of electrodes at the area of signal abnormality on the auditory cortex. The first results in
patients with unilateral pure-tone tinnitus have shown statistically significant tinnitus suppression, without suppressing white or narrow band noise in individuals who responded to TMS with decreased tinnitus [63]. More recent trials also suggested that narrow band tinnitus is suppressible with novel stimulation designs consisting of closely spaced spikes of very high frequencies [70].

Reafferentation of the auditory thalamocortical system after it has been deprived of input can also be achieved by cochlear implants (see Chap. 77). Almost, immediately after the introduction of cochlear implants for hearing improvement, it was noted that the electrical intracochlear stimulation ameliorated tinnitus in a large proportion of individuals [71, 72]. Multiple studies since then have replicated these results indicating that cochlear implants inserted for hearing improvement can also modulate tinnitus [73–77], not only unilaterally but also bilaterally in a majority of individuals [78]. A recent study using cochlear implant insertion in patients with incapacitating tinnitus and ipsilateral complete hearing loss and contralateral preserved hearing demonstrates similarly promising results [79]. Using promontory stimulation as a preoperative non-invasive test in this selected group of patients might predict good outcomes in tinnitus suppression.

A limit to this technique is that it can only be used in patients with unilateral complete hearing loss. This could potentially be extended to people with high-frequency hearing loss but preserved low-frequency hearing, as a recent paper has shown that short hybrid cochlear implants can preserve low-frequency hearing [80]. Another option is to use extracochlear stimulation for tinnitus suppression. The first attempts for developing extracochlear electrical stimulation have been made [76, 81] as well.

### Conclusion

Stimulated by recent developments in our understanding of the pathophysiology of tinnitus, treatment has shifted from purely otological approaches to brain-based approaches. Therefore, neurosurgeons should become more involved in treating this elusive symptom.

Tinnitus actually consists of two entirely different entities with a different pathophysiology, different clinical symptoms, and different treatment. Before tinnitus patients are told “to learn to live with their tinnitus” it can be suggested to look for possible causes for both non-pulsatile and pulsatile tinnitus as this can result in an otoneurosurgical treatment. In patients with non-pulsatile tinnitus, non-invasive trials with promontory or TMS can potentially help select candidates for a permanent implant as a treatment for tinnitus. Neurosurgeons should be involved not only in the surgical treatment of operable causes but also in the exploration of possible pathophysiological mechanisms, making use of their unique ability to

<table>
<thead>
<tr>
<th>Table 28.1</th>
<th>Surgically treatable causes of tinnitus</th>
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<tbody>
<tr>
<td><strong>Pulsatile tinnitus</strong></td>
<td><strong>Non-pulsatile tinnitus</strong></td>
</tr>
</tbody>
</table>
| Venous | Vestibular schwannoma  
(acoustic neuroma) |
| Benign intracranial hypertension | Other cerebellopontine angle lesions |
| Chiari malformation | Arachnoid cyst |
| High jugular bulb | Menière’s disease |
| Sigmoid sinus diverticulum | Otosclerosis |
| Sigmoid-transverse aneurysm | Microvascular compression |
| Aberrant veins aneurysm | Chiari malformation |
| Arterial | Brain tumor |
| Carotid stenosis | Symptomatic |
| Aberrant carotid artery | Cochlear implant |
| Glomus tumor | Brainstem implant |
| Vascular lesions of petrous bone/skull base | Auditory cortex implant |
| Arteriovenous malformation | Pseudopulsatile tinnitus |
| Aneurysm | Palatal myoclonus |
| Canal dehiscence | Middle ear myoclonus |
| Benign intracranial hypertension | Patulous eustachian tube |
| Carotid-cavernous fistula | |
record activity directly from the brain when performing intracranial surgery.

References

functional magnetic resonance imaging paradigm suitable for clinical use. Invest Radiol 41:87–96.
Keypoints

1. Patients with tinnitus are *prima facie* beyond the responsibility of dentists.
2. Studies of the prevalence of tinnitus in people with temporomandibular disorders (TMD) (give values from 2 to 59%) and the prevalence of TMD in patients with tinnitus ranging from 7 to 95%. Evidence about the relationship between TMD and tinnitus is conflicting and it is not known if it is causal or coincidental.
3. Patients with TMD-related tinnitus can benefit from TMD therapy but TMD therapy in patients with tinnitus without any signs of TMD is not recommended.

Keywords  TMD • Tinnitus • Dentist

Abbreviations

CMD Craniomandibular disorder(s)
MPD Myofascial pain dysfunction
TMD Temporomandibular disorder(s)
TMJ Temporomandibular joint

Introduction

Tinnitus is generally regarded as a symptom of the ear or an auditory disorder. Therefore, patients with tinnitus are *prima facie* beyond the responsibility of dentists or maxillofacial surgeons. Additionally, patients suffering from tinnitus do not primarily consult a dentist, and most patients will not relate their “ear symptoms” to possible stomatognathic or temporomandibular disorders. The understanding of tinnitus symptoms and knowledge on the pathophysiology of different forms of tinnitus has, however, changed in recent years. Tinnitus researchers have benefited from learning from other fields of medicine, from cooperating with other disciplines, and from “thinking outside the box” [1]. Today, tinnitus is seen as a symptom presenting in many forms, and the contribution of dental science to a better understanding of tinnitus is appreciated by “traditional tinnitus therapists”, such as otolaryngologists, audiologists, psychologists, and psychiatrists.

Temporomandibular Disorders

Dentists and maxillofacial surgeons have long known that tinnitus symptoms are not uncommon in patients with temporomandibular joint (TMJ) and masticatory muscle disorders – also referred to as Costen’s syndrome – [2], craniomandibular disorders (CMD), myofascial pain dysfunction (MPD), temporomandibular dysfunction, or temporomandibular joint syndrome [3]. Nowadays, these terms are summarized under the heading “temporomandibular disorders” (TMD) [4, 5]. TMD are considered as a cluster of various joint and muscle disorders and a subgroup of general musculoskeletal and rheumatologic disorders, but should be regarded as a distinct group of diseases and symptoms [6]. The complex signs and symptoms of TMD are generally described as pain or tenderness in the
region of the TMJ or the masticatory muscles (myofascial pain), limitation or disturbance of mandibular movements, joint sound (clicking and crepitation), locking, oral parafunction, masticatory muscle hyperactivities (bruxism, clenching, and rocking of teeth), and fatigue in the jaws [3, 6]. Unfortunately, since the classification of the different forms of TMD is still not agreed upon, numerous ways of categorizing TMD have been proposed [7]. In general, TMD can be classified as a joint disorder (including structural deviations, mechanical derangements, and inflammatory disorder or arthritis), muscle disorder, and a combination of both [4, 6]. Clinicians who treated patients with TMD as a main complaint have noted that these patients often present with ear symptoms as a secondary complaint. Therefore, related conditions such as tinnitus were improved and often eliminated after treatment of their TMJ problems [8–11]. Tinnitus and TMD symptoms show many parallels in their clinical appearance. Knowledge of the etiology of both symptoms and disorders is limited. Thus, valid and reproducible diagnostic criteria are lacking. As a result, conflicting opinions exist on therapeutic proceedings for patients with tinnitus and TMD. Success rates of specific therapies remain unpredictable, which in turn transforms patients of both groups into an “unpopular” group of patients.

Prevalence of Temporomandibular Joint Disorders

The literature contains conflicting evidence about the prevalence of tinnitus in individuals with TMD as a main complaint (ranging from 2 to 59%), but most studies report a much higher prevalence of tinnitus in patients with TMD than in the general population. Unfortunately, most of the presented studies are mainly descriptive and have not been designed to compare between patients with symptoms and a reference group (Table 29.1). Studies of the general population showed prevalence of tinnitus from 14.2 to 20.1% (please see Chap. 5). Vice versa, information on the prevalence of TMD in patients with tinnitus is also incongruent (ranging from 7 to 95%) (Table 29.2). However, incidence of TMD was found to be higher in patients with tinnitus than in the general population, where tinnitus occurred in 16–59% for reported symptoms and in 33–86% for clinical signs [12].

Relation Between TMD and Ear Problems

Many different manifestations lead to the diagnosis of TMD, and a discrepancy exists between reported

<table>
<thead>
<tr>
<th>Table 29.1</th>
<th>Studies reporting tinnitus in patients with TMD as the main complaint</th>
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</thead>
<tbody>
<tr>
<td><strong>Source</strong></td>
<td><strong>Prevalence of tinnitus, no (%)</strong></td>
</tr>
<tr>
<td></td>
<td><strong>Patients with TMD</strong></td>
</tr>
<tr>
<td>Bernstein et al. [35]</td>
<td>36/86 (42%)</td>
</tr>
<tr>
<td>Bush [8]</td>
<td>35/105 (33%)</td>
</tr>
<tr>
<td>Bürgers (unpublished)</td>
<td>30/82 (37%)</td>
</tr>
<tr>
<td>Camparis et al. [36]</td>
<td>54/100 (54%)</td>
</tr>
<tr>
<td>Cooper et al. [37]</td>
<td>301/837 (36%)</td>
</tr>
<tr>
<td>Dolowitz et al. [38]</td>
<td>200/338 (59%)</td>
</tr>
<tr>
<td>Gelb et al. [39]</td>
<td>311/742 (42%)</td>
</tr>
<tr>
<td>Gelb et al. [40]</td>
<td>71/200 (36%)</td>
</tr>
<tr>
<td>Goodfriend [31]</td>
<td>24/168 (14%)</td>
</tr>
<tr>
<td>Hankey [41]</td>
<td>6/68 (9%)</td>
</tr>
<tr>
<td>Koskinen et al. [42]</td>
<td>9/47 (19%)</td>
</tr>
<tr>
<td>Myrhaug [32]</td>
<td>436/1,391 (31%)</td>
</tr>
<tr>
<td>Parker et al. [9] and Chole et al. [28]</td>
<td>199/338 (59%)</td>
</tr>
<tr>
<td>Rubinstein et al. [43]</td>
<td>93/376 (25%)</td>
</tr>
<tr>
<td>Tuz et al. [10]</td>
<td>91/200 (46%)</td>
</tr>
<tr>
<td>Upton et al. [44]</td>
<td>72/989 (7%)</td>
</tr>
<tr>
<td>Wedel et al. [45]</td>
<td>8/350 (2%)</td>
</tr>
<tr>
<td>Wright et al. [46]</td>
<td>101/267 (38%)</td>
</tr>
</tbody>
</table>
symptoms and clinical findings. Therefore, epidemiological studies on TMD (as well as on tinnitus) should not be compared without restrictions. Nevertheless, the simultaneous occurrence of tinnitus and TMD has led to the assumption that there may be a relationship between the two conditions. The initial claim relating tinnitus symptoms, temporomandibular joint, and masticatory muscle disorders was made by Costen in 1934, who described a syndrome of ear and sinus symptoms relating to disturbed TMJ function [2]. Although Costen’s structural and mechanical theories on the correlation of TMD and tinnitus have now been discarded, his considerations started numerous scientific efforts to reveal the linkage between both symptoms [13–16]. At this point, many questions on this topic remain unexplained. We still do not know whether ear symptoms (such as tinnitus and TMD) are coexistent, independent, or unrelated [8, 17–22] or whether both diseases have a causal connection [23–30]. Since TMD and tinnitus occur frequently in humans, their coincidence may not mean these two diseases have common causes or common risk factors. Authors reporting causal associations between tinnitus and TMD have based their conclusions mainly on clinical, epidemiological, anatomical, and histological investigations [23–27].

For example, the simultaneous occurrence of bruxism (grinding of teeth) as a symptom of TMD and tinnitus may be explained by two different ways: patients with bruxism (TMD as a main complaint, shifting therapeutic responsibilities toward otorhinolaryngologists, audiologists, psychiatrists, etc.) process ear symptoms through grinding their teeth nightly. In addition to these causal explanations, these symptoms may occur without any causal relationship, or the presence of a third “disease” such as mental pressure, physiostress, or specific medication can act as a shared reason or a collective trigger causing TMD and tinnitus as secondary complaints [9, 19, 20]. Parker and Chole assumed that the relationship between TMD and tinnitus may be that both are responses to emotional stress [9]. However, attempts to find such a specific collective trigger for tinnitus and TMD symptoms remain speculative [21]. TMD-related tinnitus has been classified as objective tinnitus in most studies [26, 31, 32]. In contrast, Shulman and co-workers considered TMD-related tinnitus as subjective idiopathic tinnitus that was thought to directly or indirectly extend from a temporomandibular joint dysfunction on the auditory system [33, 34].

Besides epidemiological studies on TMD-related tinnitus and the steric adjacency of the Porus acusticus and the TMJ, a causal relationship between both symptoms has been observed. Ren and Isberg, for example, stated that in 53 patients with unilateral tinnitus and anterior disk displacement, disk displacement was found to be present in the ipsilateral joints in all patients, whereas the contralateral joint was asymptomatic in 50 patients (94%) [27]. In some patients, the intensity and quality of tinnitus can be altered (in most cases an enhancement) by mandibular movements, by pressure applied to the TMJ, or by biting [3, 13, 18, 24]. These alterations may indicate that increased activity of the masticatory muscles

### Table 29.2  Studies reporting TMD in patients with tinnitus as the main complaint

<table>
<thead>
<tr>
<th>Source</th>
<th>Prevalence of TMD, no (%)</th>
<th>General population</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bernhardt et al. [47]</td>
<td>18/30 (60%) &gt;2 TMD symptoms</td>
<td>697/1,907 (37%)</td>
</tr>
<tr>
<td>Bosel et al. [17]</td>
<td>129/340 (38%)</td>
<td>–</td>
</tr>
<tr>
<td>Kempf et al. [48]</td>
<td>110/138 (80%)</td>
<td>–</td>
</tr>
<tr>
<td>Linsen et al. [49]</td>
<td>17/22 (77%)</td>
<td>–</td>
</tr>
<tr>
<td>Morgan [23]</td>
<td>19/20 (95%)</td>
<td>–</td>
</tr>
<tr>
<td>Peroz [13]</td>
<td>TMJ sounds 9/40 (23%)</td>
<td>1/35 (3%)</td>
</tr>
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<td></td>
<td>Muscle tenderness 27/40 (93%)</td>
<td>8/35 (23%)</td>
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<td></td>
<td>Bruxism 25/40 (63%)</td>
<td>13/35 (37%)</td>
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<tr>
<td>Rubinstein et al. [43]</td>
<td>47/102 (46%)</td>
<td>–</td>
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<tr>
<td>Tullberg et al. [50]</td>
<td>101/120 (84%)</td>
<td>–</td>
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<tr>
<td>Upton et al. [44]</td>
<td>72/989 (7%)</td>
<td>–</td>
</tr>
<tr>
<td>Vernon et al. [18]</td>
<td>69/1,002 (7%)</td>
<td>–</td>
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</table>

*Patients with inner ear dysfunction
or pressure on the TMJ increases or even causes the perception of tinnitus, which in turn corroborates the theory that TMD is the causal trigger of tinnitus [3]. Nevertheless, up to now, no conclusive explanation exists for this phenomenon. It should be mentioned that some authors could not find any epidemiological correlation between TMD and tinnitus symptoms [8, 18, 22]. It should also be mentioned that the innervations of the TMJ and adjacent tissue project to cells in the upper part of the spinal cord and the trigeminal nucleus, which in turn project to cochlear nucleus (see Chaps. 8 and 9). This may explain why some individuals with TMD also have tinnitus.

From a dental perspective, tinnitus is possibly a secondary complaint of TMD or vice versa. Therefore, evaluation of possible involvement of the TMJ and masticatory muscle disorders seems feasible in all patients with tinnitus, as well as using TMD therapy in patients with TMD symptoms (TMD-related tinnitus). In contrast, TMD therapy in patients with tinnitus but without any signs of TMD is not based on scientific evidence.

References

13. Peroz, I. Dysfunctions of the stomatognathic system in tinnitus patients compared to controls. HNO, 2003;51:544–549
20. Laskin, DM, Block, S. Diagnosis and treatment of myofacial pain-dysfunction (MPD) syndrome. J Prosthet Dent, 1986;56:75–84
Chapter 30
The Pharmacologist

Ana Belén Elgoyhen and Carla Vanina Rothlin

Keypoints

1. One in 10 adults has subjective tinnitus, and for 1 in 100 adults, tinnitus severely affects their quality of life.
2. Despite the significant unmet clinical need for a safe and effective drug targeting tinnitus relief, there is currently not a single FDA-approved drug on the market.
3. Since in some individuals, tinnitus causes irritability, agitation, stress, depression, insomnia, and interferes with normal life, even a drug that produces a small but significant effect would have a huge therapeutic impact.
4. A glimpse of hope is appearing in the near future, as some pharmaceutical industries now have compounds targeting tinnitus in their pipeline.
5. If these compounds finally reach the market, they will set a new era that will revolutionize the treatment of tinnitus.

Keywords  Tinnitus • Phantom sound • Animal models • Lead compounds • Drug discovery

Tinnitus: A Clinical Unmet Need

Despite the significant unmet clinical need for a safe and effective drug targeting tinnitus relief, there is currently not a single FDA-approved drug on the market. For the majority of tinnitus sufferers who seek medical advice, the treatment goals are aimed at symptomatic relief (i.e. reduce or eliminate the tinnitus that is referred to as inside the head and/or ears). Symptomatic treatment is usually justified, because serious underlying pathologies are rare (see Sect. 30.2). Over four million prescriptions are written each year for tinnitus relief in Europe and the US, but these are all off-label prescriptions from a wide variety of therapeutic drugs, many of which are associated with considerable side effects or are ineffective in relieving tinnitus. There is, therefore, a large need for an effective drug therapy targeted at tinnitus, with minimal side effects compared to current medications prescribed off-label. Since in some individuals, tinnitus causes irritability, agitation, stress, depression, insomnia, and interferes with normal life, even a drug that produces a small but significant effect would have a huge therapeutic impact. However, disappearance of tinnitus should be the ultimate goal.

Tinnitus can be Pharmacologically Targeted

While the initial lesion might affect the peripheral organ of the auditory system, the neural correlate of the perceived sound is most likely in the central auditory circuitry [1] and there is growing evidence that changes in neuronal activity in different parts of the auditory pathway, including the dorsal cochlear nucleus, inferior colliculus, thalamus, and/or auditory cortex may be involved in tinnitus pathology [2–9]. Neuronal excitability can be modulated by different neurotransmitters, neuromodulators, and voltage-gated channel acting compounds [10–14], so there is no reason to believe that activity-driven changes underlying...
tinnitus cannot be pharmacologically targeted. The fact that a local anesthetic, the voltage-gated sodium channel blocker lidocaine [15], given intravenously, leads to the temporary disappearance of tinnitus or a major change in the nature of the tinnitus in 70% of patients [16–22], indicates that Pharmacologic agents can have beneficial effects on many forms of tinnitus.

**Challenges Toward Developing a Tinnitus Drug**

The quest for effective tinnitus therapies faces significant challenges. First, tinnitus is only a symptom that might be the manifest of different underlying pathologies. Differential diagnosis of triggering events and temporal onset should allow for a more rational and effective pharmacological approach. Therefore, the careful classification of tinnitus patients together with the search for drugs that can successfully target each underlying pathology becomes a priority. Moreover, the current limited understanding of the neural substrates of tinnitus, together with the lack of adequate animal models that can faithfully recapitulate its pathology, hampers the screen for new molecules in preclinical studies. Finally, because the first tinnitus drugs are yet to be approved, regulatory agencies such as the Food and Drug Administration (FDA) or the European Medicines Agency (EMEA) lack standardized protocols for their approval process. The often considerable placebo effect is another obstacle in selecting new substances for tinnitus treatment.

Many pharmacological agents have been used off-label to treat individuals with tinnitus. These include anticonvulsants, anxiolytic, antidepressants, NMDA antagonists, cholinergic antagonists, antihistamines, vasodilators, and antipsychotics, to name a few (see Chap. 78) [23, 24]. Some drugs have been reported to provide moderate relief of symptoms in a subset of patients. Careful clinical observations along with data from clinical trials have provided useful clues for deciding on a rational course of drug therapy for selected patients. However, most drugs have not proven sufficient effectiveness in randomized controlled clinical trials in order to be marketed specifically for tinnitus, highlighting the importance of selectively targeting the underlying pathological cause of tinnitus.

The first step toward designing a successful strategy in the search for tinnitus drugs would most likely include finding criteria by which to stratify tinnitus patients included in trials. As previously discussed, tinnitus often occurs as a result of insults to the ear, such as from noise exposure or administration of specific pharmacologic agents. It can also be caused by ear or head injuries, some diseases of the ear, and ear infections [1, 25]. In some cases, the causative agent remains unknown. Therefore, the identification of the triggering cause should aid in selecting the most adequate pharmacological approaches. In addition, tinnitus sounds can take a variety of forms, such as buzzing, ringing, whistling, hissing, or a range of other sounds. It can be a benign sound that is heard only occasionally or it can be devastating roars that occur 24 h a day, which prevent its sufferers from sleep or the ability to do intellectual work. All degrees of subjective tinnitus occur in between these extremes. Tinnitus is also often associated with other symptoms, such as hyperacusis and distortion of sounds [25]. Affective disorders, such as anxiety, phonophobia, and depression, often accompany severe tinnitus, and that form of tinnitus can lead to suicide. With such differences in etiology and symptoms, heterogeneity within tinnitus patients is expected. Thus, the tinnitus drug discovery endeavor faces the “one drug won’t fit all” situation. The fact that a subgroup of patients who have intermittent tinnitus that sounds like a typewriter, popcorn, or ear clicking receives significant benefit from carbamazepine [26, 27] indicates that “subtyping” tinnitus is highly needed for successful treatment. Efforts toward establishing subgroups of tinnitus are under way [28] and will most likely aid the selection of patients in future clinical trials.

An additional challenge in the design of drugs for the treatment of tinnitus derives from the fact that the neural substrates underlying tinnitus are far from being fully understood. An increase in spontaneous firing rates or neuronal synchrony in different parts of the auditory pathway as well as changes in cortical tonotopy have been proposed as potential correlates of tinnitus [1, 29]. Modern drug discovery is mostly centered on the identification of new lead molecules that interact with discrete molecular targets. This is a reductionistic approach that mainly focuses on sites of drug action. Although it has been useful in developing molecules such as statins (inhibitors of HMG CoA reductase) and HIV protease inhibitors [30], central nervous system
acting drugs owe their clinical effectiveness to actions at multiple molecular targets [31]. Thus, this reductionistic approach is most likely inadequate for a central nervous system disorder such as tinnitus.

Although a well-defined neuronal target would ease the path toward drug discovery, the empirical approach that has been used for most central nervous system disorders should not be precluded in the case of tinnitus. The importance of this approach in central nervous system drug discovery can be appreciated in the case of morphine and barbiturates, whose mechanisms of action were unknown when these drugs were introduced for human use [30]. In fact, most central nervous system acting drugs were discovered serendipitously. Thus, for example, valproic acid was used as an organic solvent in research laboratories for eight decades, until the observation of action against pentylenetetrazol-induced convulsions in rodents was made [32]; chlorpromazine was used to enhance recovery from surgical anesthesia before it was found to alleviate some symptoms of schizophrenia [33]; gabapentin was first developed as an anticonvulsant and is now used for treating neuropathic pain [34]. Thus, following these past experiences with central nervous system acting drugs, the search for drugs to alleviate tinnitus should not wait until the neural correlates are identified.

Before a compound is judged suitable for testing in humans, it must first demonstrate safety and efficacy in animal models. A drawback in the development of a tinnitus drug is the lack of validated animal models in which to test or screen for compounds. The basic dilemma faced by the animal researcher who wants to study tinnitus is whether or not the animals have the disorder. The experimenter has to find a way by which a rodent tells him about the ringing in its head. Several animal models are being developed, which are based either on noise exposure or on the administration of salicylate (see Chap. 16 and [35–37]). An additional challenge is imposed by the fact that, in humans, tinnitus is accompanied by the activation of a distress network that involves the limbic system [38–40]. This is probably not recapitulated in the animal models. However, animal models that have been developed for complex central nervous diseases such as depression or schizophrenia do not completely recapitulate the disease itself. Moreover, they are only of limited value for predicting treatment efficacy in humans [41]. However, in spite of all these drawbacks, these animal models have proven useful. In addition, in psychiatric diseases, empiric pharmacology has driven science. Thus, the serendipitous observation that central nervous system acting drugs like chlorpromazine calmed inmates of a psychiatric asylum has given way to the dopamine theory of schizophrenia and to the serotonin theory of depression and anxiety [41, 42]. These theories remain the pillars of the animal models used for preclinical validation, in spite of the fact that there is more to the major psychoses than alterations in these two neurotransmitter systems. Thus, the search for drugs to treat tinnitus should not wait for the refinement of the animal models. Moreover, the identification of compounds that alleviate tinnitus would not only lead to a better treatment but would also serve as a possible starting point for the understanding of the neural correlates of this condition, and thereof for the generation of better animal models, which target these neural substrates.

Finally, since no drug having tinnitus as its primary indication has been approved so far, there are no standardized protocols for the approval of a tinnitus drug by regulatory agencies like the FDA and EMEA. Therefore, the first pharmaceutical industry to develop a tinnitus drug will have to pave the way. In addition, tinnitus being a subjective phenomenon, assessment of outcome is probably the single most important factor in conducting a clinical trial. Widespread recognition that consistency between research centers in the ways that patients with tinnitus are assessed and how outcomes following interventions are measured would facilitate more effective co-operation and more meaningful evaluations. At the first Tinnitus Research Initiative meeting held in Regensburg in July 2006, which gathered worldwide tinnitus experts, an attempt was made to establish a consensus both for patient assessments and for outcome measurements [43].

**Tinnitus and the Pharmaceutical Industry**

Pharmaceutical companies are aware of the fact that there is a large market for a drug indicated for tinnitus relief. Evidence for this exists in the scores of patents that have been filed worldwide on potential drugs that may offer relief. Furthermore, tinnitus can be found attached to long lists of indications in many more patents filed on molecules aimed at a range of diverse
therapeutic classes. As indicated above, in spite of the fact that there is a significant unmet clinical need for a safe and effective drug targeting tinnitus relief, there is no FDA-approved drug currently on the market. The Royal National Institute for Deaf People, in the UK, estimates that a novel tinnitus drug could have a product value of US $689 million in its first year of launch [44]. However, there are very few pharmaceutical and/or biotechnology companies with tinnitus compounds in their R&D pipeline. A search carried out in the investigational drug databases Pharmaprojects (http://www.pharmaprojects.com), AdisInsight (http://www.adisinsight.com), Prous DDR (http://www.prous.com), and IDdb3 (http://science.thomsonreuters.com) shows that the following companies are developing a compound for tinnitus: Epicept, a lidocaine patch at phase II; Sound Pharmaceuticals, ebselen, a glutathione peroxidase mimetic and inducer at phase II; Auris Medical, AM101, an NMDA receptor antagonist, at phase II; Ipsen, a ginkgo biloba extract, at phase I; Merz, neramexane, an NMDA antagonist and an α9α10 nicotinic cholinergic receptor blocker [45], at phase III; and GSK, vestipitant, a neurokinin 1 receptor antagonist [46], at phase II.

From the above, it can be concluded that there are a few companies with tinnitus compounds in their pipelines in spite of the existence of such a huge market for this clinically unmet need. This most likely derives from the existing challenges described in the previous section. The lack of serendipitous discoveries of effective treatments for tinnitus has severely limited insight into disease pathology, which is often gained by such fortuitous pharmacological findings. It is the absence of a fully determined mechanism for tinnitus that makes research into this area potentially very high risk. However, if any of the above compounds reaches the market, they will set a turning point both in the treatment of tinnitus as well as in the development of future compounds.

**Potential Pharmacological Targets**

The search for drugs that target tinnitus is hampered by the lack of a deep knowledge of the underlying neural substrates of this pathology. Initially considered an inner ear pathology, it is now clear that at least chronic tinnitus is a central nervous system disorder. As indicated above, changes in cortical tonotopy, as well as increase in spontaneous firing rates and neuronal synchrony in different parts of the auditory pathway, have been proposed as potential correlates of tinnitus [1, 29].

After noise trauma induced hearing loss, one of the main causes leading to tinnitus, changes in tonotopic organization in the cortex are observed. Cortical neurons with characteristic frequencies in the frequency region of the hearing loss no longer respond according to their place in the tonotopic map, but reflect instead the frequency tuning of their less affected neighbors [47–49]. Magnetic source imaging studies confirm this reorganization in human patients [50]. This suggests that reorganization of the cortical tonotopic map and tinnitus are correlated. Interestingly, providing an acoustically enriched environment spectrally matching the hearing loss prevents this reorganization [51, 52]. Thus, preventing neuronal reorganization by an acoustically rich environment might become a treatment strategy to prevent the establishment of the long-term plastic changes that follow exposure to noise trauma. However, most clinicians are faced with the problem of treating tinnitus patients when tinnitus is most likely a chronic condition in which tonotopic rearrangements along the auditory pathway are already established. Can established tonotopic rearrangements in the auditory cortex be reversed? Experiments in laboratory animals that combine sound exposure with electric stimulation of certain neuronal pathways/circuits show promising results. In the primary auditory cortex, dopamine release has been observed during auditory learning that remodels the sound frequency representations [53]. The stimulation of dopaminergic neurons in the ventral tegmental area of rats, together with an auditory stimulus of a particular tone, increases the cortical area and selectivity of the neural responses to that sound stimulus in the primary auditory cortex while it decreases the representations of nearby sound frequencies [54]. In addition, episodic electrical stimulation of the nucleus basalis of rats, paired with an auditory stimulus, results in a massive progressive reorganization of the primary auditory cortex in the adult rat. Receptive field sizes can be narrowed, broadened, or left unaltered depending on specific parameters of the acoustic stimulus paired with nucleus basalis activation [55]. The nucleus basalis contains both cholinergic and gabaergic neurons [56, 57]. Thus, taken together, these results indicate that sound therapy coupled with drugs that can modulate the neurotransmission of the
pathways/circuits involved in the described plastic events would be an interesting avenue to investigate.

Additional neural correlates of tinnitus include neuronal spontaneous hyperactivity in the reorganized region and increased neural synchrony [48, 52, 58]. Neuronal hyperactivity can be modulated by many multiple drugs that target either voltage-gated ion channels or neurotransmitter receptors. However, examples of such drugs like benzodiazepines, anticonvulsants, NMDA antagonists, and calcium antagonists, although effective in some patients, have not proven effective in double-blind placebo-controlled clinical trials [23]. Recently, in a preliminary report using a rat behavioral model, the potassium channel modulator Maxipost (BMS-204352) reduced behavioral evidence of salicylate-induced tinnitus in a dose-dependent manner [59]. This compound is a KCa1.1 (BK) and a Kv7 positive modulator [60, 61]. Since potassium ion channels play an important role in regulating the resting potential and spontaneous and evoked neural activity, potassium channel modulators represent potential important compounds for tinnitus therapy.

The above are only some few challenging ideas concerning ways to revert altered neuronal activity, synchrony, and tonotopy observed in the auditory pathway in tinnitus. However, it is a reductionistic approach, since it only takes into account changes observed in the auditory pathway. As has been shown in the somatosensory system, auditory cortex activation is essential, but probably not sufficient for auditory conscious perception [62, 63]. Moreover, for most patients, tinnitus is more than mere changes in the auditory pathway and implicates the activation of a distress network [38–40]. This brings us back to the notion that central nervous system acting drugs, in particular, owe their clinical utility to actions at multiple molecular targets [31]. This is most likely the scenario we are facing in the search of a drug to alleviate tinnitus.

The Time is Right

For many years, the standard of care for dealing with tinnitus patients has been, “You need to learn to live with it.” Although we are far away from fully understanding tinnitus, the chances for a solution are much brighter than they were a decade ago. The development of behavioral measures of tinnitus in animals combined with physiological, biochemical, molecular, and imaging techniques are likely to provide important insights into the underlying causes of tinnitus. Tinnitus animal models will provide a way to screen for drugs that can suppress the disorder. The potential market for an FDA-approved drug to treat tinnitus is huge. Several existing drugs have been reported to provide significant relief from tinnitus in subsets of patients. Looking toward an exciting future, patients and clinicians may finally receive encouraging news if the compounds under development by several pharmaceutical industries finally reach the market. If they do, they will set a new era that will revolutionize the treatment of tinnitus.

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References


Chapter 31
The Neuroscientist

James A. Kaltenbach

Keypoints

1. This chapter reviews the current state of knowledge of tinnitus from the neuroscientist’s perspective.
2. Tinnitus is viewed as a disorder involving changes in the rate and timing of spontaneous discharges at multiple levels of the auditory system.
3. Its mechanisms vary, depending on etiology, but most commonly the disorder stems from increases in the excitability of neurons in the central auditory system.
4. Most of the available data suggest that this increase is synaptic in origin, caused by shifts in the balance of excitatory and inhibitory inputs to neurons.
5. However, other mechanisms, such as shifts in the expression of ion channels that determine the resting membrane potential of neurons, may also play a contributing role.
6. Since these changes occur at multiple levels of the auditory system, it is likely that new therapies that will prove most effective will be those that take a system-wide approach rather than those that target specific generator sites.

Keywords Tinnitus • Dorsal cochlear nucleus • Plasticity • Excitotoxicity • Neurodegeneration • Inferior colliculus • Auditory cortex

Abbreviations

DCN Dorsal cochlear nucleus
GABA Gamma amino butyric acid
IC Inferior colliculus
LTD Long-term depression
LTP Long-term potentiation
NMDA N-Methyl-d-aspartate
rTMS Repetitive TMS
TMS Transcranial magnetic stimulation

Introduction

Over the past 20 years, a great deal has been learned about tinnitus mechanisms from neuroimaging studies in humans and neurophysiological studies in animals. We now have substantial literature examining where and how activity in the auditory system is altered by tinnitus-inducing agents. Coupled with the growing number of behavioral studies demonstrating that animals develop tinnitus after exposure to various tinnitus-inducing agents, the available evidence provides us with compelling reasons to suspect that some of the reported changes in activity underlie the percepts of tinnitus. This chapter reviews the current state of knowledge of tinnitus from a neuroscientist’s perspective.

Is Tinnitus Primarily a Peripheral or Central Problem?

The term “ringing of the ears” implies that tinnitus is largely a problem of the ear. However, we now have a considerable body of evidence that the major changes underlying tinnitus can occur peripherally or centrally. House and Brackman [1] found that tinnitus persisted in 62% of patients in whom input to the brain from the auditory nerve was surgically abolished. In many of
these patients, the post-surgical tinnitus was worse than the pre-surgery tinnitus. Other studies have reported that tinnitus develops secondarily following surgical removal of eighth-nerve tumors (vestibular Schwannoma [1–3]), a procedure that can lead to major impairment of the auditory nerve. These findings point to the central auditory system as an important source of tinnitus, although there is little doubt that tinnitus in most cases begins with trauma in the auditory periphery. Thus, although agents such as noise or aminoglycoside, which cause hearing loss, often also cause tinnitus (see Chaps. 37, 42), they either have a weak long-term effect on spontaneous activity in the auditory nerve or cause this peripheral activity to decrease [4–6].

At the same time, it is important to acknowledge that in 38% of House and Brackman’s patients, tinnitus was abolished by eighth-nerve section. Although spontaneous activity is reduced following noise or aminoglycoside treatment, other alterations have been found in the auditory nerve, such as increase spontaneous bursting activity (see next section), which could potentially be tinnitus producing. Moreover, some studies suggest that sodium salicylate can cause increases in spontaneous activity and changes in the timing of spontaneous spikes in the auditory nerve that could generate tinnitus percepts [7–11]. Thus, it seems likely that some forms of tinnitus may originate peripherally, although, as discussed in the following section, most contemporary studies of tinnitus have focused on the central auditory system for the reasons given above.

**Neurophysiological Correlates of Tinnitus**

The most commonly reported effects of tinnitus-inducing agents on neurons in the auditory system are increases in spontaneous activity, bursting activity, and synchronous discharges. Chronic increases in spontaneous activity can be induced in the dorsal cochlear nucleus (DCN) [12–18], inferior colliculus (IC) [15, 19–26], and auditory cortex [27–31] using exposure or treatment conditions that have been shown in a variety of other studies to induce tinnitus in animals [17, 21, 32–39]. Increased spontaneous activity occurs in the IC and auditory cortex after salicylate treatment [40–45]. There is evidence for increased spontaneous activity in the DCN following treatment with cisplatin [44] (see Chap. 16) and in the auditory cortex following treatment with quinine [46]. Both salicylate and quinine have also been shown to cause tinnitus in animals at doses known from other studies to cause increased spontaneous activity in the auditory system [47–49].

That the increase in activity is likely to be perceptually sound evoking is supported by the following:

1. The hyperactivity displays similar spatial and temporal distribution patterns as increases in activity evoked by tonal stimulation.
2. It is well established from electrophysiological studies that increases in activity are observed throughout the central auditory system during sound stimulation, so there can be little doubt that sound percepts are linked to increases in discharge rates.
3. Cochlear and central auditory prosthetics are based on the notion that auditory percepts can be evoked by stimuli designed to increase discharge rates of auditory neurons.
4. Increased activation has been observed in the IC and auditory cortex of individuals with tinnitus [50–59].
5. Stimulation of the somatosensory system via the trigeminal nucleus or cervical nerves modulates both spontaneous activity in central auditory centers [60–63] and tinnitus [64–67]. Taken together, these findings give strong support to the view that tinnitus is linked to changes in discharge rates in the central auditory system.

However, just increased discharge rates, per se, may not be the whole story. Noise exposure, and salicylate cause increases in a specific type of activity called bursting discharges in the auditory system. Chronic increases in bursting activity have been observed in the auditory nerve following noise exposure [6], in the DCN following noise exposure [14], and in the IC following salicylate and noise exposure [23, 41]. No increased bursting has been found in the auditory cortex following noise exposure, salicylate, or quinine [23, 46, 68, 69]. Increases in bursting activity, even if limited to the auditory brainstem, may be an important correlate of tinnitus. Bursts of spikes carry an important feature that is likely to signal the presence of sound, namely, periodicities, and brief clusters of spikes with nearly identical interspike intervals. Of these, periodicities in firing are critical to the ability of neurons to encode the frequency of sounds [70, 71]. If bursting is increased, then periodicities in a restricted frequency range would probably also be
increased, and this could lead to perception of a tinnitus-like sound in a correspondingly restricted pitch range.

In addition to increased discharge rates and increased bursting activity, there is evidence for an increase in synchrony of discharges among neurons in the IC following noise exposure [23] and in the auditory cortex following noise or quinine administration [27, 30, 69]. Increased synchrony of auditory nerve fibers following salicylate treatment is suggested by increases in the amplitude of 200 and 900 Hz peaks in the frequency spectrum of ongoing ensemble activity [9–11]. This means that instead of impulses being more or less randomly related across the neural population, the impulses become increasingly coincident. This is sometimes referred to as temporal coherence (see Chaps. 12 and 13). Neurons showing increased synchrony occur in frequency bands of the hearing loss that are also the areas in which tonotopic map reorganization occurs. Increased synchrony has been hypothesized to be a neural correlate of tinnitus [72, 73] (see Chaps. 12 and 13). Pitch percepts corresponding to frequency regions with increased synchrony might be enhanced, leading to the often pitch-like percepts of tinnitus.

In summary, central auditory nuclei and cortical areas develop some of the types of changes following cochlear trauma that are also evoked by acoustic stimulation. Issues that will be addressed next are what the underlying triggers of changes in spontaneous activity might be as well as what mechanisms underlie their induction.

**The Triggers of Tinnitus-Related Activity**

**The Role of Deafferentation**

Tinnitus is often viewed as a deafferentation disorder triggered by loss of normal input from the auditory periphery. Evidence for a deafferentation mechanism of tinnitus comes from a wide range of clinical and experimental observations. Tinnitus is most commonly associated with hearing loss. Between 80 and 90% of tinnitus patients have an associated hearing loss [74] (see Chap. 5). Tinnitus can be induced by surgical damage to [75, 76] as well as compression or tumors of the eighth nerve [2, 3, 77, 78] (see Chap. 39). Tinnitus is also sometimes seen in association with conductive hearing loss [79–81] (see Chap. 83). All these conditions involve impairment of peripheral auditory functions, so there is good reason from human observations alone to suspect that loss of peripheral function and peripheral input are key triggers of tinnitus. Animal models have also yielded evidence consistent with a deafferentation-induced mechanism of tinnitus. Tinnitus percepts in animals and tinnitus-related changes in activity in the IC have been found to be associated with loss of spiral ganglion cells [23]. The induction of tinnitus-related hyperactivity in the dorsal cochlear nucleus has been found to be correlated with loss of outer hair cells [45]. This is consistent with reports that tinnitus is often found to be associated with defects in outer hair cell function, as reflected by alterations of transient-evoked or distortion product otoacoustic emissions (see review of [82]). It has been hypothesized that loss of outer hair cells may induce hyperactivity in the dorsal cochlear nucleus by causing loss of peripheral input to the granule cell system [45]. This hypothesis builds on the facts that the granule cell domain in the cochlear nucleus receives input from type II spiral ganglion neurons, which originate from outer hair cells [83–85], and there is some evidence that granule cells are among the recipients of type II input [86]. Moreover, activation of granule cells influences the level of activity of the principal cells of the DCN, the likely generators of tinnitus signals [13, 41, 60, 87].

Deafferentation can also involve loss of input to auditory structures from non-auditory areas. This possibility is raised by the fact that many subjects with tinnitus possess disorders of other systems. For example, many cases of somatic tinnitus (such as that experienced by people who can change the loudness or pitch of their tinnitus by manipulations of head and neck musculature) occur in people with somatic pathologies of the head and neck, including craniofacial anomalies, temporomandibular joint disorders, or inflammatory conditions of the neck muscles [64, 65, 88]. Furthermore, Levine [65] found that in his patients with somatic tinnitus, when the tinnitus was monaural, it was usually on the same side as the somatic disorder. Lastly, an increasing number of articles suggest that tinnitus can be induced or exacerbated by emotional conditions such as stress and anxiety [89–91]. There are several levels of the auditory pathway where auditory centers receive input from non-auditory areas. The best described example, in terms of circuitry, is the dorsal cochlear nucleus, whose output is modulated
by the cochlear granule cell system. This system receives input not only from auditory sources but also from cuneate and trigeminal nuclei and ganglion of the somatosensory system [61, 92–94] (see Chap. 9) and a variety of other pathways [87]. Since activation of the granule cell system is known to affect the level of spontaneous activity [13, 60, 95], conditions in which inputs from these areas are impaired or damaged could affect output of the dorsal cochlear nucleus via their effects on the granule cell system.

The Role of Plasticity

There are two general mechanisms by which deafferentation might induce tinnitus-related activity in the central auditory system by activating neural plasticity (see also Chaps. 12 and 13). The most frequently hypothesized mechanism is a shift in the balance of excitatory and inhibitory synaptic inputs to central target neurons toward the side of excitation. Such a shift could involve direct loss of inhibitory inputs (disinhibition) and/or an increase in excitatory inputs.

Several lines of evidence indicate that both a loss of inhibition and an increase in excitation occur centrally after loss of auditory nerve input and that such changes involve plasticity. First, loss of primary afferent input leads to loss of inhibitory influence in brainstem auditory nuclei, as signaled by reductions in glycinergic and GABAergic neurotransmission [96–104]; these reductions change over time, suggestive of a temporal or possibly homeostatic plasticity mechanism [105]. Second, there are suggestions of up-regulations of excitatory synapses — for example, cochlear ablation, noise exposure, and conductive hearing loss trigger up-regulations of cholinergic and glutamatergic systems in the central auditory systems [106–113]. Some of these adjustments vary over time. Third, degeneration of second-order neurons in the brain following noise exposure [114] is followed by regrowth of excitatory and inhibitory terminals, but a more complete return of excitatory than inhibitory synapses, indicating a reorganization of synaptic connections that favors excitation [115].

A second mechanism that could lead to tinnitus-related activity is an increase in excitability of neurons caused by alterations in their intrinsic membrane properties. Such alterations might involve up- or down-regulations of specific ion-conductance channels. Studies pointing to changes in the intrinsic membrane properties of cochlear nucleus neurons following cochlear deafferentation have been published. Cochlear ablation was found to cause increases in membrane resistances of neurons in the ventral cochlear nucleus (Francis and Manis, 2000). Hearing impairment has also been found to be associated with decreases in the expression of the two-pore domain potassium channels and reductions of Kv3.1 channels in central auditory neurons [116, 117]. Changes in spike waveform have been observed in the dorsal cochlear nucleus after noise exposure [14]. The relationship between these changes and alterations in spontaneous activity has not yet been determined.

Non-deafferentation Triggers of Tinnitus Induction

Deafferentation is not the only triggering mechanism by which tinnitus-related activity could be induced. Some inducers of tinnitus may act through non-deafferentation mechanisms, such as excitotoxicity or activity-dependent plasticity.

Excitotoxicity

Excess release of excitotoxic neurotransmitters in the brain caused by acoustic overstimulation could lead to degeneration of second-order neurons, many of which may be inhibitory. Glutamate is the most common excitatory and most powerfully excitotoxic neurotransmitter in the nervous system. It is also the excitatory transmitter of hair cells, auditory nerve fibers, granule cells of the cochlear nucleus, and the main projection neurons that make up the ascending auditory pathway. Normally, toxicity of this transmitter is prevented by its reuptake following its release by the presynaptic membranes. However, under certain conditions, such as when there is excessive sound stimulation, glutamate is released in excess, and this excess can sometimes overwhelm the reuptake mechanism. This leads to its accumulation in the synaptic cleft. Excess glutamate binds to N-methyl-D-aspartate (NMDA) receptors, which stimulates excess calcium influx into postsynaptic neurons via the calcium channels of NMDA receptors; the excess calcium stimulates intracellular enzymes that are damaging to cells and can culminate in apoptosis.
A case for excitotoxicity acting through excess glutamate release in the auditory system is suggested by the following: Overstimulation would be expected to cause excess release of glutamate from excitatory terminals in and beyond the cochlear nucleus. An increase in glutamate release and a decrease in glutamate uptake have been found to occur in the cochlear nucleus and persist for at least 5 days following acoustic overstimulation [110]. This would be expected to result in an accumulation of glutamate in the synaptic cleft and thereby trigger excitotoxic injury. Evidence consistent with this hypothesis is the finding that degeneration occurs in broad areas of the cochlear nucleus well beyond zones of peripheral deafferentation [114, 118]. These findings have been interpreted as possibly resulting from excitotoxic injury in the central auditory system [110, 118]. The loss of second-order neurons by this mechanism would be expected to shift the balance of excitation and inhibition in the central auditory system in ways that could be tinnitus inducing.

Activity-Dependent Plasticity

One of the most commonly described mechanisms by which synaptic excitability of neurons is chronically shifted is long-term potentiation (LTP). This is a long-lasting enhancement in synaptic transmission between two neurons that results from stimulating them synchronously. LTP results in a sensitization of neurons to their inputs, which is manifest as an augmentation in the response of the postsynaptic neuron to its excitatory inputs. Another manifestation of LTP is an increase in spontaneous activity [119]. If LTP occurs in the auditory system, it seems likely that the affected neurons would become hypersensitive and spontaneously hyperactive. A related, but opposing process is long-term depression (LTD), which is manifest as a reduction in the response of neurons to their inputs. These activity-dependent phenomena were originally discovered in the hippocampus and have been implicated as neural mechanisms of long-term memory. They are now known to be ubiquitous throughout the brain.

The question at hand is whether inducers of tinnitus can cause LTP in auditory neurons. There is evidence that LTP can be induced in various auditory centers by synchronous stimulation of pre- and postsynaptic neurons. LTP has been demonstrated by this method in the dorsal cochlear nucleus [120–122], inferior colliculus [123, 124], and auditory cortex [125, 126]. Thus far, it is not known whether tinnitus inducers can cause LTP in these same brain areas. However, it has been hypothesized that noise might increase the probability of synchronous firing of pre- and postsynaptic firing and thereby cause induction of LTP [127]. This possibility seems plausible since acoustic stimuli increase the frequency of firing and the occurrence of coincident spikes in the auditory system [29]. Induction of tinnitus by LTP and excitotoxicity offers an explanation of why tinnitus often occurs without any accompanying hearing loss.

Why Tinnitus Does Not Always Accompany Hearing Loss

If tinnitus is the result of increases in neuronal activity (increased discharge rate and bursting) and/or increased synchrony triggered by loss or overstimulation of afferent input to the auditory centers of the brain from the ear, and also possibly involving non-auditory inputs to these centers, then why do many people with hearing loss have no tinnitus? [128, 129] The simplest explanation is that the direction of the shift in the balance of excitation and inhibition following cochlear injury may depend on the pattern of cochlear injury. Tinnitus induction would be expected to occur when there is more degeneration centrally of inhibitory than excitatory neurons, causing disinhibition and an increase in excitation. However, it is conceivable that certain patterns of peripheral injury may not be sufficient to shift the balance of excitation and inhibition or could even favor a shift toward the side of greater inhibition. Support for this concept is demonstrated by the finding that tinnitus-related hyperactivity is initially absent following induction of noise-induced threshold shift but emerges slowly over several days following the noise exposure, only after a transient decline of activity [16]. Moreover, it has been shown that when cochlear injury induced by cisplatin is restricted to outer hair cells, there is a strong relationship between the degree of centrally recorded hyperactivity and the amount of outer hair cell loss, but when the outer hair cell loss is accompanied by mild damage to the inner hair cells, particularly disarray of their stereocilia, activity is not elevated centrally. However, when the inner hair cell injury becomes
more severe or outer hair cell loss is accompanied by inner hair cell loss, hyperactivity is clearly apparent [45]. This suggests that the effect of peripheral injury on central auditory activity depends on the balance and type of injury to the two hair cell populations and their connecting primary afferents.

Implications for Tinnitus Treatment

The state of knowledge on tinnitus mechanisms has provided a much-needed theoretical framework for conceiving and testing new therapeutic treatments for tinnitus over the past decade. Among the various modalities that have received the most attention are drug therapy, electrical stimulation, and transcranial magnetic stimulation. Efforts also continue to improve treatment through sound therapy and psychological counseling.

Drugs that are attracting interest as potential tinnitus agents are those that decrease neural activity. Initial studies with gabapentin were suggestive of a tinnitusolytic effect in animals and some human subjects [130]. However, more recent clinical trials showed that when the effects are compared with placebo across a sample of patients, no significant difference was observed [131, 132]. Thus, if gabapentin has a tinnitusolytic effect, it may be that only a small proportion of patients who have been treated with gabapentin experience benefit. Agents that activate the inhibitory receptors for GABA$_\alpha$ and GABA$_\beta$ receptors (e.g., benzodiazepine and baclofen, respectively) have been found to have a suppressive effect on tinnitus-related activity in animals [133, 134]; studies with these agents in clinical trials have yielded mixed results. While baclofen was not found to have a significant effect on tinnitus [135], there are indications that administration of benzodiazepines, benefits many patients suffering from tinnitus [136] (see review of Gananca et al. [137]) (see also Chap. 30). In some patients, the benefit may be achieved primarily by reducing the severity of the emotional reaction to tinnitus, but there is usually a subgroup that also experiences a decrease in the loudness of tinnitus.

There has been growing interest in targeting NMDA receptors, which are implicated in plasticity for tinnitus treatment. The data thus far are preliminary, but there are indications that NMDA receptor antagonists (acampresate, caroverine, ifenprodil) have tinnitusolytic effects in animals [36, 138–140]. Preliminary results suggest that the NMDA receptor antagonist, neramexane, may reduce tinnitus-related activity in the DCN of animals [141]. A recent clinical trial with neramexane yielded results suggestive of a significant tinnitusolytic effect in human subjects [142]. The drug is now being tested in a phase III clinical trial: http://clinicaltrials.gov/ct2/show/ (NCT00405886) (see also Chaps. 22 and 30).

Electrical stimulation studies have been conducted in areas of the brain that have been implicated as sites of tinnitus generation. The benefits have been most remarkable for patients stimulated at the cochlear level, either transtympanically or intracochlear using a cochlear implant [143, 144] (see also Chap. 77). Stimulation of the dorsal cochlear nucleus using the auditory brainstem implant has been found to be effective in suppressing tinnitus [145], and there are some recent indications that stimulation of the auditory cortex can suppress tinnitus [146–148].

Another approach that has generated considerable interest is repetitive transcranial magnetic stimulation (rTMS). This procedure is used primarily to stimulate the auditory cortex or nearby areas (see Chap. 88). A recent review of the literature [149] concluded that rTMS is a promising approach for the treatment of patients with certain forms of tinnitus. At present, the results of both stimulation modalities vary significantly across studies and within studies across individuals. This variability may stem from differences in stimulus parameters, differences in what parameters are optimal for each patient, and differences in the precise location of the stimulating electrode(s) or magnetic field relative to the primary generator sites giving rise to the tinnitus-producing signals. The fact that tinnitus has many forms (see Chap. 2) also contributes to the variability in the results of treatments. However, the findings provide a proof of concept that stimulation of auditory areas can, under optimal conditions, bring considerable relief to a significant number of tinnitus patients.

Summary and Conclusions

The foregoing review of tinnitus summarizes the areas of the nervous system that display activity changes believed to underlie the percepts of tinnitus. The available
evidence indicates that tinnitus is associated with more than one type of change in the auditory system. At the brainstem level, increases in bursting and non-bursting spontaneous activity are clearly demonstrable after noise exposure and salicylate treatment, while at the cortical level, increases in non-bursting spontaneous activity and neural synchrony are more apparent. The literature review also indicates that tinnitus of different etiologies likely involves different structures and possibly different mechanisms. This is best demonstrated by clinical studies showing that sectioning the eighth nerve sometimes alleviates tinnitus, but more commonly tinnitus persists and is often worsened following this procedure. This suggests that there may not be a single final common path for tinnitus and supports that there are many forms of tinnitus (see Chap. 2).

Another important concept is that tinnitus of central origin emerges as a consequence of activation of neural plasticity, which alters the excitability of neurons, primarily by shifting the balance of their excitatory and inhibitory inputs, but also possibly by shifting the balance of ion channels that control the resting membrane potential.

Our current state of knowledge provides a useful framework for developing new therapeutic approaches to tinnitus treatment. The multi-tiered distribution of tinnitus-related changes suggests that the most effective treatments for tinnitus will be those that take a system-wide approach rather than those that target specific structures. Therapies that quiet resting activity throughout the auditory system without lowering the activity of other brain pathways and without compromising sensitivity to sound will bring the type of benefits desired by most patients with tinnitus. A demonstration that such effects can be achieved on a short timescale is already indicated by the brief periods of tinnitus suppression provided by residual inhibition, somatic modulation of tinnitus, and, in some cases, by lidocaine. The goal now is to exploit these mechanisms further to increase the duration of the suppression to bring a longer lasting period and possibly chronic state of relief from tinnitus. With the foundation presently in place, we have good reason to expect that this knowledge will lead to major improvements in the treatment of tinnitus.

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References

32. Heffner, HE, Harrington, IA, Tinnitus in hamsters following exposure to intense sound. Hear Res, 2002;170:83–95
36. Guitton, MJ, Dudai, Y, Blockade of cochlear NMDA receptors prevents long-term tinnitus during a brief consolidation window after acoustic trauma. Neural Plast, 2007;2007:80904


60. Kanold, PO, Young, ED. Proprioceptive information from the pinna provides somatosensory input to cat dorsal cochlear nucleus. J Neurosci, 2001;21:7848–7858


70. Evans, EF. Place and time coding of frequency in the peripheral auditory system: some physiological pros and cons. Audiology, 1978;17:369–420


72. Eggermont, JJ. Pathophysiology of tinnitus. Prog Brain Res, 2007;166:19–35


74. Vernon, JA (Ed), Tinnitus: Treatment and Relief. Boston, MA: Allyn & Bacon, 1997


78. Møller, AR, Møller, MB. Microvascular decompression operations. Prog Brain Res, 2007;166:397–400


89. Seydel, C, Reishauer, A, Hau, H, Klapp, BF, Mazurek, B. The role of stress in the pathogenesis of tinnitus and in the ability to cope with it. HNO, 2006;54:709–714


92. Wright, DD, Ryugo, DK, Mossy fiber projections from the cuneate nucleus to the cochlear nucleus in the rat. J Comp Neurol, 1996;365:159–172
96. Suneja, SK, Potashner, SJ, Benson, CG, Plastic changes in glycine and GABA release and uptake in adult brain stem auditory nuclei after unilateral middle ear ossicle removal and cochlear ablation. Exp Neurol, 1998;151:273–288
111. Suneja, SK, Potashner, SJ, Benson, CG, AMPA receptor binding in adult guinea pig brain stem auditory nuclei after unilateral cochlear ablation. Exp Neurol, 2000;165:355–369
121. Oertel, D, Young, ED, What’s a cerebellar circuit doing in the auditory system? Trends Neurosci, 2004;27:104–110
122. Tzounopoulos, T, Kim, Y, Oertel, D, Trussell, LO, Cell-specific, spike timing-dependent plasticities in the dorsal cochlear nucleus. Nat Neuroscience, 2004;7:719–725
123. Wu, SH, Ma, CL, Sivaramakrishnan S, Oliver DL, Synaptic modification in neurons of the central nucleus of the inferior colliculus. Hear Res, 2002;168:43–54
134. Zhang, JS, Kaltenbach, JA, Effects of GABAB receptor activation on sound-induced hyperactivity in the DCN of hamsters in vivo. In: ARO Abs, 2000 #5020
137. Gananca, MM, Caovilla, HH, Gananca, FF, Gananca, CF, Munhoz, MS, da Silva, ML, Serafini, F, Clonazepam in the pharmacological treatment of vertigo and tinnitus. Int Tinnitus J, 2002;8:50–53
In my wish to perhaps help others suffering from tinnitus and the constant ringing and hissing in both ears since March 16, 1996, I thought it would be appropriate to provide a historical background on how this intractable problem came about in my life. As those who are suffering from tinnitus, I am aware it is a symptom and not a disease and that it can be brought about from a number of variable causes that include: (1) use of excess alcohol, (2) caffeine, (3) aspirin in heavy doses, (4) certain medications, (5) hardening of the arteries, (6) high blood pressure, (7) infection of the ear canal or eardrum, (8) Ménière’s disease (inner ear disorder), and (9) exposure to loud noise(s). In my particular case, the latter is the cause of my tinnitus.

Let me first describe my upbringing. Upon graduation from Washington and Jefferson College in June of 1955, I was commissioned as Second Lieutenant in the United States Army in connection with having enrolled in ROTC (a 4-year Reserve Office Training Corp). In that same month and year, I was posted at the United States Transportation School at Ft. Eustis, Virginia a short distance from Williamsburg, Virginia. In February of 1956, I was assigned to serve as a part of the post-war occupation of then West Germany and stationed at the United States seventh Army Headquarters near the city of Stuttgart. Although I was attached to a Transportation Unit, we were required to take part in maneuvers every 3 months. On those occasions, as an officer, I carried a 45-caliber pistol and was required to fire it on the firing range. On other occasions, I was designated the “firing line officer” on the firing range to supervise the firing of 50-caliber machine guns by enlisted men in our unit. Occasionally, I also fired the 50-caliber machine gun. Knowledge of fire power and the use of weapons was necessary because, as I recall, we were informed of the possible menace by the Russian Army, which had divided the country into East and West Germany. In any event, I must say that the noise from firing the 45-caliber pistol and alternately firing the 50-caliber machine gun was deafening. It must also be remembered that at that time, we had no ear protectors or any other device(s) to shield us from the horrendous noise. After completing a few rotations as “firing line officer” I noticed some “ringing” in my right ear; although it was slight, I reported it to my Company Commander.

He suggested that I make an appointment with the resident Army physician, who, after a quick examination and test, stated that I had a 10/11% hearing loss but, did not issue any Order excusing me from being on the firing range. I mentioned this encounter with the military physician because the examination was so casual and did not address my problem with the onset of tinnitus. However, I was so troubled that a continuation of exposure to extremely loud noise on the firing line would aggravate the tinnitus that I again approached my Company Commander who shared my concerns. Accordingly, he issued an Executive Order excusing me from the firing range altogether.

After that, I was assigned to the Motor Pool for the remainder of my tour until June 1957 when I received an Honorable Discharge as a First Lieutenant. Thereafter, although the mild tinnitus continued in my right ear, it was more of an irritating problem but, more importantly, did not interfere with my studies at the University of Washington and Lee Law School (Lexington, Virginia), where I was admitted shortly after my discharge from the Army.
I should note, in particular, that I was not taking any medication for the mild tinnitus in my right ear. In other words, I led a normal life and had no health problems since my discharge in June of 1957. However, to my great misfortune, all that changed on the night of March 16, 1996. On that night, as a solicitor for a local municipality, I attended a meeting of the Planning Commission to review a plan for a real estate developer who was seeking a special exception of the building code for the construction of apartments in the municipality. After a lengthy review of the plans, the meeting was adjourned.

As I was about to enter my car, the fire sirens on the tower next to the municipal building sounded off and since my car doors were locked, my only option was to place my fingers in each ear in hopes of diminishing the extremely loud sirens. All I could do was wait until the sirens turned off and then enter my car because I had no other choice. In other words, had I attempted to open my car door I would have exposed my ears to even louder noise. I knew immediately that my tinnitus was greatly increased by the exposure to the sirens because as I drove home, the ringing was louder and was now not only the right ear but also in the left ear, which previous to this incident was absent of any tinnitus. When I arrived home, my wife asked me why my face was so ashen and I related the above incident. I also told her that in the few seconds that I was exposed to the screeching of these sirens, loud hissing in both ears was immediately noticeable. I was not only devastated but fell into deep depression right away because, in my opinion, the mild tinnitus prior to this incident was forever aggravated.

As a result of the increased tinnitus, now in both ears, I could not sleep. At the suggestion of my wife, I took a couple of aspirins, but this did not reduce the loud tinnitus at all nor did it help me to fall asleep. In the morning, after a completely restless night, I was frantic and wanted to see a doctor as soon as possible. My wife called a family friend who recommended an eminent otologist. When my wife called the otologist’s office, she was informed that the doctor was attending a conference in Europe. However, after explaining the urgency of the matter, the office secretary scheduled an appointment to be seen immediately after her return in 2 weeks time. In the interim, I called another otologist whose office made an appointment for me. At that visit, I related the circumstances of exposure to the sirens to the doctor and he proceeded to conduct an examination. At the end of the exam, he gave me a prescription for Zoloft to help me cope with the tinnitus. That particular doctor advised me to return to my law office and “work on a brief and forget the tinnitus.” I responded that with constant loud hissing in both ears, I could not return to my law practice since I could not concentrate on my work, and furthermore, I was extremely depressed and disheartened about my condition. Incidentally, the Zoloft I was taking at the time caused me to feel dazed and weak. Upon the return of the otologist recommended by my family friend, an appointment was confirmed. Upon my arrival, I underwent a battery of tests to determine if there was any loss of hearing arising out of exposure for the loud sirens. The tests revealed that there was some loss of hearing, especially at high-pitched sounds. However, my hearing was not greatly diminished. Also during that visit, it was decided that I should cease taking Zoloft, and Xanax was prescribed instead.

I was also advised that it might be helpful and therapeutic to see a psychologist. I was agreeable to this approach and an appointment was made with a psychologist at a nearby geriatric center. This course of action was followed for approximately 4 months. However, since the otologist concluded that this regimen was not as helpful as she had anticipated, she recommended a consultation with one of the most eminent psychiatrists at a psychiatric hospital. I must say that this particular psychiatrist was very helpful because I found him to be sincerely interested in my plight. His candid and positive assessment of my depression occasioned by my tinnitus was helpful as well. Despite those efforts, my otologist determined, through many discussions, that my depression was so deep and pervasive that I might harm myself. Accordingly, in early December of 1996, she informed me that the department of neurology at a university hospital was performing experimental surgery (referred to as microvascular decompression of the eighth cranial nerve) for those who were suffering from loud tinnitus. Although the outcome of that particular surgery was uncertain and indeed questionable, the otologist suggested that it might be a matter that I might consider in view of my intractable tinnitus. Moreover, she concluded that emotionally, I was a suitable candidate for this procedure. As I recall, I met with the surgical team to thoroughly discuss the surgery. I also remember the admonition of the chief surgeon, specifically, that the surgery (1) may reduce the tinnitus, (2) may make
the tinnitus louder, (3) may cause a loss of hearing altogether, or (4) may not accomplish anything at all. Naturally, after hearing these, fear and anxiety swept over me but, taking into consideration the debilitating and constant hissing in both ears, it was necessary for me to consider the above scenario and to make a decision about the proposed surgery. My wife and I deliberated about this and ultimately I harkened back to my days in law school to what I remembered in my class on the subject of negligence to convince me to go forward with the surgery. The rationale in one of the cases under study had to do with how one should approach a serious problem, which was enunciated as follows: “when embarking on a course of conduct, one must weigh the magnitude of the risk(s) against the utility of the conduct.” If the risk(s) outweighs the particular conduct contemplated, then the utility of the conduct must be abandoned.

“On the other hand, if the utility of the conduct outweighs the risk(s), then one must proceed with the conduct.” In my decision, which was supported by my wife, we concluded that the hope we placed in the surgical procedure outweighed the risk(s) of potential harm because of the intractable and debilitating condition arising out of the loud and unceasing ringing in both my ears. In this context, we believed we were making the correct decision in going through with the microvascular decompression surgery of the eighth cranial nerve. Thus, in late December 1996, the surgical team scheduled me to undergo a microvascular decompression operation for my left eighth nerve. The decision to operate on the left side was made by the neurological team on the premise that since it was the side which was damaged most recently, the chance of any success would be more likely than the right side, where the tinnitus first surfaced while I was on military duty in May 1956.

The surgery began with an incision just below the hairline behind the left ear as muscle and fascia were dissected. Then a burr hole was placed and extended with the craniectomy extending to the skull floor. The dura was then opened to view the eighth nerve. Finally, a piece of Teflon felt was interposed between the nerve and vessel.

I was then taken to the recovery room and when I gained consciousness, it felt like “a ton of bricks” (emphasis supplied) had struck my head. Fortunately, nurses stayed with me (in 12-h shifts), and in 4 days I was discharged.

After the surgical procedure, I had follow-up visits with various members of the surgical team as well as the otologist. At first, it appeared to me that the tinnitus had diminished slightly but after a few months, the tinnitus had returned to the level prior to the surgery. Of course, I was quite disappointed. However, I had been advised by the surgical team that this was one of the likely results of the surgery.

I consulted with my otologist about the current state of my condition and reported that I still had difficulty sleeping due to the constant hissing in both ears.

She then recommended that I purchase a “Sound Soother” with a timer. I mention this because this machine or device replicates different sounds such as water fall and the rush of ocean waves. This helped me to begin sleep, although to a small degree, but in a few hours I could hear the hissing again, which thus disturbed my sleep to such an extent that I abandoned the “Sound Soother.” When I returned to my law practice a few months after the surgery, on the advice of my otologist, I purchased a set of head phones under the name of “Viennatoners” (operated by batteries) to wear during the daytime hours and at work.

Although these “head phones” somewhat “masked” the tinnitus, I found them not only impracticable (it was difficult to make or accept office phone calls), but also of little benefit in attempting to manage the tinnitus, so I abandoned this approach after a few months.

In the meantime, even though I had not experienced any reduction of tinnitus incident to the microvascular decompression operation, I met with the surgical team and my otologist to seek their opinion about undergoing a second surgical procedure on the right side because the tinnitus was unbearable. After taking another series of tests, it was determined once again that I was physically and mentally able to tolerate such a surgical procedure. Thus, in early June of 1997, I underwent another microvascular decompression operation – this time of the eighth cranial nerve on the right side. One might question why I would subject myself to another operation when the first one was not successful. The answer, to me at least, was quite ordinary and simple, that is to say that due to suffering tinnitus 24/7, I was rather desperate and willing to accept the risk inherent in such a surgical procedure. Despite all the pain and suffering from this operation, I came out of it without any side effects and anxiously waited for some positive result even though I had...
tinnitus on that side since my military days. However, after a number of months, I was again disappointed that there was no degree of reduction of the tinnitus.

I continued to visit my otologist to take hearing tests, which indicated that although I was extremely sensitive to high-pitched sounds (termed Hyperacusis), my hearing was not diminished as a result of the tinnitus. It was of some comfort to know that my hearing was quite good under the circumstances. It should be noted that I continued visiting my psychiatrist, only instead of visits every 3 months, my appointments have been reduced to twice a year. I find, even at this present time, that the sessions were of immeasurable benefit to me. During the visits to the psychiatrist for the past 14 years, newer medications have helped me manage my tinnitus.

I currently take Effexor to calm my nerves during the day and Clonazepam has been the drug of choice before I go to bed because it begins my sleep cycle and I am able to get at least 6 h of sleep without hearing the hissing sound of tinnitus. During the early days of my tinnitus, a group of us suffering from the same nagging disability would call each other to offer comfort and support. Indeed, I found that some in our group who were experiencing very loud tinnitus were taking Clonazepam during the day and night because it was so debilitating. One unfortunate lady, who was in our group, was suffering so much from loud constant ringing and hissing (bi-laterally) that she could not sleep at all and after a number of months, in ultimate desperation, took her own life.

Of course, I must also say that suffering from the intractable effects of tinnitus has changed my lifestyle dramatically. Thus, I must be careful to avoid noisy venues and places. As a consequence, since the onset of aggravation of my tinnitus in 1996, I have not attended the symphony, movies, concerts, weddings, sports events, and crowded restaurants. Even a set of keys accidentally dropped in our kitchen, the banging of a door, or the whirling of a mixer all cause heightened noise, which, in turn, increases the hissing ring in each ear. On some such occasions, it seems to be especially loud and there is no relief until I go to bed with the aid of Clonazepam. Nor have I been able to utilize my lawn mower or my snowplow. Not surprisingly, because of the tinnitus, I have found myself speaking to friends, colleagues, and others in a low tone and invariably, they ask me to speak louder. On occasions when I cannot avoid noise (loud church choir for example), I find it necessary to place foam earplugs (which I always carry with me) in my ears for some protection. As one can see, my quality of life has been substantially changed due to my tinnitus. On the other hand, I am fortunate to have a patient and supportive wife, an otologist whose specialty in the study of tinnitus has given me encouragement to be positive and a psychiatrist who has provided me with years of counseling and drug therapy, all of whom have given me hope to continue my fractured life as best I can under stressful circumstances.

If I may offer some gratuitous advice to anyone who suffers from tinnitus, I would first urge him or her to never surrender to this nagging problem. I would also encourage them to be their own advocate by reading the vast information available in books, periodicals, and medical journals on the subject of tinnitus. The reservoir of information on the Internet is also a tremendous source of knowledge on the subject.

Lastly, I would counsel that if one is struggling with tinnitus, a specialist in otology be sought and not, with due respect, merely a family practice physician.

I am hopeful that the recitation of events that have occupied my life since June of 1996 may be of comfort and benefit to others who are suffering from this persistent and unyielding malady.
Part III
Causes of Tinnitus
There are many causes of tinnitus, and the origin of many incidences of tinnitus is unknown. A common cause of severe tinnitus is deprivation of signals to the central auditory system (see Chap. 11). This may occur through conductive hearing loss, cochlear hearing loss, and pathologies of the auditory nerve. The relation between tinnitus and hearing loss is discussed in Chap. 35 (see also Chaps. 8 and 10).

Only a few incidences of tinnitus are direct consequences of pathologies of the ear. Tinnitus is more frequently caused by deprivation of input to the central auditory pathway because of hearing loss. Many authors agree that the initial cause of tinnitus, when accompanied with hearing loss, has a peripheral origin that may trigger a series of reactions in the central nervous system, resulting in tinnitus. The anatomical location of the pathology may be the cochlea, but it is often the auditory nervous system (see Chap. 10). The finding that both deaf people and individuals with normal hearing can have severe tinnitus clearly indicates that tinnitus is not always linked to hearing loss.

Restoration of hearing can often also reduce the tinnitus, provided the restoration occurs before the tinnitus has been allowed to continue over a long period of time.

This section describes the possible causes of conductive hearing loss located in the external ear and middle ear (Chap. 34). Most incidences of tinnitus occur together with sensorineural hearing loss such as age-related hearing loss (presbycusis (Chap. 36) and noise-induced hearing loss (Chap. 37). Complications in medical treatment can cause tinnitus, such as through ototoxic antibiotics and cytostatic (Chap. 42). One of these symptoms that define Ménière’s disease is tinnitus (Chap. 38). Surgical manipulations of the auditory nerve can cause tinnitus (Chap. 40). Pathologies that affect the auditory nerve, such as vestibular schwannoma (Chap. 39), are often accompanied by tinnitus, as are surgical trauma to the auditory nerve. Pathologies that affect other parts of the auditory nervous system, such as cerebrovascular diseases (Chap. 41), can cause tinnitus but such cases are rare. The somatosensory system is involved in some forms of tinnitus (Chap. 43), and disorders such as those affecting the temporomandibular system are often accompanied by tinnitus (Chap. 44).
Chapter 34
Conductive and Cochlear Hearing Loss

Tobias Kleinjung

Keypoints

1. Any kind of hearing loss may be accompanied by tinnitus.
2. This chapter describes possible causes of conductive hearing loss located in the external ear and middle ear.
3. Pathologies of this area include neoplastic changes (e.g., tumors), inflammatory disease (e.g., otitis media), or disorders of unknown origin (e.g., otosclerosis).
4. Cochlear hearing loss of genetic origin can be classified in to syndromic and non-syndromic forms.
5. Labyrinthitis can occur due to bacterial or viral infection or in the context of immunological disease.

Keywords
Hearing loss • Inner ear • Middle ear • Otitis media • Otosclerosis • Tinnitus

Abbreviations
AIED Autoimmune inner ear disease
NSHL Non-syndromic hearing loss
SHL Syndromic hearing loss

Introduction

Virtually any pathology involving the ear appears to have the capacity to cause hearing loss and tinnitus may accompany the hearing loss at any time. Pathology involving the external ear and middle ear leads to conductive hearing loss, whereas a pathological change in the cochlea causes cochlear hearing loss. In most cases, tinnitus cannot be regarded as a direct consequence of the pathological changes, but rather hearing loss causes deprivation of input to the central auditory pathway activating neural plasticity (see Chap. 10). The tinnitus that occurs together with hearing loss is, in most cases, subjective tinnitus. Only very rarely can conductive hearing loss cause objective tinnitus, generally because of vascular turbulence. The effect of conductive hearing loss is the same as that of an earplug, and tinnitus might be interpreted as intensified perception of body sounds that occurs because sounds from the outside are reduced.

If the hearing loss is reduced or eliminated, tinnitus may also disappear after a certain period. Many forms of conductive hearing loss, in particular, can be treated successfully by surgical interventions (Chap. 83), leading to improvement of hearing and disappearance of tinnitus. If hearing loss persists, the accompanying tinnitus also usually persists. Often, the frequency of maximum hearing loss coincides with the frequency of the tinnitus.

Causes of Conductive Hearing Loss

Changes in the Territory of the External Auditory Canal

Pathologies leading to conductive hearing loss with subsequent tinnitus can be of a mechanical, inflammatory, or neoplastic nature. It may affect the ear canal or the middle ear. Genetic factors as well as exogenous noxious agents (thalidomide embryopathy [1]) may be involved in the development of auditory canal anomalies.
Obstruction of the external auditory canal by wax is a mundane reason for sudden onset of hearing loss and tinnitus. Inflammation of the external auditory canal (swimmer’s otitis) may cause very painful swelling, redness, and discharge from the external auditory canal due to bacterial (Pseudomonas aeruginosa, Staphylococcus aureus) or, more rarely, fungal infection. When the external auditory canal becomes blocked, tinnitus may develop together with the hearing loss [2]. Neoplastic disorders leading to increasing stenosis and finally occlusion of the external auditory canal may be benign or malignant. Exostoses1 of the external auditory canal are benign new bone formations that occur with an incidence of 3–6% [3]. They may cause recurrent inflammation of the external ear canal with conductive hearing loss and transient tinnitus [4]. Apart from the constitutional factors, repeated thermal irritation of the external auditory canal by frequent contact with cold water (swimmer’s ear, surfer’s ear) has long been regarded as a predisposing factor [5].

Malignant neoplasms of the external auditory canal are far less common. These may arise from skin cells (basal cell carcinoma, squamous cell carcinoma, malignant melanoma) or in the ceruminous glands (adenocarcinoma, adenoid cystic carcinoma) [6]. Congenital changes resulting in partial occlusion or atresia of the external auditory canal can cause hearing loss and possibly tinnitus. Severe forms may also be accompanied by an auricular anomaly in addition to complete atresia of the external auditory canal. As syndromic components, external ear anomalies may also occur in association with further dysmorphologies, for example, mid-facial dysplasia (mandibulofacial dysostosis, Treacher-Collins syndrome [7]), or craniofacial dysstoses (Crouzon syndrome [8]). These anomalies may be accompanied by additional anomaly of the middle ear, which can be detected by high-resolution computed tomography [9].

Pathological Conditions of the Middle Ear

Eustachian tube dysfunction leads to impairment of the function of the middle ear. Disturbances affecting opening and closure of the Eustachian tube are an important factor in the pathogenesis of many middle ear conditions because the ventilation and drainage of the middle ear no longer function properly. The sequelae may include chronic mucoid otitis media (glue ear), recurrent acute otitis media, or chronic otitis media. Disturbances affecting opening and closure of the Eustachian tube may be caused by mechanical blockage of the tubal orifice (adenoids, tumor), by inflammatory swelling of the tubal mucosa, or by muscular insufficiency such as may occur in individuals with cleft palate [10, 11]. Conductive hearing loss, sometimes accompanied by tinnitus, may develop subsequent to Eustachian tube dysfunction.

A patulous Eustachian tube is a special condition in which there is a permanently open connection between the tube and the nasopharynx. This condition may entail a variety of symptoms, such as autophony, aural fullness, and the unpleasant sensation of hearing one’s own respiratory sounds. Reduced muscle tone and weight loss are the main factors predisposing to the development of patulous Eustachian tube.

Acute otitis media often occurs secondary to rhinitis or pharyngitis. The common organisms that cause otitis media are streptococci, Haemophilus influenzae, and staphylococci. The main symptoms are earache and hearing loss. In many cases tinnitus may occur as an additional symptom. The tinnitus may be objective, having a vascular origin. It may be perceived as pulsatile pounding and buzzing sounds that occur in acute inflammatory stage by dilatation of vessels and high pulse amplitude of blood flow. Chronic otitis media is an umbrella term that covers several different middle ear pathologies. If it lasts for several years, there may be different extents of irreversible tissue destruction in the middle ear. In addition to perforation of the eardrum and defects in the ossicular chain, enzymatic degradation processes associated with cholesteatoma2 may occur and may, in particular, lead to destructive erosion of the bony walls of the middle ear toward the cranial base. The common consequence of these

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1 Exostosis: A cartilage-capped bony projection arising from any bone that develops from cartilage. Stedman’s Electronic Medical Dictionary.

2 Cholesteatoma: Squamous metaplasia or extension of squamous cell epithelium inward to line an expanding cystic cavity that may involve the middle ear or mastoid, erode surrounding bone, and become filled with a mass of keratinized squamous cell epithelial debris, usually resulting from chronic otitis media. The lesion often contains cholesterol clefts surrounded by inflammatory and foreign body giant cells, hence the name cholesteatoma. Stedman’s Electronic Medical Dictionary.
pathologies is increasing conductive hearing loss with recurrent mucous or purulent discharge. Individuals in whom cholesteatoma is associated with erosion of the bony walls of the structures of the labyrinth, may also have sensorineural hearing loss and even deafness because the pathologies have spread to the cochlea.

There are three different causes of traumatic ear-drum perforation: direct mechanical or thermal injury, a pressure wave in the external auditory canal, or an otobasis fracture. Depending on the extent of hearing loss, the possibility of a concomitant injury to the ossicular chain or the cochlea must also be considered.

Otosclerosis causes stapedia! ankylosis, which in turn causes conductive hearing loss. Otosclerosis can also affect the bony labyrinth characterized by bone resorption and remodeling processes (otospongiosis). Otosclerosis is responsible for 5–9% of all hearing losses and 18–22% of all conductive hearing losses [12]. The condition is encountered almost exclusively in Caucasians, very rarely occurring in Asians and almost never in Blacks [13]. The female-to-male ratio is approximately 2:1. The precise etiology remains unclear [14]. Alongside genetic factors [15], the role of inflammatory processes (localized measles virus infection of the otic capsule [16]), endocrine factors [17], and immunological disease [18] have been considered in the etiopathogenesis. Independently of sex and age, tinnitus is a concomitant symptom in 65–91% of individuals with otosclerosis [19, 20]. Tinnitus already develops in many individuals with otosclerosis years before the onset of noticeable hearing loss. Tinnitus sometimes persists despite an optimal hearing outcome from surgical management (see Chap. 83).

Otosclerotic processes may spread to the cochlea and that may be responsible for persisting tinnitus. This condition, known as “cochlear otosclerosis” by many authors, causes signs of sensorineural hearing loss [21–23]. It is also associated with changes in the stria vascularis, the organ of Corti and the spiral ligaments, as demonstrated in histopathological and radiological studies [24–26].

Tumors affecting the middle ear are rare. The glomus tumor (synonyms: paraganglioma, chemodectoma) is one kind of benign tumor of the middle ear, displaying destructive growth. Two locations of glomus tumors occur: glomus tympanicum tumors (which are limited to the middle ear) and glomus jugulare tumors (lesions that affect both the middle ear and the bulb of the jugular vein) [27]. In histological terms, the tumors consist of non-chromaffin paraganglionic cells along the course of cranial nerves IX and X.

The majority of glomus tumors occur in adulthood, with a female-to-male predominance of 6:1 [28]. Glomus tumors can generally be diagnosed clinically on the basis of the symptom triad of conductive hearing loss, pulsatile tinnitus, and a red middle-ear tumor that can be seen through the eardrum using otoscopy [29]. In some patients, pulsatile objective tinnitus can be detected objectively by inserting a stethoscope or microphone into the external auditory canal. The sound is probably produced by the formation of microvascular shunts within the tumor mass [30]. With increasing tumor infiltration into the jugular foramen, additional deficits related to caudal cranial nerves IX–XI may become evident.

Facial nerve schwannoma is another – very rare – benign tumor affecting the middle ear, characterized by slowly progressive facial paralysis as well as conductive hearing loss. Wegener’s granulomatosis, 3 Langerhans cell histiocytosis, 4 and sarcoidosis 5 are among the tumor-like lesions that potentially involve the middle ear and are also accompanied by tinnitus, in addition to conductive hearing loss [31].

Malignant tumors that may involve the middle ear are squamous cell carcinoma and adenoid cystic carcinoma.
Causes of Sensorineural Hearing Loss

The causal factors responsible for the development of sensorineural hearing loss can be many and varied. Apart from congenital factors, the etiology may include infectious diseases, autoimmune diseases, toxic lesions (see Chap. 42), noise-related injury (see Chap. 37), traumatic damage (see Chap. 67), or age-induced changes (presbycusis, see Chap. 36). Furthermore, fluctuating sensorineural hearing loss is one of the three signs of Ménière’s disease (see Chaps. 38 and 60) and sudden hearing loss (Chap. 56). Depending on the cause, the form of sensorineural hearing loss can be different and have varying severity. All forms of hearing loss may be accompanied by tinnitus of varying severity. The following discussion will deal with those forms of sensorineural hearing loss that are not covered in separate chapters of their own.

Sensorineural Hearing Loss of Genetic Origin

Impairment of hearing is the most common sensorineural pathology affecting humans. Approximately, one-half of all the cases of prelingual hearing impairment have a genetic cause [32–34]. A distinction is made between genetic hearing loss occurring as a component of a specific (genetic) syndrome (30%) and non-syndromic hearing loss that occurs in the absence of any other genetic diseases or developmental anomalies [35]. Syndromic hearing loss (SHL) can be inherited in an autosomal dominant, autosomal-recessive, or X-linked manner [34], and can be associated with developmental anomalies of the inner ear or petrous portion of the temporal bone like Mondini6 or Scheibe7 dysplasia; frequently there is also a link with other organic disorders, such as thyroid disease (Pendred syndrome), renal dysfunction (Alport syndrome), or eye disease (Usher syndrome) [34]. Children with Down’s syndrome are more likely to have congenital permanent inner ear hearing loss than the general population (which has an incidence of 1:1,000). From teenage years onward, they are likely to develop degenerative cochlear changes, and most will have significant hearing loss by the age of 40 years [34, 36]. Of non-syndromic hearing loss (NSHL), 80% have autosomal recessive, 18% an autosomal dominant, and 2% an X-linked or mitochondrial inheritance pattern [32]. In the majority, a single gene defect leads to the phenotypical development of hearing loss, which may not have its onset until later in life. In recent years, many gene loci and mutations have been described that are responsible for various forms of hearing loss. For example, known mutations affect the GJB2 gene – which codes for connexin-26 – and the GJB6 gene (connexin-30) [37] (see also Chap. 7).

Infections

Inflammation of the cochlea may develop together with acute or chronic otitis media when bacteria enters the cochlea through the round or oval window, or may develop together with meningitis where bacteria enters the cochlea and the vestibular apparatus via the internal auditory canal, the cochlear aqueduct, or the vestibular aqueduct. The resulting sensorineural hearing loss is often accompanied with tinnitus. Because of the involvement of the vestibular apparatus, the symptoms are often dominated by pronounced rotatory vertigo accompanied by nausea and vomiting. Meningitis is often followed by the cochlea being filled with bone (labyrinthitis ossificans or “white cochlea”), with complete obliteration of the membranous labyrinth [38]. The bacteria in borreliosis or syphilis can spread via blood to the inner ear. Many kinds of infections can cause damage to the cochlea or the inner ear as a whole. Especially serious ones are congenital rubella or cytomegalovirus infections, which may lead to severe sensorineural hearing loss or to deafness. Of the postnatal viral infections of the inner ear, epidemic parotitis (mumps) typically causes unilateral deafness without vestibular involvement [39]. Herpes zoster oticus that is caused by reinfecation with the varicella zoster virus can cause blisters in the external auditory canal and the pinna in addition to sensorineural hearing loss and

6 Mondini dysplasia: Congenital anomaly of osseous and membranous otic labyrinth characterized by aplastic cochlea and deformity of the vestibule and semicircular canals with partial or complete loss of auditory and vestibular function; may be associated with dilated vestibular aqueduct and spontaneous cerebrospinal fluid otorrhea resulting in meningitis.

7 Scheibe dysplasia: Hearing impairment due to cochleosaccular dysplasia; usually autosomal recessive inheritance. Stedman’s Online Medical Dictionary.
tinnitus, vestibular symptoms, and facial nerve palsy. When the facial nerve is involved it is known as the Ramsey Hunt syndrome [40]. In addition to infection of the nerve sheaths, the symptoms may also be caused by secretion of toxins into the perilymph spaces of the inner ear [41].

**Immunogenic Labyrinthitis**

Sensorineural hearing loss may occur together with immunological diseases as a heterogeneous group of sensorineural hearing loss types under the heading “autoimmune inner ear disease” (AIED) [42]. Possible target structures for antibodies are the stria vascularis in the organ of Corti and the blood vessels supplying the inner ear [43]. AIED is characterized by progressive, often fluctuating and usually bilateral, sensorineural hearing loss with tinnitus and vertigo, more often in women. Progression over time is too rapid to suggest presbycusis and too slow for a diagnosis of sudden deafness. AIED patients respond well to immunosuppressant corticosteroid therapy. Some patients with AIED present with a systemic autoimmune disease, such as Wegener’s granulomatosis (see footnote 3), Cogan syndrome,8 or relapsing polychondritis9 [44]. No specific test battery that will unequivocally show the presence of an autoimmune reaction to structures of the inner ear has yet been described. The recommendation is to use general laboratory tests (antineuclear antibodies, antineutrophil cytoplasmatic antibodies, etc.) to screen patients who are suspected of having an autoimmune disease for the presence of systemic signs [45].

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8 Cogan syndrome: Typical Cogan syndrome is characterized by interstitial keratitis and vestibuloluditory dysfunction. There is generally a brief episode of inflammatory eye disease (interstitial keratitis) followed by bilateral audiovestibular symptoms. The interstitial keratitis usually occurs with sudden onset and is characterized by photophobia, lacrimation, and eye pain. The vestibuloluditory dysfunction is usually bilateral, presenting with tinnitus, sensorineural hearing loss, and acute episodes of vertigo.

9 Relapsing polychondritis: a degenerative disease of cartilage producing a bizarre form of arthritis, with collapse of the ears, the cartilaginous portion of the nose, and the tracheobronchial tree; death may occur from chronic infection or suffocation because of loss of stability in the tracheobronchial tree; of autosomal origin. Stedman’s Online Medical Dictionary.

**Age-Related Hearing Loss**

Age-related hearing loss is the commonest of all forms of hearing loss (see Chap. 36). It affects more than 40% of people over the age of 65 [46]. Apart from physiological age-related processes, endogenous and exogenous factors such as hypoxia, exposure to loud noise, hypertension, hypercholesterolemia, or diabetes mellitus may cause or contribute to hearing loss in old age [47]. Consequently, excessive noise exposure and atherosclerosis contribute to the development of presbycusis in industrialized countries. The reported presence of tinnitus together with presbycusis varies between 8 and 72% [48–50]. The risk for the development of tinnitus rises with increasing age and with increasing exposure to noise [51].

**References**

Keypoints

1. Damage in the external, middle, or internal ear can contribute to the emergence of tinnitus because of the hearing loss it causes.

2. The two components of the external ear are the auricle and the outer auditory canal.
   (a) The occlusion of the ear canal produces an alteration in sound transmission that may cause tinnitus to develop.
   (b) Ear canal inflammation may cause tinnitus.

3. The middle ear is an impedance transformer and the site of several pathologies that all may cause tinnitus.
   (a) Acute otitis is accompanied by fever, strong pain in the ear, conductive hearing loss, and discharge from the ear.
   (b) Otitis media with effusion is a chronic presence of seromucous secretions in the middle ear cavity without signs of acute inflammation.
   (c) Otitis media is an inflammation of the middle ear causing conductive hearing loss.
   (d) Cholesteatoma is a mass of keratinizing squamous cells or epithelial debris that may occur in the middle ear cavity; it can erode body structures.
   (e) Otosclerosis involves a bony formation around the stapes, impeding its motion.

4. The first symptom of otosclerosis is often tinnitus.

5. Tinnitus often occurs in association with hearing loss of cochlear origin.
   (a) Acoustic trauma is one of the most common risk factors for the development of tinnitus and one of the major causes of permanent sensorineural hearing loss.
   (b) Administration of ototoxic drugs can cause hearing loss, tinnitus, and vertigo or dizziness.
   (c) Age-related changes can cause tinnitus and hearing loss.
   (d) Tinnitus is one of the three symptoms that define Ménière’s disease.
   (e) Changes (decrease) in cochlear blood perfusion can lead to cochlear damage with hearing loss and tinnitus.
   (f) Abrupt change in barometric pressure (barotraumas) can cause damage to the cochlea and may lead to tinnitus.

6. Hearing loss due to ear diseases may trigger a series of reactions in the central nervous system, which leads to the tinnitus.

7. Head trauma can lead to tinnitus, and balance disorders are very common after mild to severe head traumas.

Keywords Tinnitus • Hearing loss • External ear • Middle ear • Internal ear • Cochlea

Abbreviations

CF Characteristic frequency
CNS Central nervous system
dB Decibel
Hz Hertz
IHC Inner hair cell
kHz Kilo hertz
OHC Outer hair cell
SGN Spiral ganglion neuron
Introduction

The peripheral auditory system includes the external, the middle, and the internal ear, as well as the acoustic nerve. Damage in one of these structures can contribute to the emergence of tinnitus, associated with hearing loss (Table 35.1). Different mechanisms of the pathophysiology of tinnitus have been hypothesized from the study of the relation between tinnitus and hearing loss, as summarized in the present chapter (see also Chaps. 8 and 10).

Hearing Loss and Tinnitus Caused by External Ear Damage

The two components of the external ear are the auricle and the outer auditory canal, which conveys acoustical waves to the eardrum. The function of the auricle is to direct sounds into the entrance of the ear canal and it acts as an acoustic filter. The auricle is important for directional hearing in the vertical plane and about the direction to the sound source, either frontal or back. Comparison of the input from both ears helps to localize sounds in space.

The external ear canal is 24–27-mm long from the entrance to the ear drum. The external auditory canal amplifies frequencies between 2,000 and 4,000 Hz when the sound source is located in front of an observer [1].

Table 35.1 Main pathologies in the ear which may cause tinnitus

<table>
<thead>
<tr>
<th>Subjective tinnitus</th>
<th>Causes</th>
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<td>Pathologies: middle ear</td>
<td>Acute otitis media</td>
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<td>Otitis media with effusion</td>
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<td>Suppurative otitis media</td>
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<td>Cholesteatoma</td>
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<td>Otosclerosis</td>
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<td>Pathologies: internal ear</td>
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<td>Noise induced hearing loss</td>
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<td>Presbycusis</td>
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<td>Ménieré’s disease</td>
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<td>Alteration in blood flow</td>
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<td>Barotrauma</td>
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<td>Head trauma</td>
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Occlusion of the Ear Canal

Occlusion of the ear canal causes hearing loss on the affected ear. If an obstruction occurs in only one ear, it causes perception of annoying echoes and distortion that may develop into tinnitus. The most common cause of occlusion in the ear canal is the presence of ear wax (cerumen). Ear wax is produced by particular sweat glands located in the ear canal wall. Its function is to protect the ear skin from infections and the ear canal from the entry of foreign bodies. Normally, it moves out of the ear canal automatically. Accumulation of wax in the ear may lead to a wax plug, which is caused by an increased secretion of ear wax, change in the composition of the wax, or anatomical changes in the ear canal (e.g., stenosis, osteoma, esostosi) that prevent the normal movement of wax thus causing buildup. Common symptoms due to the presence of a wax plug are hearing loss, aural fullness, and tinnitus, which typically worsen when water penetrates into the ear. (Ear wax is not completely solvable in water and has a hygroscopic compound.) The presence of a foreign body in the ear canal is another cause of occlusion. These include cotton, gauzes, or even living organisms such as insects, which can be very annoying. The appropriate therapy is the removal of the foreign bodies.

Cancers of the ear canal skin are rare; occlusion-induced symptoms include tinnitus. After a proper assessment, therapy may either be surgical or pharmacological.

External Otitis

Ear canal inflammation is called external otitis. It is always accompanied by itching or pain due to the rich innervation of the skin in the ear canal, and sometimes by fever. Additional symptoms may be discharge, hearing loss, and tinnitus. Signs may include redness and edema with the possible presence of secretions [2].

Ear canal infections can be caused by bacteria or fungi. Anatomical changes in the ear canal can increase the likelihood of infections; frequent and aggressive bathing and bath detergents are such factors. Others include changes of pH in the ear canal. Pharmacological treatment is very important to prevent the spread of infections to neighboring areas. Tinnitus usually disappears with the recovery from inflammation.
Compromised sound transmission to the internal ear by an obstruction of the ear canal causes sound deprivation, which enhances the existing tinnitus and may cause tinnitus in individuals who did not previously have the disorder. This is because deprivation of sound activates neural plasticity (see Chap. 12). Most individuals who are placed in a silent environment will experience tinnitus [3–5].

**Hearing Loss and Tinnitus Caused by Damage to the Middle Ear**

The middle ear consists of the eardrum and the ossicular chain (malleus, incus, and stapes), which transmits the sounds that reach the eardrum to the cochlear fluids through movements of the stapes footplate located in the oval window. The main function of the middle ear is to optimize energy transfer from the air to the cochlear fluids. It does so by acting as an impedance transformer [6]. Direct transfer of sound energy to internal ear fluids is inefficient because of large differences in the impedance of air and the fluid of the cochlea [7].

The ossicular chain is kept in place by four ligaments and two muscles, the tensor tympani and stapedius muscle. The tensor tympani muscle, with more tonic fibers, is innervated by a branch of the mandibular nerve (a branch of the trigeminal nerve). The stapedius muscle, with more phasic fibers, is innervated by the facial nerve. In humans, contraction of the stapedius muscle can be elicited by a strong sound, whereas the tensor tympani muscle contracts when swallowing and yawning. The acoustic stapedius reflex extends the dynamic range of human hearing, improving word intelligibility of loud sounds [8] and improves discrimination in noise by alternating low frequency sounds that can mask high frequency sounds.

Myoclonus (repetitive abnormal contractions) of the muscles may cause objective tinnitus [9], which may either be perceived as a rhythmic clicking or buzzing [10]. A single instance of continuous and high-frequency tinnitus has been described [11].

The middle ear is connected to the rhinopharynx through the Eustachian Tube. The tube is normally closed, except during swallowing and intense efforts when opened by the peristaltic muscles to allow air to enter the tympanic cavity, thus permitting replacement of air and normalization of pressure to ambient pressure.

**Acute Otitis Media**

Inflammatory diseases of the middle ear can cause tinnitus; damages to the epithelium in the tympanum and Eustachian tube dysfunction prevent the physiological absorption of air and promote stagnation of secretions, causing alterations in the function of the middle ear. Acute otitis is mainly caused by bacterial infections originating in the nasal cavities. It is accompanied by fever, strong pain in the ear, and conductive hearing loss [12]. It may progress into perforation of the ear drum and discharge from the middle ear.

**Otitis Media with Effusion**

Otitis media with effusion is characterized by a chronic presence of seromucous secretions in the middle ear cavity without signs of acute infection but with the presence of conductive hearing loss, aural fullness, and sometimes tinnitus. The associated dysfunction of the Eustachian tube prevents normal ventilation of the middle ear cavity causing negative pressure in the middle ear with secretion of fluids from the mucosa [13]. Treatment is either pharmacological or surgical.

**Chronic Otitis Media**

Chronic otitis is the consequence of protracted inflammation of the middle ear with alterations of the eardrums, usually perforation. The ossicular chain may be affected, calcified, or destroyed. Conductive hearing loss and tinnitus are generally also present [14].

In all these disorders, the normal functions of the middle ear has deteriorated and sound transmission to the cochlea is compromised causing conductive hearing loss often accompanied by aural fullness and tinnitus, which is described by patients as a broadband noise in most cases.

**Cholesteatoma**

Cholesteatoma is a mass of squamous cells or epithelial debris, a keratinizing lesion that occurs in the middle ear. It may be either congenital or acquired [15].
Cholesteatoma may grow in a way that involves the entire middle ear cavity by eroding the mastoid bone. Clinical presentation is similar to that of chronic otitis, and the common treatment is surgical and involves removal of the pathologic tissue.

**Otosclerosis**

Otosclerosis is a disease of the cochlear capsule characterized by the formation of soft, vascular bone around the stapes. This causes fixation of the stapes footplate in the oval window [16], resulting in conductive hearing loss and often tinnitus. A genetic factor plays a role in the occurrence of otosclerosis [17, 18], as evident from the fact that it is more common in some families and occurs more often in women. Hearing loss is initially conductive and is greatest for low frequencies, but it increases with progression of the disease. After some time, the disease progresses to the cochlea where it causes ossification causing sensorineural hearing loss and involving high frequencies. Often the first symptom of otosclerosis is tinnitus, which is typically described as ringing, whistling, or roaring. The common treatment of otosclerosis consists of replacement of the stapes by a prosthesis inserted in the footplate, which can restore hearing to near normal and often results in resolution of the tinnitus [19]. A hearing aid is an alternative option, which also can reduce the tinnitus because it restores the ability to hear environmental sounds (see Chap. 74).

**Hearing Loss and Tinnitus Caused by Cochlear Damage**

The cochlea is a coiled tube containing the sensory cells of audition. The cochlea separates sounds according to frequency before they are transduced by the sensory cells into a neural code in auditory nerve (see Chaps. 8 and 10).

Tinnitus often occurs in association with hearing loss and a common pathophysiology has been hypothesized. Both tinnitus and hearing loss are often associated with other conditions, such as noise trauma, ototoxic drugs, head trauma, Ménière’s disease, cochlear hydrops, presbycusis, and genetics alterations and syndromes. Several studies have clearly described how these conditions are associated with hearing impairment, but the exact mechanisms generating tinnitus are still under investigation. Several proposed potential mechanisms are complex and controversial.

In the past years, hearing impairments have been regarded as the effect of injuries to hair cells, mainly OHC. However, it has become evident that disorders of the cochlea can indirectly influence the function of the CNS. Symptoms such as tinnitus and hearing loss are likely to have components that originate in the CNS, causing deprivation and changed balance between inhibition and excitation [1, 20]. Some symptoms may also promote an expression of neural plasticity, which in turn can cause symptoms such as tinnitus, hyperacusis, changed dynamic range, or redirection of information (see Chap. 12).

This new approach to understand the pathophysiology of hearing impairment has blurred the distinction between cochlear and nervous system disorders. Sensorineural hearing loss related to cochlear damage is often characterized by alterations in speech perception and recruitment of loudness; in the past, they were regarded to be related only to damage of hair cells, whereas recent studies have shown that they may also be caused by a change in the function of the central nervous auditory system. In conclusion, hearing impairments may both be the result of a combination of deficits in the auditory periphery and the effect of changes in the CNS. In recent years, evidence has accumulated that plastic changes in the central auditory nervous system can cause tinnitus.

Any factor able to alter the level of the spontaneous activity in the auditory nerve may theoretically result in tinnitus [21]. Bauer [22] proposed that tinnitus is the result of reversible OHC dysfunction and decoupling stereocilia from the tectorial membrane. Schwaber [23] theorized that the basic pathophysiology of tinnitus involves an alteration in stereocilia stiffness that results in an increase in the discharge rate of hair cells. Möller [24] analyzed the link between tinnitus and hearing loss and concluded that tinnitus is not directly related to the degree of the hearing loss. Tinnitus may be the result of a combination of causes.

Factors involved in damage of the cochlea, most commonly related to tinnitus and hearing loss, are discussed in separate chapters (Chaps. 36, 37 and 42).
Ménière’s Disease

Ménière’s disease [25] is a progressive disorder defined by periodic attacks of vertigo, fluctuating sensorineural hearing loss, tinnitus, and aural fullness [26] (see Chaps. 38 and 60). Several authors have agreed that Ménière’s disease is not a single disorder, but rather a family of different disorders. It has been customary to include patients who do not show all symptoms in the term Ménière’s disease. For instance, patients who only have tinnitus with aural fullness and a mild temporary hearing loss limited to low frequencies may also be classified as having Ménière’s disease.

In the early stages of the disease, hearing loss only affects low frequencies and is almost completely reversible, but as the disease progresses residual hearing worsens definitively and spreads to higher frequencies. Symptoms are first unilateral, but in many patients they will, in the course of the disease, involve the contralateral ear after 10 or 15 years. Tinnitus may persist between acute attacks, but typically increases in 1 or 2 days before vertigo (Regarding contemporary hypothesis about Ménière’s Disease, see Chaps. 38 and 60.).

Alterations in Blood Flow in the Cochlea

The cochlea needs a correct blood supply to preserve its function [27]. The labyrinthine artery is an endartery, and the inner ear has no collateral blood supply. Even minimal alterations in blood perfusion can lead to cochlear damage. Additionally, vulnerability of the cochlea causes high-energy consumption from the blood supply required to keep the cochlear fluids in perfect balance.

It has been hypothesized that the entity and the frequencies involved in hearing loss may depend on the location of the vascular accident; the district of the cochlea permeates the thrombosis downstream or the bleedings will be affected by damage.

Many systemic diseases that cause hearing loss are also accompanied by tinnitus such as those that are caused by circulatory and microcirculatory alterations [28, 29]. Many hypotheses have been proposed to explain the pathology of sudden hearing loss that occurs without any additional symptoms. One hypothesis regards vascular impairment [30]. It has been suggested that viral infections can also cause sudden hearing loss [31] through a viral attack of endothelial cells, even though no conclusive evidence has been yet found. Hearing loss may be profound at its onset and often accompanied by tinnitus. Hearing may deteriorate within a few hours; fortunately, it is almost always limited to one ear and the chance of spontaneous recovery is good.

Hearing function is generally recovered spontaneously in one-third of patients; it is improved in one-third and is maintained with no change in the remainder of patients. In some cases hearing will improve, while tinnitus will remain the same. Different therapeutic approaches have been used such as administration of steroids, antiviral agents, or hyperbaric oxygen treatment [32, 33].

Barotrauma

Tinnitus can present after diving in water or during a flight, especially in conjunction with cold or allergy symptoms. In these cases, a barotrauma may have caused the symptoms because abrupt change in barometric pressure has affected the cochlear fluids. Tinnitus may either be temporary or persistent. If the trauma has been so severe that it caused a rupture of the eardrum or the round windows (fistula), resulting in hearing loss and/or vertigo, tinnitus often occurs. Adequate medical or surgical treatment is necessary to restore hearing and relieve the tinnitus.

Head Trauma

Tinnitus and balance symptoms are common after both mild and severe head trauma. A temporal bone fracture that may occur after head injury can lead to cochlear damage with severe hearing loss, vertigo, and tinnitus [34]. Acute vertigo typically lasts for several days and then resolves, but dizziness may persist. Hearing loss is often permanent. Tinnitus may decrease or persist together with hearing loss.

Head trauma without a temporal bone fracture can cause cochlear damage [35] with symptoms such as dizziness, tinnitus, and sometimes hearing loss. These symptoms may be temporary or occur sometime after the injury. The symptoms may last for several months
and then gradually abate, while tinnitus often remains. The mechanisms that lead to hearing disorders by concussion are not fully known; it has been suggested that the higher nervous structures are involved.

Head trauma can also cause blood effusion into the tympanic cavity, as well as rupture of the eardrum and dislocation of the ossicular chains, resulting in conductive hearing loss and tinnitus. These injuries often require adequate surgical treatment.

**Conclusion**

Many studies have focused on the relationship between tinnitus and other symptoms, such as hearing loss. However, no clear explanation has been published regarding the cause of tinnitus. Many authors agree that the initial cause of tinnitus, when occurring together with hearing loss, has a peripheral origin that may trigger a series of reactions in the CNS, resulting in tinnitus. The anatomical location of the pathology may be the cochlea, but it is often the auditory nervous system (see Chap. 10). The finding that both people with hearing loss and individuals with normal hearing may have severe tinnitus clearly indicates that tinnitus is not always linked to hearing loss.

**References**


Chapter 36
Cochlear and Non-cochlear Age-Related Hearing Loss and Tinnitus

Aage R. Møller

Keypoints
1. Age-related changes are some of the most common causes of disorders of sensory systems.
2. The most common age-related change in hearing is elevation of the hearing threshold beginning at the highest audible frequencies, progressing toward lower frequencies while deepening.
3. Age-related changes in hearing are often, but not always, accompanied by tinnitus.
4. Age-related changes in hearing function may be caused by:
   (a) Degeneration of sensory receptor cells, in the cochlea
   (b) Change in the conduction velocity of sensory nerve fibers
   (c) Change in the access to neural transmitters, such as gamma amino butyric acid (GABA), and subsequent increases in GABA receptor sites
5. Change in processing of information may also occur, causing deterioration of speech comprehension.
6. Animal studies have shown that the progression of age-related changes in hearing might be affected (slowed down) by exposure to sound (“enhanced sound environment”) indicating expression of neural plasticity plays a role in some age-related changes of sensory functions.
7. The large individual variability in age-related changes in hearing has many causes, such as exposure to loud sounds, environmental factors, genetics, different expression of genes (epigenetics), and unknown factors.

Keywords Presbycusis • Age-related hearing loss • Tinnitus • Neural plasticity

Abbreviations
ARHI Age-related hearing impairment
EPSP Excitatory postsynaptic potentials
GABA Gamma amino butyric acid

Introduction
Age-related impairment of hearing (presbycusis) is the most common disorder of the auditory system. The most obvious changes in hearing that occur with increasing age are an elevated hearing threshold for high frequencies. Presbycusis normally refers to the elevation of hearing threshold. In addition, the elevation of hearing threshold and impaired processing of sound, known as phonemic regression may occur. Many individuals acquire tinnitus in old age, and it often accompanies presbycusis. However, it may also occur together with minimal hearing loss. Most elderly people have tinnitus when placed in a silent room, such as a room used for audiologic tests.

1Presbycusis: Loss of hearing associated with aging; manifest as reduced ability to perceive or discriminate sounds; the pattern and age of onset vary (Stedman’s Electronic Medical Dictionary).
2 Phonemic regression: a decrease in intelligibility of speech out of proportion to the pure tone hearing loss associated with aging (Stedman’s Electronic Medical Dictionary).
Epidemiology of Presbycusis

Normally, hearing loss increases gradually with age, as shown in many studies. Spoor et al. [1] have reviewed the literature and presented average audiograms for different age groups from eight different population studies based on a total of 7,617 ears – including both men and women (Fig. 36.1). This classical study of age-related hearing loss included the effect of environmental factors, such as noise exposure, and did not show the individual variations.

There is a distinct difference between hearing loss in men and women (Fig. 36.1), but that may be at least partly a result of different degrees of noise-induced hearing loss. It has been preferentially men who were working in industries with heavy noise exposure. This effect of noise exposure is particularly prominent with participants in the older studies, such as those summarized by Spoor with the hearing loss depicted in Fig. 36.1. Some of the individual variations in presbycusis may thus be attributed to environmental factors, mainly the varying degree of exposure to sounds.

Large individual variations were mentioned in several studies. One study [2] quantified these variations (Fig. 36.2). This study showed the individual variations in hearing loss and in speech discrimination. Also, this study included individuals who had exposure to noise that caused hearing loss, affecting mostly men.

It seems likely that genetic factors also play a role. In fact, a gene that affects age-related hearing loss has been identified in a mice strain [3]. There are many genetic disorders that affect hearing in general [4], but not specifically regarding the deterioration of hearing with age. A study that specifically addressed genetic predisposition for age-related hearing loss [5] found that approximately half of the variance of Age-Related Hearing Impairment (ARHI) is attributable to environmental risk factors. The other half is linked to genetic factors.

Gates and co-authors [6] described the results of a large population study (Farmingham). Hearing sensitivity and word recognition tests in 1662 men and women between the age of 60 and 90 showed that the pure-tone thresholds increased with age at a rate that did not differ by gender. However, men had poorer hearing threshold in general. This means the result of noise exposure had its full effect on hearing thresholds before a person reaches the age of 60, which is the age at which this population study began. Maximum word recognition ability declined with age more rapidly in men than in women and was also poorer in men than women at younger ages.

One more recent study [7] found that the hearing threshold increased approximately 1 dB per year in individuals of 60 years and above. Females of 70 years and above had a faster rate of change in hearing threshold at 0.25 to 3, 10, and 11 kHz than females in the age group of 60–69 years.

Other authors [8] found a true gender difference in hearing threshold, including a difference in older individuals where women have less age-related hearing loss. Jerger et al. also referred to the hypothesis about

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**Fig. 36.1** (a) Average hearing loss in different age groups of men. Combined results from eight different published studies based on a total of 7,617 ears. (b) Average hearing loss in different age groups of women. Results from eight different published studies based on 5,990 ears. Reprinted from Møller AR (2006) Neural plasticity and disorders of the nervous system [1] with permission.
cardiovascular diseases promoting hearing loss [8], or perhaps, the same genetic factors that promote development of cardiovascular disorders also promote hearing loss. For example, animals studies of rats with predisposition for hypertension also acquire more age related hearing loss and more hearing loss from noise exposure [9, 10]. Other animal experiments have shown the progression of the age-related elevation of the hearing threshold can be arrested by appropriate sound stimulation [11].

**Epidemiology of Age-Related Tinnitus**

Tinnitus almost always occurs together with hearing loss (see Chaps. 35 and 37). Tinnitus is one of the three symptoms that define Ménière’s disease (see Chaps. 38 and 60). Tinnitus is also often associated with presbycusis, but different studies of the prevalence of tinnitus have arrived at different results. The reported concomitant presence of tinnitus varies between 8 and 72% [12–14]. The risk for the development of tinnitus rises with increasing age and with increasing noise exposure [15]. In the age group of 55–65 years, one study found that tinnitus occurred in 19.3% or 11.8%, depending on the questions asked in such studies [16]. Other studies have found very varying incidences of tinnitus together with age-related hearing loss [13, 14], but it is generally agreed that the incidence of tinnitus increases with age [15].

Tinnitus is related not only to the size of the hearing loss but also to the shape of the audiogram, being more common in individuals with a high-frequency, steeply sloping audiogram than in individuals with a flat audiogram [16].

Tinnitus cannot be measured in a similar way as in the case of hearing loss. The evaluation depends on the individual’s own assessments of the severity of his or her tinnitus. This adds uncertainties to epidemiologic studies of the prevalence of tinnitus and is the main cause of the differences reported by different authors.

Most people above the age of 60 have experienced some form of tinnitus, but these epidemiologic studies have only included individuals with tinnitus of a certain severity. Some of the causes of variations between the studies of individual investigators are diverse definitions of the different degrees of tinnitus, such as “bothersome tinnitus.” Most epidemiologic studies are performed using written questionnaires. The outcome of such epidemiological studies is affected not only by the definitions used for the level of severity, but also in the way the questions are phrased about the participant’s perception of his or her tinnitus.

The use of common medications that are associated with tinnitus such as certain diuretics increases with age [17] and this may count for some of the observed age-related increase in the incidence of tinnitus.

As is the case for presbycusis, environmental factors such as noise exposure, exposure to chemicals, and other environmental factors, thus similar reasons for causing more hearing loss in men than women, influence the occurrence of tinnitus. This was confirmed by the findings that tinnitus is more common in males than in females [16].
Pathology of Presbycusis

Many studies have shown that hair cells, especially outer hair cells, are injured in individuals with presbycusis [18] and that these injuries correspond to the hearing loss, as it is reflected in a person’s audiogram.

The hearing loss, as it is described by the pure tone audiogram, has been attributed to impairment or loss of cochlear hair cells – mostly affecting outer hair cells. Outer hair cells function to amplify the basilar membrane vibration (act as motors), but the outer hair cells probably do not participate in the signal transduction; that is done by inner hair cells [19] (see Chap. 8). The fact that the morphological changes in the cochlea are so apparent has made investigators and clinicians focus on this aspect of aging in hearing. More recent studies have shown evidence that hair cell damage is not the only cause of presbycusis.

Although morphologic changes in the cochlea of individuals with presbycusis are convincing, it is not the only reason for presbycusis. Other changes in the auditory system that occur normally with age also contribute to the loss of hearing. The abundant efferent innervation of especially outer hair cells makes it possible for the function of outer hair cells to be modulated by signals from the central nervous system (see Chap. 8). Plastic changes that affect the auditory nervous system may thereby affect the transduction process in the cochlea by changing the amplification in the cochlear amplifier. This means that the pathology causing hearing to deteriorate with age is located not only in the cochlea but also in the CNS.

The nervous system is involved in noise-induced hearing loss, as confirmed by the finding that noise-induced hearing loss can be reduced by pre-exposure to moderately strong sounds [20, 21].

It is difficult to distinguish between the deterioration of the hearing from noise exposure from that caused by age-related factors, although the shape of the audiogram of age-related hearing is different from the common noise-induced hearing loss (Chap. 37). The age-related hearing loss is greatest at high frequencies, whereas noise-induced hearing loss is, as a rule, greatest around 4 kHz [19].

The complexity of presbycusis is supported by the results of animal studies where different kinds of rats’ hearing loss during their lifetime were studied while the rats were housed under different conditions; with and without noise exposure.

In this study, three groups of rats were exposed to 85 dB, 105 dB, and no noise for 8 h every day during their lifetime [9]. Each group of rats consisted of normotensive and spontaneous hypertensive\(^3\) rats. The hearing loss from noise exposure in the 85 dB group was minimal when compared with those that, were not exposed to noise. The animals in the group that were exposed to 105 dB noise acquired considerable hearing loss. However, it was different for normotensive rats compared with spontaneous hypertensive rats, acquiring an average hearing loss of 30 dB and 60 dB, respectively [22]. The larger hearing loss from exposure to noise in the spontaneous hypertensive rats did not seem to be caused by the elevated blood pressure as such because hypertension induced by ligation of a kidney artery in normotensive rats did not cause larger noise-induced hearing loss [23]. Ligation of one kidney artery caused similar elevation of blood pressure as observed with age in the spontaneous hypertensive rats.

The results of these studies point to a genetic cause of the larger age-related hearing loss in spontaneous hypertensive rats when compared with that of normotensive rats. The genetic cause of hypertension also resulted in greater noise-induced hearing loss in spontaneously hypertensive rats rather than the effect of the high blood pressure as such. These animal studies have thus supported the hypothesis that genetic predisposition for hypertension also promotes hearing loss. The genetic abnormality of spontaneous hypertensive rats also predisposed the rats for acquiring larger than normal elevation of hearing threshold with age. That one genetic abnormality or risk factor can predispose, for more than one pathologic sign is not uncommon.

It has been hypothesized that female reproductive hormones may be involved in causing hearing loss [24]. Estrogen affects auditory neural function, as evidenced from its effect on auditory brainstem responses (ABR) [25]. It is known that female reproductive hormones can modulate the function of GABA receptors [26]. That may be the basis for the effect of female reproductive hormones on hearing loss.

The increased release of the afferent transmitter glutamate can exert a direct as well as an indirect neurotoxic effect at higher concentrations [27]. The age-related reduction in dopamine receptors may also be involved in the effect of age on the incidence of tinnitus [28].

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\(^3\)Spontaneous hypertensive rats: Rats with genetic predisposition for greater increase in blood pressure with age than normal.
Loss of the inhibitory neural transmitter, GABA, that occurs with age may promote presbycusis [29] and perhaps, in particular, tinnitus. The change in female reproductive hormones with age may therefore affect the development of presbycusis, and this effect may cause some of the differences between the development of presbycusis in men and women. The pathology of presbycusis is far more complex than just damaged hair cells.

It was mentioned above that the central nervous system could influence the function of the hair cells in the cochlea. Injuries to cochlear hair cells can also influence the function of the auditory nervous system. The auditory nervous system can influence how normal hair cells (especially outer hair cells) are damaged or get an abnormal injured function.

While injuries to cochlear hair cells can themselves cause symptoms, pathologies of hair cells can also promote expressions of neural plasticity, which can cause symptoms of hyperactivity, redirection of information, etc. (see Chap. 12). This may explain why injuries to cochlear hair cells are not the sole reason for the symptoms of age-related changes. Hearing loss that occurs when hair cells are injured is therefore not only caused by these injuries as such, but the function of the central auditory nervous system pathways may also be altered. This contributes to hearing loss caused by cochlear pathologies.

In a similar way, the fact that tinnitus is often associated with injuries to hair cells does not mean that it is the hair cells that generate the abnormal neural activity that causes tinnitus. The anatomical location of the abnormal function that causes these symptoms is thus not only the cochlear hair cells, but changes in the function of specific structures of the auditory nervous system may also contribute to some forms of tinnitus.

Presbycusis and age-related tinnitus are caused by a complex combination of deficits in the cochlea and changes in the central auditory nervous system [30, 31]. Advances in our knowledge about the disorders of the auditory system have now blurred the distinction between cochlear and nervous system disorders.

Problems to understand speech, even after that the loss in hearing sensitivity has been compensated for by amplification is common in elderly individuals. Age-related changes in the auditory nerve, where the variation in diameter of auditory nerve fibers increases with age [32] (Fig. 36.3), might contribute to hearing problems. Greater variation in the diameter of auditory nerve fibers in turn causes the conduction velocity to vary. Subsequently, the arrival time of neural activity at the cochlear nucleus will vary with the degree and kind of injury. This result in a temporal dispersion can have different effects on activation of cochlear nucleus cells [33] (Fig. 36.4). It is evident from Fig. 36.4 that increased temporal dispersion can cause both decreased excitation of target neurons or increased excitation. The latter may be a cause of some forms of tinnitus. Processing in the other nuclei and the cerebral cortex of the auditory system may also change as a result of age-related changes, contributing to difficulties in understanding speech.

**Epidemiology of Age-Related Tinnitus**

As has been pointed out in other parts of this book, data on epidemiology of tinnitus in general are sparse, and epidemiologic data on age-related are few. A study of the prevalence of tinnitus in children and the elderly [34] found the incidence of tinnitus in presbycusis to be 11%. A study in Sweden of 153 individuals from age 70 to 79...
showed that the incidence of tinnitus increased from 31% at age 70, to 44% at age 79. A few participants (11) in this study had less tinnitus at age 79 compared with what they had at age 70, thus some form of remission [35, 36].

**Relationship Between Hearing Loss and Tinnitus**

While there are individuals with tinnitus who have normal or near normal hearing, most forms of tinnitus are associated with hearing loss. A study has shown a clear relationship between hearing loss at 4 kHz and the odds of having tinnitus [37] (Fig. 36.5).

It should be noted that 4 kHz is the frequency of greatest hearing loss from noise exposure, and it can be assumed that a noticeable portion of the hearing loss of many of the participants in this study comes from noise exposure (see Chap. 37).

There are also individuals with considerable hearing loss who do not have tinnitus. Hearing loss may therefore not be regarded to always cause tinnitus, although hearing loss – including conductive hearing loss – may be associated with tinnitus, because deprivation of sound activates neural plasticity (see Chaps. 11 and 12).

Individuals with low-frequency tinnitus tend to have more severe hearing loss than people with high-frequency tinnitus [38] (see Fig. 36.6). Tinnitus in connection with age-related hearing loss can have several causes. It can be caused by activation of neural plasticity because of reduced input to the nervous system from the ear (deprivation of sensory input is a strong promoter of plastic changes) (see Chap. 12). It can be caused by aging factors other than those that cause hearing loss. The reduced GABA activity that occurs with age [39] reduces inhibition in general in the nervous system and that may promote hyperactivity, which can cause tinnitus.

It has become evident that risk factors for age-related deterioration of CNS functions causing disorders, such as different forms of dementia, overlap with risk factors for cardiovascular diseases [40]. Little is known about the relation between dementia and hearing loss or about the risk factors for presbycusis and various forms of dementia. However, it has been found that many of the changes that occur with age can be slowed down or prevented by appropriate exposure to sound [11].
Conclusion

The cause of tinnitus is complex, as discussed in several chapters in this book. Although the likelihood of having tinnitus increases with age, as does hearing loss, the casual relationship between hearing loss and tinnitus is complex and many other factors than hearing loss are involved in causing age-related tinnitus.

References

Chapter 37
Noise-Induced Hearing Loss: Implication for Tinnitus

Donald Henderson, Eric C. Bielefeld, Edward Lobarinas, and Chiemi Tanaka

Keypoints

1. Noise-induced hearing loss (NIHL) is often associated with tinnitus.
2. The shape and depth of the audiogram in patients with NIHL varies considerably.
3. Characteristics of tinnitus (sensation level, pulsatile versus continuous, perceived pitch) also vary widely across individuals.
4. The relationship between the pattern of hearing loss and the characteristics of the tinnitus is complex and a relevant topic of research.
5. This chapter focuses on three topics relevant to NIHL and tinnitus:
   
   (a) The relationship between the parameters of a noise exposure and the resulting hearing loss.
   
   (b) The cochlear pathologies underlying permanent hearing loss and temporary hearing loss and how they differ.
   
   (c) Noise-induced tinnitus and the animal modeling of tinnitus used to study the relationship between noise and tinnitus.

Keywords Temporary threshold shift • Permanent threshold shift • Kurtosis • Noise interactions • Tinnitus

Abbreviations

ATS Asymptotic threshold shift
CNS Central nervous system

EAM External auditory meatus
GPIAS Gap-prepulse inhibition of the acoustic startle
IHC Inner hair cell
NBN Narrow band noise
NIHL Noise-induced hearing loss
NIOSH National Institute for Occupational Safety and Health
OHC Outer hair cell
OSHA Occupational Safety and Health Administration
PTS Permanent threshold shift
TTS Temporary threshold shift

Introduction

Hearing loss from exposure to noise can either be temporary or permanent, depending on the level or duration of the exposure. The audiological symptoms associated with both noise-induced temporary threshold shift (TTS) and permanent threshold shift (PTS) include an elevation in hearing thresholds with particular vulnerability in the 3–6 kHz region; decreased frequency resolution and increased vulnerability to masking; abnormal growth of loudness; compromised temporal processing (i.e. decreased temporal summation of acoustic power and increased forward masking); and, of course, tinnitus (see review by Henderson et al. [1]).

There have been scores of studies on the relationship between noise exposure, the resultant hearing loss, changes in cochlear tuning, and the pathological basis for the corresponding audiomeric symptoms (see Review articles by Saunders et al., Lieberman, Henderson and Hamernik [2–4]). However, our understanding of the biological basis of tinnitus is not as well understood (see review by McFadden and Wightman [5]).
Tinnitus is a particularly interesting problem because noise exposures primarily damage the auditory periphery (cochlea) while evidence of tinnitus is often clearly central in origin. A fundamental question is what the changes in the operation of the cochlea that leads to a phantom perception generated in the central nervous system (CNS).

**Acoustic parameters of Noise-Induced Hearing Loss (NIHL)**

A review of the relationship between the parameters of noise exposure and temporary or permanent hearing loss is a reasonable place to begin an examination of the relation between noise exposure and tinnitus.

**Temporary Threshold Shift (TTS)**

Exposure to loud sound can lead to acute TTS, or if the noise is loud enough or long enough the hearing loss can be PTS. The most comprehensive study of TTS was done in the 1940s and 1950s by Hallowell Davis [6] and his distinguished colleagues. They systematically studied the relationship between the acoustic variables of *frequency*, *intensity*, and *duration* and the perceptual correlates of loudness changes, pitch coding, and tinnitus.

A summary of their findings is schematically illustrated in Fig. 37.1 and includes the following results: (1) Exposure to pure tones or noise above 90 dB SPL can shift an individual’s hearing threshold; (2) The magnitude of the hearing loss caused by a specific tone depends on the frequency of the tone, i.e. high frequencies such as 2,000 and 4,000 Hz caused a larger threshold shift than low frequencies (500 Hz). Note that the 500 Hz tone caused a broad hearing loss that was roughly equal in magnitude to the 2,000 and 4,000 Hz tones, but that the 500 Hz tone required a 32-min exposure while the 2,000 and 4,000 Hz tones required only 4-min exposures to elicit the same threshold shifts (Fig. 37.1a); (3) The peak of threshold shift in the audiogram was typically 1/2 to 1 octave above the frequency of the exposure (Fig. 37.1a–c); (4) The magnitude of TTS grew

![Fig. 37.1 Pattern of TTS from exposure to tones and noise. (a) Average TTS following exposure to either 500, 2,000, or 4,000 Hz; (b) growth of hearing loss for 2000 Hz tone at 120 dB SPL for 1, 4, or 16 min; (c) average hearing loss from exposure to band of noise (insert) at 130 dB SPL for 32 min; (d) individual subject’s exposure to 1000 Hz at 130 dB SPL. Adapted from Davis et al. [6]](image-url)
with duration of exposure; (5) There was substantial inter- and intra-subject variability (Fig. 37.1b). One subject develops less than 15 dB of TTS after 16 min while another subject develops 50 dB after only an 8-min exposure to the 1,000 Hz tone. The variability across subjects is especially puzzling given that they all had the same pre-exposure audiogram and received exactly the same noise exposure; (6) Wide band noise caused a pattern of hearing loss with a “notch” or peak ranging between 3 and 6 kHz. Since the external auditory meatus (EAM) acts like a ¼ wave resonator, the actual location of the notch (3, 4, or 6 kHz) partially depends on the length of the subject’s EAM. Larger subjects with longer EAMs tend to have notches at lower frequencies, while smaller subjects with shorter EAMs tend to have notches at higher frequencies.

The authors used binaural loudness balancing techniques to compare the loudness between an exposed and non-exposed ear. They reported a change in loudness with TTS (i.e. the degree of loudness shift is greater at low sensation levels, but the difference is reduced or disappears at high levels of stimulation). This phenomenon has been termed ‘recruitment’ [7]. With regard to frequency coding, they reported a diplacusis (i.e. for the same stimulus, the normal and ear with TTS develop different pitches). Finally, without analyzing the observation, they reported that a number of the subjects developed a buzzing or ringing in their ears which has become known as tinnitus. The tinnitus following a pure tone exposure was reported to have a much more consistent and defined pitch than the tinnitus following a noise exposure. Most of the subjects completely recovered. However, several were left with a permanent hearing loss. The results of Davis et al. [6] on the development of TTS have been expanded and confirmed by a number of investigators (series summary by Ward [8]). The early collection of research raises several questions. What is the relation between TTS and PTS in cases of more extreme exposures? What are the underlying changes in cochlear anatomy and physiology that lead to the constellation of symptoms associated with TTS and tinnitus?

Given that the audiological symptoms are essentially the same for both the TTS and PTS, it is reasonable to assume that the underlying changes in the cochlea are similar between the two conditions. However, this assumption ultimately proves to be too difficult to confirm or deny. An interesting perspective on TTS and PTS is provided in the literature on asymptotic threshold shift (ATS) [9–11]. ATS refers to the phenomenon where hearing loss grows over the first 8–24 h of a noise exposure. Hearing then stabilizes at an asymptotic level and remains at the same level for weeks or months of a constant noise exposure. If subjects are studied at different time points (i.e. 24-h exposure to 60 days), an interesting trend emerges (see Fig. 37.2). Both the 24-h subjects and the 60 day subjects have the same magnitude of threshold shift. When the 24-h subjects are removed from the noise, they begin to recover to normal sensitivity and suffer no PTS or cochlear damage. However, for subjects exposed to 60 days of noise, even though they have the same magnitude of threshold shift, when they are removed from noise they recover slowly and only partially [12]. The transition from TTS to PTS illustrates how the conditions produce the same apparent threshold shift on the audiogram but with significant differences in the underlying pathology.

Pathology of TTS

The term “TTS” suggests the pathological changes might be insignificant. However, in cases of TTS the cochlea can suffer a fairly wide spectrum of possible anatomical changes, from substantial temporary pathological damage to subtle, non-symptomatic pathological changes. Nordmann et al. [13] have shown that TTS exposure can be associated with a disconnection between the tallest outer hair cell (OHC), stereocilia, and the tectorial membrane due to changes in the structure of the organ of Corti. The assumption is partial recovery
results from structural recovery of the supporting cells and eventual reattachment of the stereocilia to the tectorial membrane. Also, the VIII nerve dendrites under the inner hair cells (IHCs) suffer excitotoxicity \[14, 15\], leading to de-afferentation of the IHC (Fig. 37.3). However, studies of IHC/VIII nerve fiber excitotoxicity with kainic acid (which mimics the effects of noise) show that the swollen VIII nerve dendrites recover and become functional again \[16\]. Consequently, part of TTS is likely due to the repairable excitotoxicity. Finally, the cochlea can sustain permanent losses of OHCs that are not sufficient in number to impair threshold detection. Collectively, the pathology associated with TTS may be repairable, or the permanent changes are too minor to be detected with typical audiological measures, so PTS would not be observed audiometrically.

**Permanent Threshold Shift (PTS)**

The relationship between the parameter of a noise and PTS are similar to TTS, but the levels required to cause PTS are higher or the durations are longer. For humans the threshold for causing PTS with years of daily repeated noise exposures is approximately 85 dBA \[17\]. The assumption is that repeated daily exposure for 5–10 years will lead to PTS. The predictive course of NIHL prepared by ISO1999 is associated with large degrees of variability, consequently making the prediction for an individual questionable. The underlying assumption of the ISO1999 procedure is that the degree of HL is related to the total energy of the exposure. The U.S. Occupational Health and Safety Administration (OHSA) considers 85 dB(A) to be the “action” level where workers are monitored and 90 dB(A) is permissible for 8 h. For each 5 dB increase in level, there is a halving of the duration (for example 95 dB(A) for 4 h is equivalent to 100 dB(A) for 2 h). In 1995, the National Institute for Occupational Safety and Health (NIOSH) prepared a recommendation for a noise standard that has a maximum tolerable exposure of 85 dB for 8 h and a 3 dB trading ratio (88 dB(A) for 2 h equals 91 dB(A) for 1 h), but the NIOSH amendment has not been enacted into law.

The effects of continuous and impulse/impact noise are different. For example, in the relationship between the noise level and ATS or PTS, for either laboratory studies or in large demographic studies, hearing loss grows at the rate of 1.7 dB for each dB of noise above the threshold for damage \[18\]. The relationship between the noise level and hearing loss changes dramatically with exposure to impulse, impact, and high level bursts of continuous noise. To illustrate, chinchillas were exposed to impact noises of equal energy, i.e. the impacts’ peak levels \(\times\) the number of repetitions were counterbalanced so that each group had equal amounts of acoustic energy (102–135 dB SPL) (Fig. 37.4) \[19\]. As seen in Fig. 37.4, the hearing loss was approximately the same for exposure to impacts of 102–119. However, above 119 dB, the hearing loss increased dramatically in spite of the equal energy that each exposure had. The interpretation of these results is that for the lower levels 99–119 dBA, the impact noises caused the same cochlear damage and HL because the ear was responding to the total energy of the exposure. However, at higher levels the hearing loss is more related to the peak level of the impact. This can suggest direct mechanical damage. This and a number of experiments with high level exposure \[11, 20\] lead/led to the formulation of the “critical level” hypothesis \[21\].
which assumes that high level exposures damage the ear causing direct mechanical failure.

The threshold of direct mechanical failure or “critical level” depends on the duration of the exposure. For example, for gunfire with peak levels of approximately 140–165 dB pSPL and impulse durations of approximately 1 ms, the “critical level” is between 150 and 155 dB pSPL peak level. For impact noise with duration of 200 ms, the “critical level” for mechanical failure is approximately 120 dBA. Short duration impulse and impact noises are shown in Fig. 37.5 [22].

When a noise exceeds the “critical level”, damage to the cochlea is immediate and direct, as seen Fig. 37.6. This figure illustrates a number of pathologies associated with exposure to “gunfire” and the resultant mechanical failures. These failures range from dramatic damage as seen in Fig. 37.6a [23], where the organ of Corti is ripped from the basilar membrane (Note the split of the cuticular plate between first and second row of OHC; this type of damage allows endolymph to bathe the OHCs and cause their death), to a more subtle damage where OHCs are separated from their Deiters’ cups (Fig. 37.6b).

When a continuous noise exposure is terminated, recovery of function proceeds almost immediately in the affected cochlear region and hearing sensitivity recovers to baseline or to a stable level of PTS. However, with exposure to high level impact/impulse noise, the time course of recovery may be complicated and tri-phasic. For example, there is a rapid recovery for 15 min–1 h, a rebound where hearing loss increases over a 2–6 h period, and then finally a slower recovery to a stable level of hearing or hearing loss.

**NIHL and Tinnitus**

There is no question about the strong correlation between NIHL and tinnitus. In the review of the clinical and experimental literature on noise and tinnitus for the military, it is stated, “... noise doses associated with hearing loss are likely to be associated with tinnitus”. However, they were not specific about the exact relationship between HL and tinnitus, i.e. the percentage of people with hearing loss that suffer tinnitus or the magnitude of the hearing loss and tinnitus. They did report that exposure to impulse noise is more likely to produce tinnitus than exposure to continuous noise. More recently, a study evaluating soldiers exposed to blast trauma in Iraq and Afghanistan found that 49% of combat personnel exposed to blasts developed tinnitus. Moreover, tinnitus ranked as the chief audiologic complaint. These new findings provide direct evidence...
of noise overexposure and subsequent tinnitus. However, more studies are needed to characterize the persistence and features of this tinnitus.

The correlation between NIHL and tinnitus remains far from perfect. There is still a large gap in our knowledge on how peripheral damage (by noise) in the cochlea leads to abnormal neural activity in the brain and the false perception of tinnitus. Two possible causes of tinnitus may be secondary neural degeneration (i.e. VIII nerve to cochlear nucleus, etc.) in the CNS or changes in the balance of excitation and inhibition in auditory pathways. Morest and colleagues [24] have reported neural degeneration in the auditory system secondary to cochlear degeneration caused by noise. The implication of the noise-induced CNS degeneration for perception is not clear. TTS, which presumably does not cause CNS degeneration, can also cause tinnitus. The alternative hypothesis for tinnitus and NIHL is a change in balance of excitation-inhibition. Salvi and colleagues [25] have experimental data showing rapid changes in the inferior colliculus and auditory cortex after NIHL. After traumatic noise exposure, the spontaneous activity of the VIII nerve remains normal, but the spontaneous activity of the cochlear nucleus can increase with “bursts” of neural responses [26]. In addition, studies of evoked potentials (inferior colliculus, auditory cortex) after acute noise exposure show an elevation of threshold as well as enhancement of the amplitude of the evoked potential [25, 27]. These findings suggest that the hearing loss caused a release of inhibition. With the development of animal models of tinnitus, we can expect more information for the relationship between cochlear pathology, changes in neural firing patterns, and tinnitus.

### Animal Models of Tinnitus

Jastreboff was the first to develop an animal model of tinnitus over 20 years ago. The initial studies looked at the effects of high doses of sodium salicylate and the development of transient tinnitus in rats. The model used a creative and straightforward lick-suppression paradigm that required discrimination between real sound and quiet. When the animals are exposed to high-dose salicylate, they fail to discriminate between quiet conditions and audible sound conditions. Since the animals failed to perceive the quiet intervals they continued to drink in the presence of a quiet/calm state. The inability to perceive the quiet state is interpreted to mean that the animals are experiencing tinnitus induced by the salicylate.

A similar technique was developed by Heffner. However, there were a number of notable differences. Heffner used an operant food-reinforced behavioral technique whereby gerbils could avoid shock if they refrained from responding during quiet intervals. Responding was allowed during sound. More importantly, Heffner used
varying levels of unilateral tone trauma (10 kHz, 124 or 127 dB SPL, for 0.5, 1, 2, or 4 h) to induce tinnitus. The key findings were threefold. First, regardless of sound intensity or duration not all animals developed tinnitus, highlighting individual differences in susceptibility of tinnitus. Second, the probability of tinnitus increased as a function of sound intensity. Finally, only long duration, high intensity tone trauma resulted in tinnitus. Tinnitus was seldom reported for low intensity or short duration trauma. Thus, there is a direct relationship between the trauma duration or level and the probability of developing tinnitus.

The most recent animal model of tinnitus relies on the acoustic startle response to a brief startling broadband or band-pass noise. Presentation of this stimulus reliably induces a large motor startle in rats that can be measured on a pressure sensitive plate. However, when a brief low intensity signal is presented before the startling sound, a significant reduction in startle amplitude is observed. This is known as pre-pulse inhibition. The acoustic signal preceding a startling sound that is audible serves to reduce the startle response. Another way of inhibiting the startle is by presenting a silent gap in a low-level continuous background noise before a startling stimulus. In this paradigm, there is always a background band-pass noise running throughout the session. At random intervals, startling sounds are presented and elicit large startle responses. On some trials, silent gaps are embedded in the continuous noise 100 ms before the startle sound. If these are detected, the amplitude of the startle response is decreased. This is known as gap-prepulse inhibition.

We have performed a number of preliminary studies to evaluate the suitability of the gap-prepulse inhibition of the acoustic startle (GPIAS) model on detecting the presence of noise induced tinnitus. When we pooled the results across a number of preliminary studies we found a direct correlation with the level of noise trauma and the probability of chronic tinnitus. When animals were unilaterally exposed to 123 dB SPL (NBN centered at 12 kHz, BW = 100 Hz, duration of 1 h) to 75% at 126 dB SPL (NBN centered at 12 kHz, BW = 100 Hz, duration 1 h). Pharmacologically-induced transient tinnitus with a high dose of sodium salicylate (250 mg/kg, 1 h pre-session, i.p.) yielded evidence of tinnitus in all the animals tested. Group sizes were 12, 12, and 24 rats (Harlan SASCO Sprague Dawley, adult males, mean body weight 375 g).

In addition to the duration of tinnitus, we were also interested in the pitch of noise-induced tinnitus. Evidence from human studies suggests that there is a relationship between the frequency of the maximal hearing loss and the pitch of the tinnitus. When animals were unilaterally exposed to 12 kHz noise at 123 dB SPL, tinnitus was observed between 12 and 16 kHz (Fig. 37.8). Immediately after the noise exposure, however, animals failed to detect gaps at multiple frequencies. This effect disappeared within 24 h, but evidence of tinnitus remained in the 12–16 kHz region. Increasing the level of the unilateral 12 kHz NB noise to 126 dB SPL led to a nearly complete loss of gap-induced prepulse inhibition at 16 kHz (Fig. 37.9). Changing the center frequency of the noise from 12 to 16 kHz resulted in the maximum loss of gap-induced prepulse inhibition occurring at 20 kHz instead of 16 kHz (Fig. 37.10). Audiometrically, these changes in the “pitch” of the tinnitus would seem to be related to a shift in the location of maximal OHC trauma in the cochlea, but that relationship has yet to be confirmed anatomically.
One limitation of the GPIAS model is that the startle response is dependent on binaural hearing. If the unilateral acoustic trauma is excessive, the startle stimulus is less effective at producing a strong startle response. Because of this, it is advantageous to limit the hearing loss to the high frequencies. There is also clinical value of limiting the NIHL as tinnitus induced by noise tends to be perceived at higher frequencies. The startle stimulus can also be moved so that it is a band-pass noise within the audible range of even the exposed ear. This can increase the effectiveness of the startle stimulus following the exposure.

Despite the gaps of knowledge that still exist regarding the biological basis for tinnitus and the basis for tinnitus susceptibility, a number of research groups have been steadily narrowing the gaps. Progress is likely to accelerate as animal models continue to be developed and act as a platform for basic science and pre-clinical drug therapy models. However, challenges still remain for understanding tinnitus. However, NIHL is known to be one of the key catalysts for the development of chronic tinnitus. A concerted effort using animal models, human and animal imaging studies, physiological, behavioral, and pharmacological studies will likely enhance our knowledge base and move us closer to providing strategies to reduce the impact of tinnitus.
Conclusions

The hearing loss caused by exposure to noise can be either temporary or permanent. In addition to a loss of hearing sensitivity, traumatic noise exposure degrades signal detection in background noise, reduces the dynamic range of loudness, and can induce tinnitus. The deleterious effects of noise are related to each of the primary dimensions of sound: frequency, intensity, and duration of exposure. Our current noise standards are over 40 years old (from 1968), and do not reflect modern scientific research or our understanding of the effects of noise. For example, research has shown that certain types of noise exposure (combinations of continuous noise with impulse/impact noise) or noise combined with ototoxic solvents pose an increased risk to hearing compared with simple continuous noise exposures. Since the initial noise legislation of 1968, much has been learned about the mechanisms through which noise causes hearing loss, and in the last 10 years, much progress has been made in unraveling the mystery of noise-induced tinnitus.

References

Chapter 38
Tinnitus and Ménière’s Disease

Yu-Lan Mary Ying and Moises A. Arriaga

Keypoints

1. Ménière’s disease is characterized by a triad of symptoms: fluctuating hearing loss, attacks of vertigo, and tinnitus. Some authors have included aural fullness.
2. Patients often seek treatment for the severe vertigo attacks, but may have had other otologic symptoms for some time prior to the onset of vertigo.
3. Tinnitus that occurs in Ménière’s disease is best characterized as a low pitched, narrow band of noise, resembling a “roaring sound.”
4. The tinnitus may change with the fluctuations in hearing loss, and tinnitus increases as hearing loss worsens with the progression of the disease.
5. During the active phase of Ménière’s disease, the vertigo can be debilitating and dominating the symptoms.
6. As the disease stabilizes, the tinnitus can become a serious and a severe problem.
7. It is believed that endolymphatic hydrops (imbalance in volume of the fluid systems of the inner ear) is the cause of the symptoms of Ménière’s disease, but there is still uncertainty regarding many aspects of the pathology of the disease.

Keywords  Tinnitus • Cochlear implants • Promontory stimulation • Treatment

Abbreviations

AAO-HNS  American Academy of Otolaryngology – Head and Neck Surgery
DHI  Dizziness Handicap Inventory
HHIA  Hearing Handicap Inventory for Adults
PA  Pure tone average
THI  Tinnitus Handicap Inventory

Introduction

Ménière’s disease is defined by the presence of three symptoms (or four): intermittent vertigo, fluctuating hearing loss, and tinnitus with aural fullness occurring on one side. Some authors have added aural fullness as a fourth symptom. It was first described by Prosper Ménière in his original publication in 1861. Its diagnosis is largely based on the clinical history and hearing tests. Individuals with Ménière’s disease experience incapacitating attacks of vertigo, associated with nausea and vomiting lasting for hours [1]. The sudden attacks of vertigo last anywhere from 30 min to several hours, with unilateral hearing loss occurring together with tinnitus; often aural fullness is present as well. Audiological findings include fluctuating low frequency and progressive sensorineural hearing loss with tinnitus. The course of Ménière’s disease is unpredictable and highly variable among individuals and can be accompanied with periods of remission. Disequilibrium may persist for 24–72 h after the attack before resolving completely.

Tinnitus may be the first symptom of Ménière’s disease and may precede the remaining symptoms by months or years. Fluctuating cochlear signs, such as tinnitus, hearing impairment, and/or fullness in the ear...
were present prior to onset of the first vertigo attack in more than 50% of patients in a study by [2].

Criteria for the diagnosis of Ménière’s disease can be divided into four categories as possible, probable, definite, and certain (Table 38.1). Furthermore, scales of dizziness [i.e., Dizziness Handicap Inventory (DHI)], hearing loss [i.e., Hearing Handicap Inventory for Adults (HHIA)], and tinnitus [i.e., Tinnitus Handicap Inventory (THI)] have been developed to quantify the symptoms associated with Ménière’s disease.

The Committee on Hearing and Equilibrium of the American Academy of Otolaryngology – Head and Neck Surgery (AAO-HNS) published its most updated guidelines for defining, reporting, and interpreting results of the treatment of Ménière’s disease in 1995 [3]. (For detailed discussion see Chap. 60)

### Table 38.1 Diagnosis of Ménière’s disease

<table>
<thead>
<tr>
<th>Certain Ménière’s disease</th>
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<tr>
<td>Definite Ménière’s disease, plus histopathologic confirmation</td>
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<table>
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<tr>
<th>Definite Ménière’s disease</th>
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<tbody>
<tr>
<td>Two or more definitive spontaneous episodes of vertigo 20 min or longer</td>
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<tr>
<td>Audiometrically documented hearing loss on at least one occasion</td>
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<tr>
<td>Tinnitus or aural fullness in the treated ear</td>
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<td>Other causes excluded</td>
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<table>
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<tr>
<th>Probable Ménière’s disease</th>
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<tbody>
<tr>
<td>One definitive episode of vertigo</td>
</tr>
<tr>
<td>Audiometrically documented hearing loss on at least one occasion</td>
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<tr>
<td>Tinnitus or aural fullness in the treated ear</td>
</tr>
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<td>Other causes excluded</td>
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<tr>
<th>Possible Ménière’s disease</th>
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<tr>
<td>Episodic vertigo of the Ménière type without documented hearing loss or Sensorineural hearing loss, fluctuating or fixed, with disequilibrium but without definitive episodes</td>
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<td>Other causes excluded</td>
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The prevalence increases linearly with age up to 60 years. There is a slight female preponderance, and the typical age of onset is 30–60 years. Bilateral Ménière’s disease incidence ranges from 10 to 70%, increasing the frequency with time [14]. Genetic predisposition has been reported in families with Ménière’s disease [15, 16]. Factors such as diet, weather changes, as well as emotional and physical stress can also precipitate vertigo attacks and make any symptom complex and worse. Some of the variance in the incidence reported by different investigators may have been caused by differences in the definition of Ménière’s disease.

The symptoms of Ménière’s disease are signs of an imbalance in the volumes of the fluids in the inner ear, known as endolymphatic hydrops [17]. Cochlear hydrops causing fluctuating low-frequency hearing loss, tinnitus, and aural fullness without associated vertigo may precede Ménière’s disease with eventual development of the full syndrome occurring in 37–42% of patients [18]. Lermoyez’s syndrome is a variant of Ménière’s disease, where hearing loss and tinnitus precede an attack of vertigo by days or months, with improvement of hearing after vertigo episodes. Patients with severe and long-term Ménière’s disease are at risk of developing Tumarkin otolithic crisis or drop attacks of falling because of loss of lower-extremity muscle tone without loss of consciousness. The incidence is reported as 7% of patients with Ménière’s disease to as high as 72% in one report [19]. The cause is thought to be a sudden stimulation
of the vestibular end organs by shift of the utricular macula or rupture of inner ear membranes, but the exact cause is unknown [20].

The frequency of vertigo attacks varies widely with a mean of 6–11 episodes per year [21]. Spells tend to change in severity over a period of time, becoming milder but still unpredictable. The decrease in hearing and tinnitus can occur before or during the vertigo attack. Typically, the disease eventually “burns out” with the decline and cessation of vertigo, but there is progressive deterioration of hearing. Hearing fluctuates in the early course of disease but eventually becomes progressively worse, stabilizing at about 50 dB pure tone average (PA) and 50% word discrimination score [18]. The hearing loss in Ménière’s disease begins at low frequencies, thus, different from many other causes of hearing loss such as noise induced hearing loss, hearing loss caused by ototoxic substances, and presbycusis that mainly affect hearing at high frequencies. Eventually, hearing loss in individuals with Ménière’s disease involves all frequencies. Tinnitus that occurs in Ménière’s disease is often described as a harsh roaring machine-like sound that is more pronounced during vertigo attacks. Hyperacusis (decrease tolerance to sounds, see Chap. 3) and distortion of sound in the affected ear can also be present. Tinnitus and aural fullness prevail during life in the majority of individuals with Ménière’s disease [22].

Pathophysiology

Despite the long history of Ménière’s disease, the etiology and pathophysiology of this condition are still unknown. Most hypotheses of the pathogenesis of Ménière’s disease include anatomical abnormalities of the endolymphatic fluid system, but other hypotheses involve viral infection, autoimmune disease, allergy, and activation of neural plasticity. That endolymphatic hydrops is the cause of the symptoms of Ménière’s disease has been supported by histopathological findings, although not all patients with the histopathology have the typical symptoms [23, 24]. The hydrops are believed to be caused by mechanical obstruction to endolymphatic flow or by intrinsic malfunction of the endolymphatic system, resulting in an overabundance of endolymphatic volume and/or pressure [25]. Dysfunction of the spiral ligament fibrocytes, which interferes with the recycling of K+ ions and results in osmotic imbalance, can cause expansion of the endolymphatic compartment [26].

Schuknecht developed a theory of gradual distention of the endolymphatic system that leads to a rupture of membranous labyrinth and sudden release of a large volume of endolymph into perilymphatic space [20]. It is suggested that sensory and neural structures are injured from exposure to the potassium-rich endolymph, resulting in vertigo and hearing loss. As the rupture heals and homeostasis is restored in the inner ear compartments, the symptoms subside. Dohlman (1980) [27] suggested that increase of potassium occurs in the perilymph during a Ménière’s attack and that potassium-rich fluid surrounding the vestibular nerve is the cause of the experienced vertigo. Zenner (1987) [28] found that perilymphatic potassium intoxication leads to a longitudinal contraction of the outer hair cells. This results in their decoupling from the tectorial membrane. Dulon et al. (1987) [29] demonstrated that small changes in the osmolarity of the surrounding in vitro medium induce fast contractions (hypo-osmotic solution) or elongations (hyper-osmotic solution) in isolated outer hair cells. However, the hypotheses that assume that Reissner’s membrane ruptures before an attack occurs have been questioned.

Another theory proposed by Gibson and Arenberg (1997) [30] is a disturbance in longitudinal flow of endolymph from the cochlear duct to the endolymphatic sac due to a narrow vestibular aqueduct, resulting in hydrops.

It has been suggested that longitudinal flow was involved in maintaining endolymph homeostasis. However, measurements of the dispersal of markers in the endolymph [17] have failed to support these hypotheses. These measurements were interpreted to suggest that the normal state of the endolymph is maintained without a significant involvement of volume flow at all [17]. The situation is different in abnormal states, such as is assumed to be present in Ménière’s disease where the volume of inner ear fluid is abnormal. In such situations, the longitudinal volume flow of the endolymph may contribute to homeostasis.

The role of the endolymphatic sac is complex and poorly understood. It seems to act as a “bidirectional overflow” system that responds to the endolymph volume disturbance [17].
In other hypotheses, the sac is postulated to actively regulate the flow by maintaining an osmotic gradient and secreting glycoproteins that attract movement of the endolymph toward the sac. The sac produces sac-cin, a hormone thought to increase the volume of endolymph, which may promote faster flow [30].

It has been suggested that the endolymphatic sac is primarily responsible for the immuno-defense of the inner ear [31]. It is hypothesized that a viral infection leads to an inflammatory immune and microvascular-mediated injury. Circulating immune complexes and serum auto-antibodies to inner ear antigens are greater in Ménière’s disease patients than in controls [32–34]. This suggests that circulating immune complexes may be involved in the pathogenesis of Ménière’s disease, either as a direct cause of damage or as a by-product of an underlying autoimmune abnormality [33]. This hypothesis is supported by the clinical experience of beneficial effect of treatment with corticosteroids [35].

As early as in 1923, Duke (1923) [36] proposed an allergic theory for Ménière’s disease. However, it was not until 1970s that studies showed an improvement in the symptoms of this disease after desensitization to inhalant allergens and an elimination diet for allergies to food.

Pulec (1973) [37], in discussing Ménière’s disease, reported allergies were related to the sensorineural hearing loss and symptoms of Ménière’s disease among 36% patients. Fourteen percent of his Ménière’s disease patients responded to allergy treatment. In a case-control study, Derebery and Valenzuela (1992) [38] found an inhalant allergy in 41.6% and a food allergy in 40.3% of patients with Ménière’s disease answering a self-reported questionnaire, in comparison with rates of 27.6 and 17.4% in their control population.

Furthermore, a significant percentage of patients with Ménière’s disease and allergy showed improvement in both allergy and Ménière’s symptoms when treated with desensitization and diet control [39]. Hence, symptoms of food allergy should be questioned for patients with endolymphatic hydrops and fluctuant hearing loss, as suggested by Shambaugh and Wiet (1980) [40].

Inspired by the benefits from treatment with air-pressure (pressure chamber) [41–43] by applying air-puffs to the inner ear (using the Meniett device), it has been suggested that activation of neural plasticity may be involved in creating the symptoms of Ménière’s disease [44].

### Tinnitus Associated with Ménière’s Disease

Tinnitus in Ménière’s disease is best characterized as a low pitched, narrow band of noise, usually described as a “roaring sound”, corresponding to the low-frequency sensorineural hearing loss [45–47]. In the early stages of the disease, tinnitus may be intermittent. As the disease progresses, tinnitus becomes permanent, but its intensity fluctuates. Hearing loss and tinnitus normally increase over time. After a long time, the end state of Ménière’s disease “burnt-out” where the effects of vertigo attacks have ceased, tinnitus may become the most disturbing complaint.

As Stahle (1988) [48] described in his results of an epidemiologic study in Sweden, the tinnitus quality fluctuated in its intensity and paralleled the control of vertigo symptom and ear blockage. In a separate group of patients with control of their chief vertigo complaint, the ear blockage persisted, as did the tinnitus. Herraiz et al. (2006) [49] found a statistical association between tinnitus intensity and worse hearing loss or hyperacusis in 102 patients with Ménière’s disease, uninfluenced by the number of vertigo spells. In the initial phases of Ménière’s disease, tonal-tinnitus is usually not present, as opposed to in the later stages of the disease where tonal-tinnitus is described by a number of patients [50].

### Pathophysiology of Tinnitus in Ménière’s Disease

Tinnitus in Ménière’s disease may be caused by similar mechanisms as other forms of tinnitus that are related to injuries of the cochlea (see Chap.10). Hearing loss may cause tinnitus through the effect of deprivation of input to the auditory system that activates neural plasticity (see Chap. 12).

### Management and Treatments

There is no known cure for Ménière’s disease, and treatments are aimed at reduction of its symptoms. Vertigo is often the most debilitating symptom of
Ménière’s disease, and most treatments focus on relieving this symptom. The tinnitus of Ménière’s disease may well remit with improvement in low-frequency hearing as a result of medical or surgical treatment. For a detailed discussion of treatment of Ménière’s disease, see Chap. 60.

Treatments specifically directed toward tinnitus in Ménière’s disease are similar to treatments of other forms of tinnitus described in the chapters in Part VI of this book.

Neural Plasticity

The reason that overpressure can relieve symptoms of Ménière’s disease is not known, but it has been hypothesized that neural plasticity is involved in at least one or more of the symptoms of Ménière’s disease [51]. These symptoms are assumed to be caused by an imbalance of the volumes of the fluid in the inner ear [17], the causes of which are unknown. The finding that applying air-puffs to the inner ear can ameliorate the symptoms, thus stimulating the vestibular sensory cells, indicates that functional abnormalities may be involved in causing the symptoms of Ménière’s disease and an activation of neural plasticity may be involved.

References

Chapter 39
Tinnitus and Vestibular Schwannoma: Overview and Clinical Correlations

Jason May, Virginia Ramachandran, and Anthony T. Cacace

Keypoints

1. Historical overview considering aspects of medical and audiological evaluation of patients with vestibular schwannoma and tinnitus.
2. Treatment modalities and outcomes are discussed together with a review of relevant postsurgical issues.
3. The value of asymmetrical hearing loss, vestibular complaints, and facial nerve problems in establishing the index-of-suspicion for medical and allied-health professionals regarding the presence of vestibular schwannoma.
4. It is indicated that it is difficult to predict the likelihood of tinnitus following microsurgery for tumor resection.
5. Stereotactic radiosurgery does not appear to have a substantial influence on tinnitus.
6. Atypical forms of tinnitus may occur as postsurgical complications, particularly when hearing is lost completely and abruptly during surgery.
7. Tinnitus after severing the auditory nerve causing unilateral deafferentation of the auditory periphery causes a cascade of reactive changes in the peripheral and central nervous systems that can result in anomalous forms of cross-modal plasticity.
8. Gaze-evoked, gaze-modulated, and other forms of somatic (cutaneous-evoked) tinnitus can result.

Keywords
Vestibular schwannoma • Acoustic neuroma
• Tinnitus • Gaze-evoked tinnitus • Cutaneous-evoked tinnitus • Audiology • Neuro-otology • Microsurgery
• Neurosurgery • Gamma knife radiosurgery

Abbreviations
CPA Cerebellar pontine angle
ABR Auditory brainstem response
IAC Internal auditory canal
CT Computerized tomography
MRI Magnetic resonance imaging
fMRI Functional magnetic resonance imaging
LINAC Linear accelerator
PET Positron emission tomography
GET Gaze-evoked tinnitus
Cm Centimeter
Mm Millimeter
TCR Trigemino-cardiac reflex
NF2 Neurofibromatosis type 2, NF2

Introduction

Background

Vestibular schwannoma, earlier known as acoustic neuroma or vestibular neuroinoma, is a benign tumor of the VIIIth cranial (cochlear–vestibular) nerve that constitutes between 6 and 10% of all intracranial tumors and 80–90% of all tumors of the cerebellopontine angle (CPA [1]).

While vestibular schwannoma has been recommended as the official nomenclature of this disease [2], the terms acoustic tumor, acoustic neuroma, and vestibular neurinoma remain interchangeable among clinicians.
The incidence of vestibular schwannoma is estimated at 17.4 cases per million [3]. These tumors originate from the nerve sheath and consist of Schwann cells in a collagenous matrix; typically arising from within the bony portion of the internal auditory canal (IAC) at the myelin–glial junction (Obersteiner-Redich zone) and growing outward from the porus acusticus to the CPA [1]. They are generally well-circumscribed and produce symptoms by displacing adjacent anatomical and neural structures without invasion.

Vestibular schwannomas are typically slow growing [average rate-of-growth: 0.2 cm per year], but can grow as rapidly as 2 cm per year. Initially, symptoms arise due to VIIth and VIIIth nerve compression, but as the tumor size increases (2–3 cm), the fourth ventricle can become compressed and hydrocephalus can result from total obstruction; trigeminal symptoms can also occur once the tumor exceeds 3 cm. If growth continues, brainstem compression, cerebellar-tonsillar herniation, and death can result.

In this class of tumors, 95% are unilateral, non-hereditary, and manifest in individuals between the ages of 40–60 years [1]. The exception is Neurofibromatosis type 2 (NF2), a disease characterized by bilateral vestibular schwannoma, usually presenting before 21 years-of-age. Neurofibromatosis type 2 can be inherited through an autosomal dominant transmission or as a result of de novo mutations. The genetic mutation responsible for NF2 is caused by a defect located on chromosome 22, band q11-13.1. Additionally, these patients also have higher occurrence rates of meningiomas, ependymomas, and Schwannoma of other cranial nerves.

Presenting Symptomatology

The most common presenting symptom of vestibular schwannoma is asymmetrical sensorineural hearing loss. Loss of pure tone hearing sensitivity typically progresses slowly over many years and is often, but not always, accompanied by reduced performance on tests of monosyllabic word recognition, absence and/or decay of the acoustic-stapedius reflex, and abnormalities of the auditory brainstem response (ABR). A harbinger of this disease is the occurrence of word recognition scores that are much worse than would be expected based upon the pure-tone average (0.5, 1, 2 kHz, respectively) or the roll-over effect for monosyllabic words, if a complete psychometric function (percent correct word recognition vs. stimulus level) is generated (e.g., [4]). Decreased pure tone hearing sensitivity and other audiometric abnormalities are thought to occur as a consequence of direct injury to the cochlear-vestibular nerve, interruption of the cochlear blood supply, brainstem involvement, or a combination of factors [5]. Indeed, unilateral tinnitus can be the initial and only presenting symptom signaling tumor presence [6]. Recent estimates indicate that approximately 10% of individuals with vestibular schwannoma will present themselves in this manner [7]. The presentation of unilateral tinnitus, in and of itself, is an indication for further work-up.

Although vertigo is not a commonly presenting symptom, disequilibrium or unsteadiness can be seen in 40–50% of patients. Presumably, if destruction of vestibular nerve fibers is sufficiently slow, most patients will compensate over a period of time. Headaches are found in 50–60% of patients, but are rarely observed as a presenting symptom. Similarly, facial-nerve weakness is also rare, occurring less than 1% of the time as a presenting symptom [7].

Management Options and Their Relationship to Tinnitus

The treatment options for vestibular schwannoma include planned observation, surgical extirpation, and radiation. With respect to the first option, because these tumors are typically benign and slow growing, they can be carefully observed over a period of time and re-evaluated to determine if more aggressive treatments are necessary. This strategy is often applied in older adults that are poor candidates for surgery, in individuals with small tumors, in instances when the involved side is the only hearing ear, and/or if tumor growth is less than 2–3 mm per year.

Since the late 1800s, surgical procedures have undergone vast improvements. The current goals of surgery are: (1) complete tumor extirpation to cure the underlying disease, (2) preservation of facial nerve function, and when possible, (3) preservation of hearing. However, depending on size and growth patterns, these ambitious goals cannot always be attained and consequently, surgical co-morbidities remain as a distinct reality.
To appreciate the evolution of this field, we provide here some historical background and relevant vignettes based on the more detailed review of Machinis and colleagues [8]. Briefly, the first documented surgery performed by von Bergmann (1890) was unsuccessful as the patient died prior to localization of the tumor. Sternberg (1900) is credited with the first accurate pathological description of the tumor. In a two-stage operation circa 1894, Sir Charles Ballance described the finger dissection of vestibular schwannoma, which he called an “encapsulated fibro-sarcoma” (Ballance 1907) [9]. While the first decade of the twentieth century saw continued, albeit limited attempts at tumor resection, the failure rate of surgery was alarmingly high. Apparently, the high failure rate was due to small craniotomies leading to cortical herniation and eventual death.

With respect to the suboccipital approach, several key individuals (e.g., Woolsey, Fraenkel, and Krause) made definitive advancements [10]. Subsequently, Panse (1904) [11] proposed, but never performed, a surgical technique through the petrous bone, which would later be described as the translabyrinthine approach (see [12] for additional historical insights). However, at this early juncture, the translabyrinthine approach did not gain popularity because it did not provide adequate access to the CPA and because there was a high rate of meningitis postoperatively. Consequently, it was not until Cushing described the bilateral suboccipital approach that complications and mortality rates approached what might be considered acceptable levels [13].

The surgical approach advocated by Cushing aimed at subtotal intracapsular resection. Initially, morbidity and mortality rates were high (approximately 75% and 40, respectively); however, after 30 operations, Cushing reduced the mortality rate to 20% and with more experience, the mortality rate dropped to 7.7% (based on a series of 176 cases). Cushing advanced the field by avoiding herniation and medullary compression during surgery and by making a large curvilinear incision between both mastoid processes [13]. Nevertheless, a disappointing 5-year survival rate of 50% could not be overcome, due in large part to tumor recurrence.

Subsequently, Cairns (1931) reported the first complete removal of a vestibular schwannoma with preservation of facial-nerve function. Progress in this area continued as Olivecrona reported an impressive 40% facial nerve preservation rate, with recovery of facial-nerve function in another 20% [8]. Improvements in diagnosis, surgical care, and advances in the suboccipital approach further improved mortality rate to approximately 2.4% [14].

Introduction of the operating microscope was a key element to advancing surgical outcome, preservation of hearing and facial-nerve function and reduction of other morbidities [8]. This instrumental innovation also allowed for the perfection of the translabyrinthine approach and for the introduction of the middle cranial fossa approach. Other advancements came with the advent of the polytime Pantopaque imaging technique [14], which led to the identification of small tumors in the IAC. With the available imaging techniques and use of the operating microscope, House and colleagues lowered the mortality rate to the range of 0.8–5% [15]. Use of the auditory brainstem response (ABR), use of the ABR, computerized tomography (CT), and magnetic resonance imaging (MRI) further revolutionized diagnosis. Thus, based on these historical trends and innovations, three surgical approaches have withstood the test-of-time and are considered the mainstay of the surgeon’s arsenal. These include the translabyrinthine, retrosigmoid/suboccipital, and middle cranial fossa approaches, each having specific advantages and disadvantages.

In the contemporary surgical literature, William House is credited with perfecting the translabyrinthine approach [7]. Because no brain retraction is used, this technique is considered by some to be the safest approach. In the absence of a high riding jugular bulb and/or an anteriorly placed sigmoid sinus, the translabyrinthine approach provides excellent access to the lateral CPA. Thus, with the fundus and lateral portion of the IAC completely exposed, the facial nerve can be dissected easily. The disadvantage of this approach is that hearing is always sacrificed during surgery.

The retrosigmoid/suboccipital and middle cranial fossa approaches are considered the procedures-of-choice in cases where hearing preservation is attempted. While the retrosigmoid/suboccipital approach can be applied to all vestibular schwannoma, of the available procedures, it has the advantage of providing the widest exposure and best operative field visualization of the posterior fossa. However, limited access to the lateral CPA near the IAC, poor visualization of the VIIth and VIIIth cranial nerves, and the potential for cerebrospinal fluid leaks, both peri- and postoperatively, are clear disadvantages.

The middle cranial fossa approach, also introduced and perfected by William House, is considered the
procedure-of-choice for small intracanalicular tumors. Advantages of this approach include access to the lateral third of the IAC and the fact that it is an extra-dural procedure. The disadvantages include risk to the facial nerve, limited access to the posterior fossa, potential for dural laceration (particularly in patients over 65), and the need for temporal lobe retraction. As the facial nerve lies on the anterior surface of the canal, tumor removal can be hindered by this approach.

**Stereotactic Radiosurgery**

In addition to the surgical procedures noted above, stereotactic radiosurgery has become a viable management tool. This treatment modality can be used either alone or in combination with microsurgery, particularly in those instances where tumors are incompletely excised. While different types of radiosurgery are available [linear accelerator (LINAC) and gamma knife], gamma knife is the approach used most frequently. Gamma knife treatment involves a single-session application of collimated beams of cobalt radiation to a localized intracranial location. This approach allows for high doses of radiation to the location of the lesion while registering smaller doses to surrounding structures, thus minimizing morbidity [16]. The effectiveness of this method occurs through interference with the cellular life-cycle, thereby inhibiting growth of the tumor. Individuals, who have tumors less than 2.5–3.0 cm including the absence of or limited tumor-related symptoms, are considered candidates [17, 18]. Age, hearing status, facial movement, facial sensation, balance, vertigo, and tinnitus are other factors that are given consideration in patient selection [19]. As gamma knife radiosurgery requires only a local anesthetic and has a relatively fast recovery time, it is an appealing and often a superior option for individuals who may not be medically able to undergo surgery.

Outcomes of gamma knife radiosurgery are considered good when viewed in terms of tumor control and cranial nerve morbidity, particularly with the advent of improved dosing strategies. However, hearing and balance function may be adversely affected [20]. Direct comparison of the effectiveness of gamma knife radiosurgery to microsurgery is difficult because tumor size and patient characteristics are confounding variables and randomization of patient selection has been difficult. Because gamma knife radiosurgery is typically limited to smaller tumors, better facial-nerve function and hearing outcomes have been observed with this modality. However, severity of tinnitus and vertigo are generally unchanged following this type of treatment [21].

As with any treatment modality, complications are inevitable. With gamma knife radiosurgery, complications can include facial twitching, weakness, numbness, pain, trigeminal-nerve dysfunction, watery or dry eyes, hydrocephalus, hearing loss, tinnitus, balance disturbances, and vertigo [18, 22]. Some post-treatment effects, such as hearing loss, may have a delayed onset [16], although lower doses of radiation have been shown to reduce complication rates [23]. Gamma knife radiosurgery may also be associated with a small risk for developing radiation-induced malignancies [20], and in cases where enlargement of the tumor occurs, surgery may be necessary [24]. Unfortunately, in those instances when tumors fail to respond to radiation, management through salvage surgery may be more difficult, thus resulting in poorer outcomes [25].

**Tinnitus and Vestibular Schwannoma**

While it has been estimated that tinnitus is the sole symptom in 10% of patients presenting with vestibular schwannoma, it is present in 60% along with asymmetrical hearing loss. In their series of individuals undergoing hearing preservation surgery, Levo et al. (2000) [26] observed that tinnitus worsened postoperatively in 6–20%, remained unchanged in 25–60%, and improved or resolved in 30–50% of individuals. Based on logistic regression analysis to determine which of eight independent variables were prognostic for the presence or absence of postoperative hearing, Rastogi and colleagues (1995) [27] found that porus acousticus widening was the best prognostic indicator. In this series, the presence or absence of tinnitus did not play a significant role in their outcome data. In studying tumor size and age, Fahy et al. (2002) [28] were unable to predict tinnitus outcome postoperatively. They also failed to show a statistically significant association between changes in tinnitus and quality-of-life. Nevertheless, they found that tinnitus improved in 16%, was unchanged in 55%, and actually worsened in 29% of patients. Based on these data, it remains unpredictable which patients will have worsening tinnitus postoperatively.
There are several surgical factors that have been examined in regard to tinnitus symptom outcomes. For example, Schaller et al. (2008) [29] showed that the intraoperative occurrence of the trigemino-cardiac reflex (TCR) during surgery was a negative prognostic factor not only for hearing preservation but also for the presence of ipsilateral postoperative tinnitus. The TCR occurred in 17% of patients undergoing tumor resection; 60% of those patients had postoperative tinnitus vs. 17% of those who did not have TCR. Kanzaki et al. (1999) [30] addressed the question of surgical technique on postoperative tinnitus. Based on questionnaires obtained from 202 patients, they found an increase in tinnitus in those individuals undergoing hearing preservation surgery vs. those patients undergoing translabyrinthine surgery. In cases where hearing preservation was attempted, tinnitus was present in 78.6% preoperatively and increased to 89.3% postoperatively. Individuals who underwent a translabyrinthine approach decreased from 72.7% preoperatively to 67.3% postoperatively. The outcome of surgery to preserve hearing showed no predictive value with respect to tinnitus occurrence or estimated tinnitus loudness. This study indicates that while hearing preservation is a distinct surgical outcome, individuals remain at risk of developing tinnitus. In a series of vestibular schwannoma surgeries using the middle cranial fossa approach (n=311) where the facial nerve was preserved 99% of the time and where hearing was preserved 49% of the time (in smaller tumors), Haid (1998) [31] found that 45% of individuals also had a reduction in tinnitus severity.

**Tinnitus and Gamma Knife Radiosurgery**

Tinnitus outcomes from gamma knife radiosurgery treatments tend to be reported by the presence/absence of tinnitus or use of a visual-analog scale as a method to gauge tinnitus severity. As noted previously, while available data suggest that tinnitus tends to remain unchanged following this treatment modality, the type of outcome measures used to track these changes lack the necessary precision to demonstrate clinically significant changes. To our knowledge, no attempts have been made to measure relevant psychoacoustic parameters of tinnitus perception, such as pitch and loudness, prior to and following gamma knife radiosurgery. Based on a nonrandomized prospective study of 63 patients who underwent gamma knife radiosurgery and 28 patients who underwent microsurgery using a suboccipital approach, Myrseth and colleagues (2009) [21] found no change in tinnitus severity using a visual-analog scale for either group, despite the fact that decreases in hearing sensitivity were observed in both groups. In a retrospective questionnaire-based study and chart review comparing gamma knife radiosurgery with translabyrinthine microsurgery in individuals with non-serviceable hearing and small tumors, Coelho and colleagues (2008) [19] noted that tinnitus was present post-treatment in 7/21 (33%). This occurred in 2/12 (17%) in the gamma knife radiosurgery group; and 5/9, 56% in the translabyrinthine microsurgery group). Of those in the gamma knife group without tinnitus, 9/10 showed no post-treatment changes, while one had new onset of tinnitus. Of those with tinnitus prior to treatment, both noted post-treatment changes. Of four individuals in the translabyrinthine group without tinnitus preoperatively, one noted new onset tinnitus. Of the five that had preoperative tinnitus, one noted improvement and two had no change and got worse. While this information is of interest, the small sample size limits the generalizing power of these data. In a retrospective analysis of 123 patients with vestibular schwannoma treated with gamma knife radiosurgery, Hempel et al. (2006) [32] found that the presence or absence of tinnitus remained stable in 90% of patients. In this sample, approximately 4% reported tinnitus onset, while 6% reported tinnitus cessation following treatment. Interestingly, seven patients also reported improvement in hearing sensitivity, but there is no report of how hearing loss may or may not have correlated with changes in tinnitus. Bertalanffy and colleagues (2001) [33] found that in a group of individuals undergoing gamma knife radiosurgery with preoperative tinnitus (n=32), tinnitus improved in six cases (46%), was unchanged in five cases (38%), and worsened in two cases (15%). Tinnitus appeared as a new symptom in one patient. Régis and colleagues (2002) [34] reported a decrease in tinnitus for approximately 16% of patients following gamma knife radiosurgery. Niranjan and colleagues (1999) [35] found that of 29 patients with intracanalicular tumors who underwent gamma knife radiosurgery, 7/13 who had preoperative tinnitus continued to experience tinnitus at a long-term post-surgical follow-up. The post-treatment stability of tinnitus is interesting in light of hearing preservation outcomes with gamma...
knife radiosurgery. Based on the Gardner–Robertson classification scheme [36], hearing outcomes tend to fluctuate compared to changes in tinnitus. Given the typical high correlation of tinnitus to hearing loss, additional audiometric information may be helpful in future studies better understand these effects. Régis and colleagues (2008) [37] provided some information regarding the relationship between hearing loss and tinnitus by noting that in a sample of 184 patients followed for 3 years or longer, the presence of tinnitus preoperatively was viewed a protective factor for hearing preservation, although it was not speculated why this might be the case.

The mechanism(s) of pathology for complications of gamma knife radiosurgery may include effects of tumor swelling prior to shrinkage, cochlear-nerve toxicity, and/or damage to cochlear structures due to radiation exposure. Direct radiation-induced cochlear-nerve toxicity has been speculated to be a causative factor for hearing loss [38, 39], which suggests that the basal turn of the cochlea near the modiolus and the inferior-most extension are most susceptible to high radiation doses. Using the ABR as a physiologic measure to distinguish cochlear from retro-cochlear effects, Bertalanffy and colleagues (2001) [33] showed that that cochlear function was generally unaffected by this type of treatment.

**Atypical Forms of Postsurgical Tinnitus**

Even with the best of efforts and skills of the neurosurgeon and neuro-otologist, hearing can be lost completely and abruptly during microsurgery. While the consequence of profound unilateral hearing loss occurs as a direct result of the translabyrinthine procedure, if the tumor size exceeds 2 cm, a high probability of profound unilateral hearing loss is also expected [27], regardless of the surgical approach used. Consequently, complete loss of hearing in the course of tumor resection results in a unilateral deafferentation of the auditory periphery. This acute injury sets the stage for a cascade of reactive changes in afferent/efferent pathways that can result in gaze-evoked, gaze-modulated, and/or other forms of somatic tinnitus.

In its purest form, gaze-evoked tinnitus (GET) is a phenomenon whereby horizontal or vertical deviation of eye position, from a neutral head-referenced position, results in an auditory sensation. While exact mechanisms are unknown, it has been postulated that these cross-modal effects may occur from reactive sprouting of neurons to unoccupied (denervated) synaptic sites, unmasking of silent synapses, and/or ephaptic interactions [40, 41]. It has also been suggested that time from deafferentation to the onset of symptoms may provide insight about potential mechanisms underlying these effects. Rapid onset of GET suggests unmasking of silent synapses, whereas longer delays may be consistent with sprouting, changes in strength of existing neural connections, ephaptic interactions, or a combination of processes [41–43].

Initial descriptions documenting the phenomenology of GET were brief case reports [44–46]. In-depth series of cases were subsequently reported [40], and these observations were followed by detailed methods for quantifying the visual-spatial coordinates and psychoacoustic properties of this phenomenon with contemporary psychophysical methods [47, 48]. Gaze-evoked tinnitus is distinguished from gaze-modulated tinnitus in which there is typically an area of visual space where the tinnitus is absent; tinnitus only becomes manifest after change in eye position exceeds certain spatial coordinates of gaze. In contrast, gaze-modulated tinnitus occurs when existing constant tinnitus is altered in some way (change in loudness or pitch) by change in eye position. However, there are terminological disparities in the literature that tend to obfuscate our understanding. For example, in their sample of individuals with gaze-modulated tinnitus, Coad and colleagues (2001) [49] suggest that “eye movement” is the relevant parameter. This description is in contrast to a report from the same laboratory whereby activations were maintained by sustained lateral gaze [42]. Giraud et al. (1999) [48] also use the term “eye movement” to describe their effects. We suspect that the term “eye movement” is a misnomer, but this specific use of terminology will require clarification.

With greater recognition of this phenomenon, several groups have tried to estimate the prevalence of gaze-modulated tinnitus based on either retrospective or prospective convenience samples [43, 49, 50]. Prevalence estimates range from 1.8 to 32% [43, 49, 50]. However, the incidence of GET is unknown, but in all likelihood, it is not as commonly observed as the gaze-modulated form.

While there is no agreed upon treatment for gaze-evoked or gaze-modulated tinnitus, Sanchez and Pio (2007) [51] describe a case whereby daily eye movement
exercises had the effect of suppressing the underlying tinnitus perception. The mechanism(s)-of-action is/are unknown and theoretical accounts remain to be adequately explained. Nevertheless, this interesting observation is worthy of further investigation and replication.

**Cutaneous-Evoked Tinnitus**

Also observed following skull-base surgery for gross total excision of mass lesions (glomus jugulare tumor, vestibular schwannoma) were documented reports of cutaneous-evoked tinnitus [52, 53]. In one individual, the trigger zone for cutaneous-evoked tinnitus was a circumscribed area in the upper aspect of the hand by the wrist area. Activation of this area by stroking the skin produced a tonal tinnitus approximating 800 Hz. In another presentation, simultaneously touching the right index finger and thumb triggered an auditory noise-like sensation.

**Imaging Gaze-Evoked, Gaze-Modulated, and Cutaneous-Evoked Tinnitus with Functional (fMRI) and Positron Emission Tomography (PET)**

As individuals can either turn their tinnitus on and off and/or modulate the percept with change in eye gaze, the potential exists for localizing the source of the tinnitus-related activity through various forms of neuroimaging methodologies (e.g., fMRI or PET). In these investigations, individuals with GET serve as their own endogenous generators and act as their own controls. The first account of imaging GET was reported by Cacace and colleagues (1996) [54] using fMRI. This innovation was possible because it allowed for the on-off paradigm commonly used in fMRI studies to be applied [55]. By comparing the differences in activation between tinnitus on and tinnitus off conditions, tinnitus generator sites could in theory be localized. In one example where fMRI was successful in this regard, activations were observed in the upper brainstem and frontal cortex (superior colliculus and frontal-eye fields) [54, 56, 57]. This approach set the stage for other types of imaging studies on tinnitus and allowed other investigators to replicate and expand upon these original results (e.g., Giraud et al., 1999 and Lockwood et al., 2001) [42, 48].

With respect to cutaneous-evoked tinnitus, using fMRI and a finger tapping opposition task, Cacace and colleagues (1999) [53] were able to validate that finger tapping in one hand activated specific auditory pathways. When the finger tapping opposition task was performed with the right hand, which served to trigger the tinnitus, activation of the left temporal cortex (i.e., the superior portion of the Sylvian fissure and inferior aspect of the parietal operculum) was observed contralateral to the motor activation task. This cortical activation was in addition to the expected motor area activation sites in or near the Rolandic sulcus. Importantly, the finger tapping opposition task with the opposite hand only activated sites associated with the motor system, thus documenting the specificity of this phenomenon by using an objective test.

The occurrence of GET and cutaneous-evoked tinnitus expand our perspective on the biological basis of tinnitus by considering these phenomena in the context of lesion-induced cross-modal plasticity. Obviously, these conditions are sufficiently different from other forms of tinnitus, and these manifestations require an expansion of the existing models and frameworks to account for these phenomena.

**Conclusions**

The detection, management, and treatment of vestibular schwannoma has evolved over a period of time, whereby current treatment options have reduced mortality to near zero and minimized substantially the surgical morbidity. This current state-of-affairs is due to technical innovations in imaging and electrophysiology allowing for early diagnosis, improvements in surgical technique, use of the operating microscope, and availability of alternative treatment options. Tinnitus and vestibular schwannoma are intimately related. Unilateral tinnitus can serve as a red flag to signal the presence of this disease, prompting further evaluation and ultimately resulting in early diagnosis and better surgical outcomes. Tinnitus outcomes vary with treatment type. Following microsurgery, it is unpredictable if tinnitus will get better or worse. However, with stereotactic radiosurgery, tinnitus is generally unchanged following treatment. Lastly, in cases where hearing is lost completely and
abruptly during surgery, gaze-evoked, gaze-modulated, or other forms of somatic (cutaneous-evoked) tinnitus can result. These later types of tinnitus, which may be a consequence of cross-modal plasticity, are not accounted for by available models of tinnitus generation and require special consideration in future theories. Nevertheless, they serve to expand the biologic basis of tinnitus and provide additional insight to the complexity of this phenomenon under various conditions and circumstances [58].

References

Chapter 40
Microvascular Compression of the Vestibulocochlear Nerve

Dirk De Ridder and Aage R. Møller

Keypoints

1. Microvascular contacts or compressions of the vestibulocochlear nerve can result in tinnitus.
2. For nonpulsatile tinnitus, the contact is most often at the central nervous system segment.
3. For pulsatile tinnitus and typewriter tinnitus, the contact is at the peripheral nervous system segment. The tinnitus is unilateral and characterized by intermittent paroxysms of tinnitus.
   (a) A typical development consists of progressively more frequent bouts of tinnitus, which last longer and longer.
   (b) If bilateral vascular compressions exist, the tinnitus alternates between the left and right side, and does not occur on each side simultaneously.
4. Associated symptoms are correlated with related contacts/compressions of nearby nerves and include overt or cryptogenic hemifacial spasms, geniculate neuralgia, optokinetically induced short bouts of disabling positional vertigo, and tinnitus frequency-specific hearing loss.
5. Auditory brainstem responses (ABRs) correlate with disease progress and clinical symptoms and can be used diagnostically.
   (a) Tinnitus is causally related to a decrease in amplitude of peak II in the ipsilaterally elicited ABR.
   (b) Tinnitus frequency-specific hearing loss is causally related to prolongation of the ipsilateral interpeak latency (IPL) I–III.
   (c) Prolongation of contralateral IPL III–V occurs and is a sign of slowed signal transmission in the brainstem.
6. Magnetic resonance imaging sequences with constructive interference in steady state can visualize most vascular contacts/compressions of the auditory nerve.
7. Microvascular decompression should be performed before irreversible nerve damage is induced; clinically, the procedure should be performed before 4–5 years.

Keywords Pulsatile • Tinnitus • Vascular conflict • Microvascular compression • MVC • MVD

Abbreviations

AAO American Academy of Otolaryngology
AAOO American Academy of Ophthalmology and Otolaryngology
ABR Auditory brainstem response
CISS Constructive interference in steady state
CVCS Cochleovestibular compression syndrome
ENT Ear nose and throat
HFS Hemifacial spasm
IPL Interpeak latency
MRI Magnetic resonance imaging
MVC Microvascular compression
MVD Microvascular decompression
PNS Peripheral nervous system
TGN Trigeminal neuralgia
Introduction

Definition of Microvascular Compression

A blood vessel compressing a cranial nerve induces a nerve stimulation leading to a hyperactive cranial nerve syndrome [1, 2] with or without a loss of function. It is diagnosed almost solely based on the history taken, and a magnetic resonance imaging (MRI) is used for exclusion of other pathology and as a possible confirmation.

Primary and Secondary Microvascular Compressions

Microvascular compression (MVC) can occur as such or can be induced based on a general lack of space in the posterior fossa, such as seen in the Arnold–Chiari malformation [3–5] or associated with space-occupying lesions. This can result in a direct compression [6, 7] or indirect compression [8], but can also occur contralaterally, possibly due to a decrease in intracranial space [9–13]. In Sindou’s series [4] of 39 patients with Arnold–Chiari malformation,1 nine suffered from trigeminal neuralgia. After decompressing the foramen magnum, five of these nine individuals got rid of their pain, whereas the remaining four persons required a second microvascular decompression (MVD) operation. In this series of trigeminal neuralgias treated by MVD, the nerve was compressed between the pons and petrous bone in 3.9% of persons studied, due to the small size of the posterior fossa [14]. Removal of a tentorial meningioma can improve sudden hearing loss related to an MVC of the vestibulocochlear nerve based on the same premises [8].

Signs and Symptoms of Microvascular Compression

Examples of MVC syndromes are trigeminal neuralgia, glossopharyngeal neuralgia, hemifacial spasm [HFS], disabling positional vertigo, tinnitus, and otalgia.2 Other clinical syndromes such as spasmodic torticollis [15], cyclic oculomotor spasm with paresis [16], superior oblique myokymia [17, 18], and abducens spasm [19] may also be initiated by vascular compressions of the respective cranial nerves (nerves intermedius, spinal accessory nerve, and oculomotor and trochlear nerves). The incidence of the different MVC syndromes seems to be related to the length of the central nervous system (CNS) segment [20].

MVC of cranial nerves usually occurs unilaterally and, thus, induces unilateral symptoms [21–25] characterized by paroxysmal and intermittent spells of hyperactivity. The paroxysms typically become more frequent over time, the intermittent symptom-free periods become shorter and terminate in a constant dysfunction [26–28]. The symptoms of MVC can often be evoked by specific triggers [21, 26–28]. MVC syndromes are most common in late middle age (mean age 50 years) [21–25].

MVC of the vestibulocochlear nerve can cause any of the following paroxysmal symptoms depending on the place of compression: vertigo disabling positional vertigo [29], tinnitus [30, 31], hearing loss [32], or ear pressure (Dirk De Ridder unpublished observation). MVC rarely presents bilaterally (1–12%) [23, 25, 33, 34]; if it does, the pain or spasm alternates sides and never occurs on both at the same time. There usually is a delay between the onset of symptoms from one side and the development of symptoms of the other side [24, 35, 36], with only 2–3% of the bilateral cases starting simultaneously. Bilateral MVC has a higher incidence in familial cases [33, 35].

MVC syndromes that affect more than one cranial nerve occur rarely (incidence 2.8%) [24]. The combination may occur unilaterally (1.5%) or bilaterally (1.3%). The mean age is higher than for unilateral symptoms, 63.2 vs. 55.3 years, which is similar to bilateral MVCs (61.4 years) [24]. If one blood vessel contacts two or more cranial nerves, symptoms do not develop at the same moment in time [37]. The best-known double compression syndrome is called the tic convulsif, consisting of a combination of HFS and trigeminal neuralgia [38], which can occur even bilaterally [37, 39].

These data suggest that if bilateral tinnitus is due to MVC it is expected that the left- and the right-sided component should start at different moments in time and with a different pitch. Theoretically, true bilateral tinnitus (i.e., with same pitch) could occur if the compression is at the level of the cochlear nucleus.

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1 Arnold-Chiari malformation: displacement of the medulla and cerebellar tonsils and vermis through the foramen magnum into the upper spinal canal; often associated with other cerebral anomalies.

2 Otalgia: Earache.
Microvascular Compression of the Vestibulocochlear Nerve

Cochleovestibular Compression Syndrome

A recent meta-analysis has confirmed that blood vessels in contact with the vestibulocochlear nerve can result in otological symptoms, including hearing loss and tinnitus [40, 41].

Whereas initially it was proposed that only vascular compression of the root entry zone of a cranial nerve could cause symptoms [42], it was later suggested that any vascular contact along the CNS segment (between the internal acoustic meatus and the brainstem) could result in tinnitus [20]. Vascular loops inside the internal acoustic meatus along the morphologically more resistant peripheral nervous system (PNS) segment, however, were described to produce either typewriter tinnitus [43] or pulsatile tinnitus [41, 44]. Typewriter tinnitus consists of paroxysms of tinnitus perceived as Morse code, machine gun-like staccato, or typewriter sound and has been shown to be responsive to treatment with carbamazepine [43], thus analogous to trigeminal neuralgia.

Diagnostic Criteria of Cochleovestibular Compression Syndrome

Based on the analogy with other vascular compression syndromes, tinnitus caused by MVC would be expected to be unilateral and have short-lasting paroxysms with the tinnitus-free intervals becoming progressively shorter – ending in constant tinnitus. This kind of tinnitus would be expected to occur in middle-aged individuals and would not be anticipated to be associated with a flat hearing loss, as the typical MVC disorders (HFS and TGN) are not associated with complete weakness or complete loss of sensation in the entire distribution of the cranial nerve. Persistent compression can result in changes in the characteristics of pain and sensory impairment [26]. In a similar fashion, long-standing HFS could result in facial palsy or Bell’s palsy [45]. Similarly, chronic vestibular nerve compression can lead to hypofunctioning of the labyrinth, clinically expressed as gait instability [25].

Both parts of the vestibulocochlear nerve might be compressed at the same time, and symptoms from the vestibular nerve would be expected in individuals with tinnitus from MVC. A similar evolution is noted, with progressively more vertiginous spells and shorter symptom-free periods [25, 28]. In contrast to Ménière’s disease, the spells are shorter lasting and have no aura and no postictal period. In a chronic stage, persistent instability is noted [25, 46].

It is of interest, however, that in Ryu’s study, 73% of the patients with a MVC were diagnosed as having Ménière’s disease [28]. The main electrophysiological difference between Ménière’s disease and cochleovestibular compression syndrome (CVCS) is that in Ménière’s disease there are no abnormalities in peak II and interpeak latency (IPL) I–III of the auditory brainstem response (ABR) [47]. Two more nerves are in close relationship with the cochleovestibular nerve: the intermediate and the facial nerve. Vascular contact with the nervus intermedius is associated with geniculate neuralgia [22]. At the acute stage, intermittent paroxysmal bouts of otalgia occur; at a later stage, a deep, dull hemifacial pain develops [22].

Vascular contact with the root exit zone of the facial nerve can result in HFS [48, 49] and concomitant contact with the cochleovestibular nerve. The same vessel can cause auditory signs including low frequency tinnitus and hearing loss [31, 50].

Characteristic Features of Tinnitus as a MVC Syndrome

Selection Criteria

1. Intermittent paroxysmal spells of tinnitus lasting only seconds
   (a) Hearing loss at the tinnitus frequency
2. Associated ipsilateral symptoms from adjacent cranial nerves
   (a) Cryptogenic or overt HFSs
   (b) Bouts of otalgia or feeling pressure in the ear
   (c) Vertiginous spells: short lasting, optokinetically induced
3. Positive MRI for vascular compression
4. Positive brainstem auditory evoked potential using Møller’s criteria

Classification of Cochleovestibular Compression Syndrome

The characteristics of CVCS can be classified into four different groups based on the American Academy of
Ophthalmology and Otolaryngology’s (AAOO) (later renamed the American Academy of Otolaryngology [AAO]) criteria of Ménière’s disease [51, 52], relating to the certainty of the diagnosis of CVCS as the cause of tinnitus [53]:

- Possible CVCS: initially intermittent unilateral tinnitus spells without associated symptoms.
- Probable CVCS: possible CVCS with associated symptoms (vertigo spells; ipsilateral cryptogenic or overt HFS; ipsilateral pressure feeling in the ear, ipsilateral ear pain, or deep, dull hemifacial pain; ipsilateral frequency-specific hearing loss).
- Definite CVCS: probable CVCS with abnormal ABR and/or abnormal MRI.
- Certain CVCS: definite CVCS is surgically proven.

Pathophysiology of the CVCS

A cranial nerve has two parts, a CNS segment and a PNS segment separated by a transition zone, known as the root entry or root exit zone (for sensory and motor nerves, respectively) or Obersteiner–Redlich zone. The length of the CNS segment is different in every cranial nerve, with sensory fibers, in general, having a longer CNS segment than motor fibers [54]. For the VIIIth cranial nerve, the CNS segment encompasses the entire cisternal trajectory of the cochleovestibular nerve with the root entry zone located at the entrance of the internal auditory canal, thus the root entry zone is located at the internal auditory meatus.

Functional Anatomy

The cochlear nerve contains approximately 30,000 axons [55], 90% of which are myelinated (type I) and 10% of which are unmyelinated (type II) [56]. (For details, see Chaps. 8 and 36.) Myelinated nerve fibers represent the afferent neurons from the inner hair cells and the efferent neurons to the outer hair cells. Unmyelinated nerve fibers, on the contrary, represent the efferent neurons to the inner hair cells and the afferent neurons from the outer hair cells [56].

The average axon diameter of the PNS segment is fairly constant at ±3 μm [56] or 4.2–5.5 μm [57], suggesting conduction velocities of approximately 12 m/s [58] (11.6 ± 1.6 m/s). Whether differences exist in fiber spectrum, especially with regards to fiber diameter between apical and basal fibers in humans, is still debated, so it is not known whether a direct correlation exists between axonal diameters and tonotopy in humans; however, it has been suggested [57].

The auditory system is tonotopically organized. This means neurons sensitive to specific acoustic frequencies are topographically arranged in an orderly manner [59–62]. As the cochlea is tonotopically organized (Von Bekesy’s place theory of pitch perception) – as well as the cochlear nuclei, the inferior colliculus, and the auditory cortex – the cochlear nerve has to be tonotopically organized too [31], as shown in animal studies [63]. The cochlear nerve (as other cranial nerves) rotates as it travels through the auditory canal and cisternal segment of the subarachnoidal space toward the cochlear nucleus [64]. The tonotopy follows this rotation as well. This tonotopy has been demonstrated in humans as well as in studies of MVDs of the vestibulocochlear nerve [31]. It has also been demonstrated by means of an MRI technique using 3D reconstructions of high-resolution (0.6 mm slice thickness), heavily T2-weighted images (constructive interference in steady state, CISS) [65] also known as virtual endoscopy [66].

Pathophysiological Model of CVCS

Several hypotheses have addressed the pathology of MVC in general. Some of them concern the cranial nerve and some concern the respective nucleus. HFS has been studied extensively, and evidence for hyperactivity in the facial motonucleus has been presented [48]. There is no evidence supporting the old hypothesis that blood vessels elongate and their brain “sags” with age [2, 67–70]. It is not known whether the formation of vascular loops in the posterior fossa that can come close to cranial nerve increases with age [69]. MVC has been claimed to cause focal demyelination (see Chap. 84), but little evidence of demyelination or other morphological changes in cranial nerves in individuals with symptoms of cranial nerve vascular compression has been published.
Focal demyelination, if it exists because of MVC, could cause ectopic excitation [68, 71–73] (see Chap. 84). Such ectopic excitation might cause dysfunction of the cochlear nerve, most likely leading to a reorganization of the auditory nuclei in the auditory brainstem through activation of neural plasticity. Subsequently, the entire auditory tract, including the auditory cortex, can become hyperactive, resulting in gamma band activity, which may cause tinnitus [74, 75].

**Microvascular Compressions Can Result in Tinnitus due to Abnormal Signal Transmission**

Animal (cat) studies have described the tonotopic organization of the auditory nerve. The tonotopic organization of the human auditory nerve [31] has been related to the site of vascular contact and the frequency-specific dysfunction of the cochlear nerve revealed as the frequency-specific hearing loss and a frequency-specific tinnitus [31, 41].

Nonfrequency-specific click evoked auditory brainstem potentials are used routinely in an attempt to discover early demyelination. If the close contact with a blood vessel causes demyelination, frequency-specific ABR would be expected to detect such focal demyelination. (For details about the anatomy of the auditory nerve, see Chap. 36 and [76].)

The neural generators of the auditory evoked responses (ABRs) in humans have been determined [59, 76]. The generators of the ABR in humans are not the same as the generators of the ABR in animals, including those in monkeys [76].

Peak I in humans is generated in the distal part of the cochlear nerve; peak II is generated in its CNS segment; peak III in the cochlear nuclei; peak IV in the superior olivary complex; peak V in the lateral lemniscus; and peak VI in the inferior colliculus (see Table 40.1) [76].

The IPL I–III would therefore be expected to be increased. If the vascular compression occurs at the CNS segment [20], peak II would be expected to be affected. Evoked potentials, in general, are the result of synchronized firing pattern as a reaction to a sensory stimulus [76]. The more synchronized the nerves fire, the higher the summed amplitude will be. If MVC of the cochlear nerve creates functional impairment of some fibers, the temporal coherence of firing will decrease, resulting in a decrease of the amplitude of peak II. This hypothesis is supported by clinical findings that show a peak II decrease in individuals with tinnitus ipsilateral to MVCs with recurrence of peak II when surgical decompression is successful [53]. This suggests that the tinnitus is causally related to dysfunctional signal transmission at the site of compression in the initial stage of compression.

1. Chronic MVC results in frequency-specific hearing loss at tinnitus frequency.

In the first 2 years, no significant changes in ABR are noted in patients presenting with tinnitus and MVC [53]. Once peak II decreases are noted, IPL I–III prolongs [53]. The fact that the IPL I–III prolongation is related to the duration of the tinnitus furthermore suggests that this is a dynamically progressive pathology [53] and that the effect of vascular contact with blood vessels creates changes over time, both electrophysiologically [53] and clinically [53].

The IPL I–III prolongation seems to be significantly related statistically to the degree of tinnitus after normalization for age [53]. Postoperatively, a shortening of the IPL I–III is not related to a clinical improvement in tinnitus but to an improvement in tinnitus frequency-specific hearing loss [53].

Schwaber and Hall [46] analyzed auditory brainstem evoked potentials in cochleovestibular compressions: IPL I–III interval difference ≥0.2 ms occurs in 66% of patients with a diagnosis of an MVC syndrome. Wave II amplitude <33% (in comparison with the contralateral) occurs in 57%. Contralateral IPL III–V interval difference ≥0.2 ms occurs in 30%; the ipsilateral IPL I–III absolute interval ≥2.3 ms occurs in 24%. Contralateral IPL III–V absolute interval ≥2.2 ms occurs in 2% of patients diagnosed with an MVC syndrome. This is associated with hearing loss for high frequencies in 65% of patients, a mid-frequency hearing loss in 27% of patients, and a low frequency loss in 8% of patients. A flat hearing loss was not seen in patients diagnosed with a MVC in Schwaber’s series [46].

While the ABR changes (increased IPL I–III) indicate that the conduction velocity in the auditory nerve has decreased, intracranial recordings from patients undergoing MVD operations for severe tinnitus [77] did not find any significantly increased latencies when compared with individuals with some hearing loss who did not have tinnitus, confirming that IPL I–III is related
to hearing loss and not tinnitus, per se. When compensated for hearing loss, individuals with tinnitus do not have significant changes in auditory evoked potentials from the peripheral part (IPL I–III) of the ascending pathways but a slight change in the potentials recorded from the inferior colliculus.

Signals transmitted via the compressed nerve fibers arrive at the cochlear nuclei in delay (IPL I–III prolongs) in comparison with the contralateral input. Because auditory input arrives bilaterally, this slowing down of nerve conduction in the auditory nerve of the affected ear (ipsilateral IPL I–III) will be counterbalanced by slowing down the auditory signals coming from the contralateral ear (De Ridder, submitted). As this slowing down can only occur in the brainstem, this will result in an increase in IPL III–V in the contralateral side. As such, a pathophysiological explanation can be proposed for Møller’s criteria of MVC syndromes of the cochleovestibular nerve.

Criteria of microvascular compression of the VIIIth nerve [29]:

1. Ipsilateral IPL I–III ≥2.3 ms
2. Contralateral IPL III–V ≥2.2 ms
3. IPL I–III difference ≥0.2 ms
4. IPL III–V difference ≥0.2 ms
5. IPL I–III difference ≥0.16 ms if low or absent peak II
6. IPL III–V difference ≥0.16 ms if low or absent peak II
7. Peak II amplitude <33%

Whereas initially tinnitus is causally related to abnormal signal transmission in the peripheral part of the cochlear nerve at the site of the compression, electrophysiologically demonstrated by peak II decrease ipsilateral to the tinnitus side, chronic tinnitus might be the result of deafferentation due to hearing loss caused by slowing down of signal transmission in the peripheral part of the cochlear nerve, electrophysiologically related to IPL I–III prolongation. It is known that the most common cause for tinnitus is auditory deprivation, inducing the development of an auditory phantom percept [78]. Therefore, it is likely that when the compression has resulted in a hearing loss this will result in tinnitus, specifically at the frequency of hearing loss [31, 53, 79–81]. It has also been shown that the neural network in the brain that generates tinnitus changes with time [82], with a marked change before and after 4 years of tinnitus duration. This could explain why tinnitus that has lasted a long time is more difficult to treat by surgical decompression than acute tinnitus [28, 30, 31, 83, 84]. MVD is less successful in treatment of tinnitus that has lasted for longer than 3–5 years than tinnitus that has lasted a shorter period [31], coinciding temporally with the tinnitus-related brain network changes.

### Conclusion

It is evident from several studies that MVD operations are more successful in treating tinnitus that has not lasted too long (less than 3–5 years). Studies have shown cure rates of 30% of patients and 30% improved. Worsening of tinnitus caused by MVD operations and other complications are rare but can be severe and life threatening.

After a MVD operation, the hearing threshold of the frequency of the tinnitus may improve if IPL I–III normalizes and peak II reoccurs.
The following pathophysiological mechanism can be suggested for tinnitus: when a blood vessel comes into contact with the auditory part of the VIIIth nerve and starts interfering with normal signal transmission, initially no electrophysiological changes can be retrieved. After 2 years, when enough fibers are involved a decrease in peak II on the ABR develops. When the close contact with a blood vessel continues, IPL I–III may increase, associated with hearing loss at the tinnitus frequency. This signal transmission slowing at the side of the compression is compensated by a contralateral slowing in the brainstem (contralateral IPL III–V prolongs). When hearing loss develops, tinnitus might relate more to the deafferentation, which induces network changes in the brain based on neural plasticity, and tinnitus at that stage has become a phantom percept. These tinnitus network changes alter in time, which might explain why surgical decompression has to be performed before 4 years in order to be successful.

References

the most reliable diagnostic signs? Acta Neurochir (Wien) 140:1279–86.


Chapter 41
Causes of Tinnitus: Cerebrovascular Diseases

Miguel J.A. Láinez, Alejandro Ponz, and Anna Piera

Keypoints

1. Tinnitus can be divided into two broad groups: objective and subjective tinnitus.
2. Several layers of complexity are involved in the pathophysiology and the cause of tinnitus, and it is rarely known what causes an individual’s tinnitus.
3. Disorders that affect the brain are often accompanied by tinnitus.
4. Cerebrovascular diseases can be the cause of both objective and subjective tinnitus.
5. This chapter discusses cerebrovascular diseases as a cause of tinnitus and how it is produced.

Keywords Tinnitus • Cerebrovascular diseases • Arterial pulsatile tinnitus • Venous pulsatile tinnitus

Abbreviations

CTA Angiotomography
CVD Cerebrovascular diseases
DAVF Dural arteriovenous fistula
GJT Glomus jugular tumor
HJB High jugular bulb
MRA Magnetic resonance angiography
PT Pulsatile tinnitus
RI Resistive index

Introduction

Tinnitus can be divided into two broad groups: objective and subjective tinnitus. Objective tinnitus is caused by sound generated in the body reaching the ear through conduction in body tissues [1]. The source can be turbulent flow of blood in an artery where there is a constriction, or it can be caused by muscle contractions. Unlike subjective tinnitus, an observer using a stethoscope or a person listening to the individual at a close distance may hear the sound. Subjective tinnitus is meaningless sounds that are not associated with a physical sound and only the person who has the tinnitus can hear it.

Several layers of complexity are involved in the pathophysiology and the cause of tinnitus, and it is rarely known what causes an individual’s tinnitus (idiopathic tinnitus). Disorders of the central nervous system (CNS) or disorders that affect the function of the CNS are often accompanied by tinnitus. In the large group of cerebrovascular diseases, some cause tinnitus as an isolated symptom, but tinnitus is often associated with other symptoms. Such diseases can cause both objective tinnitus (e.g., pulsatile tinnitus in carotid-cavernous fistula) and subjective tinnitus such as those from ischemia of the inferior colliculus that can activate subcortical auditory pathways, and thereby cause tinnitus. This chapter discusses cerebrovascular diseases as a cause of tinnitus as well as the mechanisms, which cause the tinnitus. We will distinguish between pulsatile tinnitus and non-pulsatile tinnitus.

Pulsatile Tinnitus

Pulsatile tinnitus is perceived by an individual as pulsations in the tinnitus that are synchronous with the heart, and it is similar to pulsating sounds or a rushing
sound (see chapter 59). Pulsatile tinnitus can be subjective or objective. Objective tinnitus can result from blood flow through a constriction causing the flow to become turbulent. Objective tinnitus that is strongly associated with the timing of the heart beat is most likely caused by turbulent flow in arteries or veins of the head or neck area located adjacent to the ear, on the surface of the head, or just inside the head. Patients with such problems require special imaging studies and often require surgery to resolve the issues.

This type of tinnitus can be heard as several characteristic sounds including a lower pitched thumping or booming. Objective tinnitus can also be caused by respiration and heard as a blowing sound, which is coincidental with respiration. Tinnitus that sounds like clicking, rhythmic sounds can be caused by muscle contractions in the head, such as those from muscles in the palate or the middle ear muscles.

Patients with pulsatile tinnitus may need a thorough medical evaluation to locate the cause of their tinnitus. Many published studies describe methods for treating objective tinnitus [2, 3]. Vascular imaging techniques have been employed to help determine the site of lesion, but there are many forms of pulsatile tinnitus that have no known cause (idiopathic category). This can be caused by failure to properly interpret imaging studies or miss the trouble spots that may be tangled in other structures or hidden by bone or other tissue. A clinical interview is crucial to identify this type of tinnitus [4–7].

**Arterial Pulsatile Tinnitus: Differential Diagnosis**

Many different pathologies of the cerebrovascular system have been reported as cause of pulsatile tinnitus. Some are listed below.

**Cervical Arterial Stenosis**

Stenosis of the carotid or the subclavian arteries are typical causes of pulsatile tinnitus, ipsilateral, or contralateral to the side of tinnitus. Often, the intensity or the appearance of tinnitus is not related to the degree of stenosis. Doppler ultrasonography is a useful test for use in tinnitus clinics to distinguish between these causes of pulsatile tinnitus [8, 9]. When stenosis of the carotid artery is symptomatic, endarterectomy also relieves tinnitus [10–12].

**Aberrant Internal Carotid Artery and Other Morphologic Abnormalities**

Internal carotid artery morphologic abnormalities that can present with pulsatile tinnitus are mainly tortuosity (having many turns and twists) and coiling of the artery. Head bruit causing objective tinnitus can be evaluated by angiotomography or magnetic resonance angiography (MRA) of the head and neck, which can differentiate these abnormalities from other more serious vascular disorders [13, 14].

**Cervicocephalic Arterial Dissection**

Pulsatile tinnitus rarely occurs together with cervicocephalic arterial dissection. Tinnitus may occur together with ischemia caused by arterial dissection in carotid stenosis, causing Horner’s syndrome. Dissection of the vertebral artery may cause vertigo and dysgeusia. Angiography is essential if there is a high degree of suspicion of such pathologies, and delay in the diagnosis should be avoided [15].

**Fibromuscular Dysplasia of Cervical Arteries**

Pulsatile tinnitus often occurs together with stenosis of the carotid artery. Fibromuscular dysplasia of the vertebral artery can also cause tinnitus [16]. Symptoms such as tinnitus, vertigo, headache, and cervicofacial hypoesthesia might lead a person to seek medical help from a neuro-otologist. Fibromuscular dysplasia may cause pulsatile tinnitus directly because of its stenosing angiopathy and indirectly by activation of sympathetic nervous system through its effect on a sympathetic plexus.

**Dural Arteriovenous Fistulas (DAVF)**

Pulsatile tinnitus is a common symptom in individuals with intracranial dural arteriovenous fistulas [17]. The occurrence of pulsatile tinnitus is related to the location of the fistula and the location of the arteries feeding the fistula. Yeh et al. have published an interesting study in which they compared the characteristics of
DAVF and carotid duplex sonography [18]. They showed that the occurrence of pulsatile tinnitus is highly correlated with the location of the DAVF and its feeding arteries. They also showed that the resistive index¹ and the end diastolic velocity in the external carotid artery are related to the presence of DAVF in pulsatile tinnitus patients by sonography [18]. Although this technique is not considered to provide a definitive diagnosis, sonography may assess pulsatile tinnitus in patients who are candidates for angiography [19].

**Carotid-Cavernous Fistula**

Carotid-cavernous fistula is a rare vascular abnormality that may develop following traumatic injury to the skull base; it may also be spontaneous. Objective tinnitus caused by this pathology is of acute or subacute onset, and an early intervention, endovascular or surgical, is needed to prevent permanent disability. Pulsatile tinnitus occurs together with carotid-cavernous fistula — other symptoms being papillary abnormalities, proptosis, headaches, and papilledema [20–24].

**Aneurysms**

Petrus carotid aneurysms and other located aneurysms are serious causes of tinnitus. Pulsatile tinnitus may be a symptom of the compression by the aneurysm when located near auditory structures. Hemorrhage, secondary to an aneurysm, can produce tinnitus as an acute symptom in addition to other symptoms derived from the subarachnoid hemorrhage [25, 26].

**Vertebrobasilar and Carotid Dolichoectasia**

Dolichoectasia² is an angiopathy characterized by dilatation, elongation, and tortuosity of brain arteries. It most frequently involves the vertebral and basilar arteries, but involvement of both the vertebrobasilar and carotid systems is rare. A magnetic resonance angiography (MRA) imaging or a computed tomographic angiography (CTA) can show an enlarged tortuosity of these arteries, often producing compression of the cranial portion of vestibulocochlear nerve [27].

**Persistent Trigeminal Artery**

Persistent trigeminal arteries are rare and represent a remnant of the fetal carotid-basilar circulation. They typically extend from the internal carotid artery to the basilar artery. In rare instances, a persistent trigeminal artery is associated with a carotid-cavernous fistula; patients with this condition may have pulsatile tinnitus in addition to other symptoms such as proptosis, eye pain, conjunctival injection, diplopia, and decreased visual acuity [28].

**Subclavian Steal Syndrome**

This syndrome is characterized by a subclavian artery stenosis located proximal to the origin of the vertebral artery. In this case, the subclavian artery steals reverse flow of blood from the vertebrobasilar artery circulation to supply the arm during exertion, resulting in vertebrobasilar insufficiency. As the vertebrobasilar arterial system feeds both the peripheral and central auditory and vestibular systems, symptoms such as dizziness, recurrent vertigo, hearing loss, and tinnitus [29] may occur in the subclavian steal syndrome.

**Internal Auditory Canal Vascular Loops**

A vascular loop entering the internal auditory meatus can be another cause of pulsatile tinnitus. Normally, the wall of the internal auditory meatus prevents vibrations of an artery from reaching the cochlea, but structural differences between the internal acoustic meatus and pericarotid area can originate tinnitus. De Ridder et al. insulated the carotid artery preventing arterial pulsations be transmitted to the bone. Abnormalities in the surgical interpositioning of Teflon felt between the arterial loop and the cochlea can eliminate this form of tinnitus [30].

**Ischemic and Hemorrhagic Infarctions of the Posterior Circulation**

These kinds of infarcts may cause tinnitus, but it is unknown what exactly causes the tinnitus [31–34].

¹Doppler resistive index (RI) is the peak systolic velocity – the end diastolic velocity divided by the peak systolic velocity.

²Dolichoectasia: The term “dolichoectasia” means elongation and distension. It is used to characterize arteries throughout the human body that have shown significant deterioration of their tunica intima (and occasionally the tunica media), weakening the vessel walls and causing the artery to elongate and distend.
Brainstem Telangiectasias

Capillary telangiectasia (dilation of small or terminal vessels) is often found incidentally on magnetic resonance imaging because it is normally associated with only minor neurologic symptoms. There has been little evidence about whether such lesions are responsible for symptoms at all. In some individuals, telangiectasia is associated with tinnitus and sensorineural hearing loss. The auditory brain stem responses (ABR) in such individuals have abnormalities regarding waves III and IV. Another sign is asymmetry in optokinetic nystagmus that is present in some individuals [35].

Proatlantal Intersegmental Artery

Primitive carotid-vertebral and carotid-basilar anastomoses are formed early during human embryogenesis at approximately 24 days. From cephalic to caudal direction, these anastomoses are cranial extensions of the primitive internal carotid, trigeminal, otic, hypoglossal, and proatlantal intersegmental arteries. The proatlantal intersegmental artery maintains the posterior circulation until the vertebral arteries are fully developed, between the 7th and 8 gestational weeks. Normal and abnormal morphofunctional aspects of prenatal and postnatal forms of the proatlantal intersegmental artery have been described. When the proatlantal intersegmental artery fails to obliterate, it can produce symptoms affecting the function of vertebrobasilar structures such as hearing, tinnitus, and dizziness. In some individuals, these arteries do not give noticeable symptoms and are only found incidentally [36].

Venous Pulsatile Tinnitus: Differential Diagnosis

Glomus Jugular Tumor

Glomus jugular tumors are benign and slow-growing lesions that can be locally aggressive because of their proximity to lower cranial nerves and major vascular structures [37]. These lesions are known causes of pulsatile tinnitus and other symptoms by their compression of the nerves of the skull base [38]. Surgical resection is often complicated; the possibilities of using radiosurgery are limited and combinations of both localized surgery in the middle ear and gamma knife surgery have shown good results.

High Jugular Bulb

The jugular bulb is normally surrounded by a bony layer in the jugular fossa. It is named a high jugular bulb (HJB) if it is anatomically above the inferior surface of the bony annulus, extending into the middle ear or located above the basal turn of the cochlea. HJB is a frequent cause of objective pulsatile tinnitus. It may be dehiscent or aberrant. Techniques using endovascular management and surgery using ligation and embolization have been described to relieve this abnormality [39].

Sigmoid or Jugular Diverticulum

Jugular bulb diverticulum is a rare condition. It has been reported that unilateral auditory symptoms may accompany this disorder, although some individuals are asymptomatic. Some individuals with this condition are referred to clinics of neurotology centers with symptoms of unilateral sensorineural hearing loss and tinnitus. Tomographic venography is a useful tool to diagnose the condition [40, 41].

Condylar Vein Abnormalities

One of the most important venous foramina of the human skull is the condylar canal. This structure is described as the most stable and permanent venous emissary, with a prevalence of nearly 100%. It has been reported that patients with dural arteriovenous fistula of the anterior condylar vein have symptoms related to an unusual venous drainage. Pulsatile tinnitus may be the symptom of alarm for abnormalities of the jugular venous system [42].

Venous Angioma of Posterior Fossa

Venous angioma can cause tinnitus by affecting the auditory pathway [43–45] and the structures of the inner ear, thus similar to brainstem telangiectasias and
other vascular malformations of the brainstem and other posterior fossa locations.

### Sinus Thrombosis

Dural and profound sinus thrombosis commonly presents with headaches and some neurological symptoms depending on the location of the thrombosis and the surrounding edema and infarct. Dural sinus thrombosis may cause tinnitus with headaches in some individuals, more common if the course of the symptoms is subacute. The complaint from sigmoid sinus thrombosis may be unilateral head pain and unilateral pulsatile tinnitus [46–49].

### Cerebrovascular Diseases with Subjective Tinnitus

Some cerebrovascular disorders can cause pulsatile tinnitus by affecting the auditory system at different levels. For example, a brain ischemic infarct located in the inferior colliculus can produce an acute or subacute tinnitus, and even chronic tinnitus as a sequela, by its effect on the auditory pathway.

### References


34. Lee H. Sudden deafness related to posterior circulation infarction in the territory of the nonanterior inferior cerebellar artery: frequency, origin, and vascular topographical pattern. Eur Neurol 2008;59(6):302–6; Epub 2008 Apr 11
Complications to Medical Treatment

Paolo Enrico and Ron Goodey

Keypoints

1. When medical treatment is blamed, tinnitus may be harder to treat.
2. Adverse consequences are better accepted and more easily managed if the patient had been well informed before treatment started and had acknowledged and accepted the risk.
3. Ear syringing, suctioning, instrumentation, local anaesthetic injection, grommet insertion, dental treatment, hyperbaric oxygen therapy, and ototoxic ear drops are all relatively minor procedures that may be blamed for tinnitus.
4. Major ear operations may cause hearing loss and tinnitus.
5. Ototoxic drugs can cause hearing loss and tinnitus after administration systemically, intrathecally, or topically to extensive wounds or burns as well as from use as eardrops.
6. Onset of tinnitus is occasionally blamed on radiation therapy, noisy organ imaging, medical equipment accidents, neck manipulation, and general anesthetic.
7. Tinnitus can be triggered by procedures on any region of the body when there have been excessive pain and associated anxiety, fear, and anger.
8. The medical treatments most commonly accused of causing tinnitus are treatments with drugs. Usually, the tinnitus improves when the drug is withdrawn, provided there is no permanent damage to the cochlea or powerful associated factors.
9. Drugs with proven ototoxicity and that also cause tinnitus include aminoglycoside antibiotics, antineoplastic drugs, anti-inflammatory drugs, loop diuretics, antimalarials, and others. The ototoxicity may be synergistic with other agents that damage the inner ear.
10. Drugs that are not usually considered ototoxic but are sometimes blamed for causing tinnitus include lidocaine, anticonvulsants, antidepressants, cannabinoids, antihypertensives, beta-adrenergic blocking agents, opioids (buprenorphine), caffeine, and antihistamines. At times, drugs from within most of these groups are also credited with ameliorating tinnitus.

Keywords Tinnitus • Complication of treatment • Medical misadventure • Ototoxicity • Pathogenesis • Drug-induced tinnitus • Therapy-induced tinnitus

Introduction

Many differing medical treatments are thought by patients to have triggered the onset of their tinnitus [1]. Indeed, there are a variety of mechanisms and pathways by which this may occur. Medical treatment can result in reduced or abnormal stimulation through the auditory, somatosensory, vestibular, and other sensory pathways. Activity in central pathways can be affected directly. Unwanted effects of medical treatment may be temporary, but are associated with tinnitus that may persist once triggered. Medical treatment of almost any type throughout the entire body may be blamed as the trigger for the onset of tinnitus when that treatment has had powerful emotional associations and was accompanied by severe pain.

Tinnitus tends to be worse, and its management more difficult when the onset has been associated with fear or anger. Unfortunately, for the patient and therapist, when
the onset of tinnitus is perceived as being a complication of medical treatment, it is usually associated with anger and often with fear and anxiety as well. This can make management difficult. The main exception is when the possibility of tinnitus developing had been anticipated, clearly explained, and then accepted by the patient as an acceptable trade-off for life-saving treatment.

As clinicians, we may sometimes support a patient’s claim for compensation for tinnitus, which the patient attributes to medical treatment they had received. More often, however, many of us encourage our patients to disassociate their tinnitus from such emotionally charged triggers. We justify doing so on the basis that the association is unproven and that dwelling on it makes the tinnitus more intrusive and harder to manage. A review of the tinnitus literature shows that we seldom investigate a suspected relationship between the onset of tinnitus and a medical treatment, let alone report it.

This chapter is an opportunity to review not only the situations in which tinnitus is acknowledged as a complication of medical treatment but also situations that have been largely ignored in scientific literature as causes of tinnitus but which, in one author’s experience, occasionally are. The editors are to be congratulated for making it possible to consider all situations in which tinnitus may be a complication of medical treatment. Some of the sections in Part 1 of this chapter express unsubstantiated opinions acquired from Dr. Goodey’s otological practice and his discussions with colleagues. They are presented as a challenge to other colleagues for wider consideration. Part 2 of this chapter focuses entirely on drug therapy as a trigger for tinnitus. It discusses drugs with proven ototoxicity, and some of those that are sometimes accused of causing tinnitus but not considered ototoxic. Part 2 draws heavily on Dr Enrico’s extensive knowledge and experience as a neuropharmacologist.

**Part 1: Procedural Treatments that May Cause Tinnitus**

**Minor Procedures in and Around the Ear**

Often, procedures that clean the ear of wax and/or debris also reduce or eliminate any associated tinnitus. However, such procedures may occasionally trigger or aggravate tinnitus. Other procedures in the region may also trigger or aggravate tinnitus. Quite often (but not always), temporomandibular joint dysfunction may be aggravated by the same procedures and consequently aggravate the associated tinnitus.

**Ear Syringing**

Ear syringing is only occasionally mentioned in journal articles as a trigger for the onset of tinnitus [1–3]. However, it is frequently acknowledged as a trigger by patient support groups [4]. Even some of the more professional support groups find it necessary to produce brochures on the association [5, 6]. Mostly, they provide balanced and generally reassuring information. In such brochures, the triggering of tinnitus is sometimes attributed to ear syringing, but only when it is “poorly performed.” Many otologists who deal with patients troubled by tinnitus accept that some of these patients appropriately attribute the onset of their tinnitus to ear syringing.

Occasionally, syringing-induced tinnitus has been associated with rupture of the tympanic membrane (especially if it was already weakened). Rarely, there has been major trauma to the middle ear, and inner ear as well, especially if a carelessly attached nozzle came off with the pressure used. However, more commonly, any trauma attributable to syringing has been relatively minor and confined to the ear canal. The symptoms associated with the onset of tinnitus induced by syringing are pain and vertigo. Tinnitus is especially likely to have occurred and persisted if the doctor or nurse continued to syringe an ear after the patient had wanted them to stop.

Syringing should be avoided in those with a weakened or perforated eardrum (or a grommet) or with an infected ear canal. The water used must be at body temperature. The nozzle must be firmly attached; it should have a smooth and rounded tip; and it must be directed at the posterior canal wall. If pain or vertigo is induced, the procedure must be stopped immediately.

**Ear Suctioning**

Ear suctioning is often recommended as a safe alternative to syringing, and it usually is. It is the treatment of choice when there is a perforation or a grommet
Complications to Medical Treatment

However, noise levels at the suction tip are sometimes loud enough to be distressing to the patient and to trigger tinnitus [7–11], even when there is no measurable change in the audiogram.

Tinnitus is more likely to be triggered if the suction noise is excessively loud because of the material being aspirated. In this context, noise levels of 96 dB have been measured at the suction tip [12]. Tinnitus is more likely to be triggered if the commencement of the suction noise is abrupt and unexpected. If inner ear damage occurs, it may be a direct consequence of noise energy. Alternatively, inner ear damage could result from violent contraction of the stapedius muscle, as can be caused by a sound blast. However, Dr. Goodey is not aware of any patients in whom the annular ligament has been damaged and a perilymphatic fistula caused as a result of suctioning.

During suctioning, tinnitus and hyperacusis may occur and persist without any persisting change in hearing. In some of these, the situation may be identical with “acoustic shock disorder” described in comparable situations [13, 14]. Associated symptoms may include acute ear pain, muffled hearing, a feeling of fullness and numbness, and occasionally vertigo. Tinnitus and hyperacusis may persist when all the other symptoms have settled. In such situations, the inner ear may have been protected by the intermittent pattern and relatively short duration. A possible mechanism for the symptoms could be contraction of tensor tympani.

Suctioning of a mastoidectomy cavity or through a perforation often triggers vertigo. Occasionally, this is followed by persistent tinnitus, especially if suctioning was continued after the patient had become distressed.

A wise microscopist will always ask in advance whether the patient is intolerant to loud noise and always instruct their patient to tell the microscopist to stop if the suction noise is hurtful, causes vertigo, or is otherwise distressing.

Cleaning the Ear Canal Skin with Instruments

Cleaning of the ear canal with instruments often causes superficial ulceration and sometimes lacerations. Occasionally, a patient reports that it triggered their tinnitus. Ear canal injury or infection may also lead to chronic changes in the ear canal skin, which may then have a continuing effect on tinnitus.

Trauma Affecting the Middle Ear and/or Inner Ear

Clumsy instrumentation or failure to adjust to sudden head movement (such as during removal of a foreign body) can cause damage not only to the ear canal skin but also to the tympanic membrane, ossicular chain, and — through inadvertent manipulation of the chain — the inner ear. Tinnitus may result even without measurable hearing loss.

Injection of Local Anaesthetic

Injection of local anaesthetic into the ear canal in preparation for a minor surgical procedure occasionally triggers severe vertigo, which may last several hours and be extremely distressing for the patient. Accompanying tinnitus is insignificant because the vertigo is so distressing. Occasionally, tinnitus persists after nausea and vertigo have subsided. The development of effective topical anaesthetics has largely eliminated the need for injections of local anaesthetic into the ear canal for minor procedures [15].

Insertion of a Grommet

Quite commonly, insertion of a grommet to relieve Eustachian tube dysfunction or a middle ear effusion also reduces any associated tinnitus. Occasionally, however, insertion of a grommet may trigger or aggravate tinnitus, even when there has been no reaction to the local anaesthetic used and when the procedure has been gentle. In this situation, the tinnitus usually subsides or reverts to its previous level if the grommet is removed promptly, and the resulting hole was covered with a rice paper patch.

Dental Treatment

Case history questionnaires may include dental treatment as an item associated with the onset of tinnitus [16]. In Dr. Goodey’s experience, dental treatment can
be a potent trigger or aggravator of tinnitus. The tinnitus tends to be more severely affected on the side of the dental treatment and occurs more often if the procedure has been prolonged and painful and associated with anxiety. There is usually associated temporomandibular joint dysfunction and sometimes aggravation of chronic neck problems as well. However, dental treatment as a trigger for tinnitus receives little or no attention in the literature, whereas dental disorders as triggers for tinnitus do receive some attention [17–19]. Without associated factors, noise from dental drilling is seldom, if ever, loud enough and prolonged enough to cause hearing loss and tinnitus in patients. However, dentists and their assistants may occasionally suffer occupational noise-induced hearing loss and tinnitus after many years of exposure [20]. Malfunction of an air drill can cause a sudden and unexpected loud blast of noise and result in tinnitus and associated symptoms described as the acoustic shock disorder in the section “Ear suctioning” of this chapter.

Barotrauma

In the context of medical treatment, barotrauma is only likely to be blamed as the trigger for tinnitus when there has been difficulty in equalizing while hyperbaric oxygen was being used as an adjunct to therapy [21]. The incidence of barotrauma as a consequence of hyperbaric oxygen therapy has been assessed and correlated with conditions being treated [22, 23]. An associated incidence of tinnitus gets little mention. Occasional patients are adamant that their tinnitus occurred or became worse during hyperbaric oxygen treatment. If equalizing problems have occurred during a previous treatment session, or are anticipated, then a mini grommet will give complete protection during subsequent treatments. When treatment in a hyperbaric chamber is required following a diving accident, then any inner ear damage can usually be attributed to the original accident and not to the treatment.

Ototoxic Ear Drops

When the eardrum is perforated or has a grommet, there is potential for ototoxic components in ear drops to cause sensorineural hearing loss and trigger tinnitus. The incidence of this occurring has been very low considering the widespread use over a large number of years [24]. However, hearing loss and tinnitus from the use of such drops do occur. The risk is probably minimized if such drops are only used when the middle ear mucosa is inflamed. A modern clinician is unwise to allow such drugs to be used in high-risk ears or once the middle ear mucosa is healthy [25]. Fluoroquinolone antibiotic drops are now available, which are proven clinically and experimentally to be nonototoxic [26–28]. Unfortunately, they tend to be much more expensive and also less well tolerated, especially by children. Nevertheless, with expert panels in the US, Canada, United Kingdom, and Australia all advocating the use of fluoroquinolones, a clinician who continues to prescribe potentially ototoxic drops has to be prepared to justify the need for these types of medications.

Major Procedures in and Around the Ear

Stapedectomy, labyrinthectomy, tympanoplasty, simple myringoplasty (especially with an overlay graft, which involves more manipulation of the malleus), mastoid surgery, vestibular nerve section, and vestibular schwannoma surgery can all trigger tinnitus. However, these have all been dealt with in the section “Complications of surgical treatment”. Any resulting tinnitus is usually associated with additional sensorineural hearing loss.

As with the minor ear procedures, these more major operations only occasionally cause damage and tinnitus. More often, they reduce or relieve pre-existing hearing impairment and associated tinnitus or they have no effect on tinnitus.

Occasional Causes of Unexpected Tinnitus and Sometimes of Cochlear Hearing Loss

Radiation Therapy

Prior irradiation increases the incidence of ototoxicity, including tinnitus, during subsequent treatment with cytotoxic drugs [29]. Usually, the possibility of such life-saving treatment causing hearing loss and tinnitus will have been understood and accepted as a risk by patient. Occasionally, this is not the case, and the unexpected symptoms greatly increase the patient’s distress. In the past, irradiation to reduce vascularity of a glomus tumor has caused
unexpected cochlear damage and tinnitus. Irradiation is no longer used in this context. However, the inner ear is occasionally damaged during irradiation of intracranial tumors, even when cytotoxic drugs are not used. Resultant hearing loss may be accompanied by tinnitus. In Dr. Goodey’s experience, tinnitus is more likely to occur if postirradiation necrosis of the external ear canal also occurs. Presumably, this is because of the added effect of somatosensory stimulation. Subsequent care of the ear canal helps reduce the impact of the tinnitus.

Noise from Organ Imaging Equipment
Especially MRI

Patients sometimes attribute their tinnitus or its increased intrusiveness to the noise associated with having an MRI [30]. Noise levels have been measured in excess of 93 dB [30] and continue throughout the relatively lengthy procedure. There is no associated increased hearing loss. Probably, anxiety, fear, and the claustrophobic environment have contributed, even though the patient has blamed the noise alone for the onset or aggravation of their tinnitus. Any patient with troublesome tinnitus should use hearing protection during an MRI.

Medical Equipment Accidents

During otologic surgery, noise levels generated by otologic drills have been measured as 82–106 dB and by suction measured as 71–84 dB. These are considered acceptable levels. No change in postoperative bone conduction was found [31]. Others have recorded noise levels from air turbine drills of 116 dB and at suction tips of 96 dB [12]. It is widely accepted that there is a high risk of inner ear damage if a drill burr comes in contact with an intact ossicular chain or suction is applied to perilymph in the oval or round window or lateral canal fistula. A hose becoming detached from a compressed air cylinder has triggered severe hearing loss and tinnitus. Other incidents have been reported anecdotally and include a gas explosion.

Neck Manipulation

Patients regularly claim that manipulation of their neck was the trigger for their tinnitus. The resultant tinnitus can usually be modulated by neck movement suggesting proprioceptor disturbance–triggered somatosensory tinnitus. However, in some patients, neck manipulation triggered severe temporary vertigo as well as persistent tinnitus. It may be that on some occasions, neck manipulation triggers tinnitus (and sometimes vertigo) through temporary effects on the vertebral arteries. In others, radiological evidence of facet joint damage caused by manipulation has been demonstrated [32]. If a patient’s neck is to be manipulated vigorously, there should be preceding organ imaging expertly read, the therapist should be experienced, and the therapist should stop immediately if untoward symptoms start to develop.

General Anaesthetic

Tinnitus may be triggered after almost any type of surgical procedure, but mostly if the procedure was under general anaesthetic and a relaxant has been used. There may be postoperative suboccipital headache as well. In these circumstances, the tinnitus can usually be modulated by the neck. Some anesthetists maintain gentle traction on the head and neck while relaxants are wearing off and claim that this reduces the incidence of postoperative headache. In Dr. Goodey’s experience, this maneuver can reduce postoperative tinnitus as well. It is a wise precaution in a patient who already has troublesome tinnitus, especially if they blame it on a previous operation under general anesthesia.

Sometimes, postoperative tinnitus is associated with temporomandibular joint pain and can be modulated by the jaw. In these circumstances, difficulty with intubation may have been the mechanism.

General Reaction to Painful Procedures

In Dr. Goodey’s experience, distressing and painful surgery anywhere in the body can act as the trigger for the onset of tinnitus. The resulting tinnitus may be extremely distressing and difficult to manage. This occurs most often if the pain experienced has been excessive because of complications or inadequate anesthesia, and especially when there are powerful emotional associations because of the nature of the surgery and the consequences of it.
Occasionally, there may be associated sudden hearing loss suggesting microembolism, especially after breast, orthopedic, and cardiac surgery.

Most often there is no measurable change in hearing. There may be some pre-existing hearing impairment, which may have predisposed the patient to the onset of tinnitus in response to the powerful triggering effects of pain, anxiety, fear, and anger.

Part 2: Drug Therapies, Which May Cause Tinnitus

Ototoxicity from Medical Therapy

Over 150 medications and chemicals have been reported to be potentially able to induce hearing loss and/or tinnitus, possibly by acting on both peripheral and central acoustic structures [33–35]. Drug-induced ototoxicity may be reversible or irreversible and associated with both acute and long-term administration of drugs. Among the major classes of ototoxic drugs are the aminoglycosides and other antimicrobial agents, antineoplastic drugs, anti-inflammatory drugs, loop diuretics, antimalarial drugs, and others (Table 42.1). Due to their importance in clinical practice, some ototoxic drugs are discussed in more detail below.

The pharmacological and chemical heterogeneity of drugs, which share the ability to induce hearing loss and/or tinnitus, is noteworthy. Unfortunately, research in this field is often limited by several problems, among which is the lack of a good animal model. As a consequence, the neurobiological basis of drug-induced ototoxicity is still largely unknown and may involve biochemical and physiological changes in discrete parts of the acoustic system [35]. So far, there is no evidence of a common pathway leading to drug-induced damage of acoustic structures.

Chemotherapy of Microbial Diseases

Aminoglycosides

Aminoglycosides are an important group of antibacterial drugs used primarily against Gram-negative aerobic and facultative anaerobic bacteria. Streptomycin is also effective against several tubercular and nontubercular mycobacteria, including *Mycobacterium tuberculosis*, the etiological agent of tuberculosis. Aminoglycosides are bactericidal and act by binding to the 30S subunit of bacterial ribosomes, disrupting the elongation of the peptide chain; they may also impair translational accuracy resulting in misreading of the mRNA sequence. Aminoglycosides are poorly absorbed from the gastrointestinal tract and, therefore, are usually administered parenterally by injection or infusion. Aminoglycosides are well distributed into bodily fluids, except for the eye and the central nervous system. As their metabolism within the body is negligible, aminoglycosides are excreted unaltered by glomerular filtration (serum half-life of 2–3 h). They are also found in breast milk but, as they are not well absorbed orally, these drugs are considered compatible with use during breastfeeding [36]. Aminoglycosides are classified as an FDA pregnancy category D (positive evidence of human fetal risk, but the benefits from use in pregnant women may be acceptable despite the risk). Therefore, they should be used during pregnancy only when the alternatives are worse.

All aminoglycosides are able to induce both reversible and irreversible damage at cochlear, vestibular, and renal level. Nevertheless, aminoglycosides are still among the most commonly used antibiotics worldwide, mainly because of their cost effectiveness [33], but also to face the emergence of bacterial strains with advanced patterns of antimicrobial resistance [37, 38]. Aminoglycoside toxicity correlates with the total amount of drug administered and occurs in almost all patients exposed to a toxic dose. The risk of toxicity is increased if impaired renal function is allowed to cause the serum level to rise [39]. Abnormally high sensitivity to the ototoxic effects of aminoglycosides (idiosyncrasy) may also be an inherited trait, and several mutations at the mitochondrial genome level have been identified [40, 41]. Cochlear and vestibular structures appear to differ in sensitivity to aminoglycosides-induced damage. Indeed, streptomycin and gentamicin are mainly toxic at the vestibular level, while amikacin, neomycin, dihydrostreptomycin, and kanamycin act primarily at the cochlear level [34, 41]. Netilmicin appears to be as effective as gentamicin, but is less ototoxic [38, 41].

Both animal and human studies show that aminoglycosides affect outer hair cells first and later the inner hair cells. Degeneration of hair cells usually starts at the basal turn and progresses toward the apex. The mechanisms of aminoglycoside-induced ototoxicity
Table 42.1  Drugs which are claimed to cause ototoxicity and/or tinnitus

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<tr>
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<th>Ototoxic</th>
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<td><strong>Drugs acting at synaptic and neuroeffector junctional sites</strong></td>
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<td>β2-selective adrenergic receptor agonists</td>
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<td>Procaterol</td>
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<td>Nonselective β adrenergic receptor antagonists</td>
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<td>Serotonin receptor agonists</td>
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<td>Eletriptan</td>
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<tr>
<td>Ergonovine</td>
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<tr>
<td>Methyl ergonovine</td>
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<tr>
<td><strong>Drugs acting on the central nervous system</strong></td>
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<tr>
<td>Valproic acid</td>
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<td>Flecaïnide</td>
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<td>Antidepressants – Tricyclic</td>
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<td>Antidepressants – SSRI</td>
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<td>Drugs affecting renal and cardiovascular function</td>
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<td>Antiarrhythmics</td>
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<td>Dihydrochinidime</td>
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<td>Imidapril</td>
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<td>Benazepril</td>
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(continued)
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<th>Drug Class</th>
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<td><strong>Chemotherapy of parasitic infections</strong></td>
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<tr>
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<td>Quinine</td>
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<td>Sulfadoxine – pyrimethamine</td>
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<td><strong>Chemotherapy of neoplastic diseases</strong></td>
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<td><strong>Platinum compounds</strong></td>
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<td><strong>Hormones and hormone antagonists</strong></td>
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have not been fully characterized; however, several mechanisms have been proposed, including disruption of mitochondrial protein synthesis, generation of reactive oxygen species (ROS), and excitotoxicity from enhancement of the glutamatergic N-methyl-D-aspartate (NMDA) receptor function [39, 41].

**Approaches to Protection**

Due to the widespread use of these drugs, prevention of aminoglycosides-induced ototoxicity is very important. Patients should also be questioned for symptoms of tinnitus, decreased hearing, dizziness, disequilibrium, and problems of ocular fixation. Careful monitoring of serum levels together with audiological or vestibular function tests are essential components of the standard of care required to reduce the incidence of aminoglycoside ototoxicity.

Scientific research is now focused on the biological mechanisms underlying aminoglycosides-induced damage in order to develop coherent attempts at protection such as administration of antioxidants or iron chelators, interference with cell death signaling pathways, and blockade of glutamate NMDA receptor [41–44]. At present, experimental evidence shows a decrease in ototoxicity when antioxidants or iron chelators are co-administered with aminoglycosides. However, successful translation of experimental evidence to the clinic is a slow process requiring consideration of many points. Therefore, the currently more “orthodox” approach of monitoring serum drug levels and ototoxicity symptoms remains the standard of care [39, 45].

It may be impractical to monitor serum drug levels and perform audiological or vestibular function tests on all patients receiving treatment with aminoglycosides. It is essential to do so in those patients with high risk for developing ototoxicity, including those receiving prolonged treatment courses, those who have had previous aminoglycoside therapy, those with sensorineural hearing loss, or patients in whom inner ear damage would create a disproportionately major handicap. Because the incidence of ototoxicity is related to the serum aminoglycoside concentrations, it is critical to reduce the maintenance dosage of these drugs in patients with impaired renal function or who are concomitantly taking loop diuretics [46] or nephrotoxic drugs. The elderly are especially at risk from aminoglycosides, as their renal function may be significantly impaired without increase in serum creatinine.

Idiosyncratic hearing loss induced by aminoglycosides is, in theory, preventable by genetic screening to identify those at risk (e.g., individuals with the m.1555 A>G mutation). The use of such genetic screening is questioned because of the high cost of the tests. However, when the expenses of genetic screening are compared to the lifelong management of a profoundly deaf child, the cost effectiveness of genetic screening may prove very favorable [40].

**Chemotherapy of Neoplastic Diseases**

**Platinum Compounds–Cisplatin**

In theory, any drug with the capacity to destroy malignant cells should be regarded as having the potential to damage the cells of the cochlea and cause hearing loss and tinnitus. Cisplatin (cis-diamminedichloroplatinum) is an inorganic platinum coordination complex used alone or in combination with other anti-cancer agents. Its main application is in the medical therapy of malignancies including sarcoma, small-cell lung cancer, germ cell tumors, lymphoma, and ovarian cancer [36, 47]. Cisplatin disrupts DNA function in several ways. It inhibits DNA synthesis by the formation of DNA cross-links; it denatures the double helix and covalently binds to DNA bases interfering with replication and transcription [48, 49]. Cisplatin is administered parenterally either by the intravenous or by the intraperitoneal route. It is not metabolized but is excreted mainly by the kidney (>90%). A few studies have examined the excretion of cisplatin into human milk with contradictory results, and therefore, breastfeeding during cisplatin therapy should be considered contraindicated. Cisplatin is nephrotoxic, neurotoxic, mutagenic in bacteria, produces chromosomal aberrations in animal cells in tissue culture, and is teratogenic and embryotoxic in mice [50]. There are no adequate well-controlled studies in pregnant women [51], and Cisplatin is therefore classified as FDA pregnancy category D.

Cisplatin ototoxicity seems to be mediated by the generation of ROS in the cochlear tissue and has been shown to act on at least three major targets: the organ of Corti, the spiral ganglion cells, and the lateral wall [52]. Increased ROS and organic peroxide following
the administration of ototoxic doses of cisplatin would overwhelm the antioxidant potential of the cochlear cells, leading to calcium influx, which would activate the apoptotic pathway causing cell death [39, 52]. Several genetic variants have been associated with increased sensitivity to cisplatin-induced ototoxicity [52–54]. Research in this field is still in an early phase. However, it is conceivable that a better understanding of the genetic variants associated with cisplatin-induced ototoxicity may be an important step toward case selection and safer cisplatin treatment [53, 55, 56].

The clinical presentation of cisplatin-induced damage to the inner ear includes tinnitus and high-frequency sensorineural hearing loss. The hearing loss is usually modest but can be permanent and can progress to involve the lower frequencies. The tinnitus is often more irksome than the modest loss of hearing. The risk of inner ear damage is increased by prior irradiation and concomitant use of aminoglycosides.

**Approaches to Protection**

In general, patients who embark on antineoplastic chemotherapy are not only well monitored but also well informed. They are aware and have accepted the possibility of adverse consequences of drugs, including the development of tinnitus and some loss of hearing. Nevertheless, research on new methods of protection against ototoxicity (such as chemoprotection) is definitely needed. At present, the only strategy for reducing cisplatin-induced ototoxicity is based on limiting the total dose per cycle, the cumulative dose, and the dose intensity, which inevitably limits the antineoplastic effectiveness [57, 58]. Various strategies have been proposed to reduce cisplatin ototoxicity by chemoprotectants; in particular, an “upstream approach” to prevent the generation of ROS with antioxidants and a “downstream approach” using inhibitors of molecules involved in the apoptotic cell death pathway (such as caspases and p53). Indeed, the administration of several antioxidants does seem to be able to limit cisplatin ototoxicity [44, 59, 60]. Unfortunately, this approach has limited clinical usefulness because of the potential for negative interaction between antioxidants and antineoplastic drugs, resulting in reduced therapeutic effectiveness.

A particularly important issue in protection from cisplatin ototoxicity is the extensive use of this drug in pediatric patients, mainly because of its effectiveness in increasing the survival rate for children with cancer [47, 61, 62]. While new anti-cancer treatment protocols are very successful in improving pediatric patient survivals, they also subject the children to toxicities, which may profoundly affect a child’s life and development [63, 64]. The reported incidence of cisplatin-induced ototoxicity in children varies from 10 to 85% of cases. Nevertheless, the implications of hearing loss to speech and language development are very important in very young children, whereas educational and psychosocial problems are more important for older children [63].

A child’s age at treatment and the cumulative dose of cisplatin are the two most important risk factors in predicting moderate to severe hearing loss in children [62, 65]. During cisplatin therapy and a subsequent follow-up, pediatric patients should be audiometrically tested for the development of drug-induced sensorineural hearing loss [63, 66].

Several recent reports have shown a protective effect of amifostine, a thiolic cytoprotectant, in pediatric cancer patients treated with cisplatin [67–69]. However, evidence is contradictory, and more research is needed [70–72].

**Chemotherapy of Parasitic Infections**

Malaria is one of the most severe public health problems worldwide and a leading cause of death and disease in many third-world countries [73]. In Western world countries in which malaria has never existed or has been eliminated, the greater majority of cases occur either in travelers returning home or in migrants arriving from areas where malaria is endemic – “imported malaria” [74].

Each year, millions of people from malaria-free countries travel to areas where malaria is common and are therefore subjected to antimalarial chemoprophylactic treatment, which includes administration of several ototoxic drugs [75–78].

**Quinolines and Related Compounds**

Intravenous quinine dihydrochloride is currently the first-line antimalarial drug for the treatment of severe malaria in the UK [79]. Quinine is also sometimes
Complications to Medical Treatment

used for night cramps and chloroquine for arthritis; chloroquine and hydroxychloroquine are also used in the treatment of rheumatoid arthritis and lupus associated arthritis. Quinoline derivatives are thought to exert their antimalarial effect by reaching high concentrations in the *Plasmodium* digestive vacuole and preventing the biocrystallization of toxic heme released during proteolysis of hemoglobin into hemozoin. Failure to inactivate toxic heme would poison the parasite, possibly via oxidative damage to plasma membranes [36, 80]. Quinolines are well absorbed from the gastrointestinal tract and may also be administered parenterally either by injection or by infusion. Although rare in western countries, quinine and quinidine overdose may lead to severe toxicity and death related to cardiovascular and neurological effects, particularly in children [81, 82]. Although several skeletal and muscular malformations have occurred in laboratory animals, quinoline derivatives appear safe in human pregnancy and during lactation [83–85].

Quinine is known to cause reversible hearing loss and tinnitus in both humans and animal studies [86–88]. Ototoxicity also has been reported in association with the use of other quinoline-type antimalarial drugs including chloroquine, hydroxychloroquine, and mefloquine [89–91]. The biological bases of quinolines-induced ototoxicity have not been fully resolved. However, some experimental evidence suggests that quinine may affect the function of calcium-dependent potassium channels and reversibly alter the mechanical properties of outer hair cells [92–95].

**Approaches to Protection**

Quinoline derivatives cause hearing impairment and tinnitus without vestibular disturbance. Both the hearing loss and the tinnitus are usually reversible, but the changes can progress to cochlear degeneration, permanent hearing impairment, and increased likelihood that the tinnitus will persist [96, 97]. Young and unborn children are probably more susceptible to quinoline-induced hearing loss [98, 99]. The ototoxic effects of quinine may be potentiated by doxycycline, an antibiotic, which is sometimes used with quinine in the prophylaxis or treatment of malaria [100]. On its own, doxycycline is not thought to be ototoxic. It has been reported that chloroquine-induced damage to the cochleovestibular system can recover if the medication is stopped and appropriate therapy is instituted with steroids and plasma expanders [89].

Mefloquine is also ototoxic, but in addition to hearing impairment and tinnitus, it may also cause vestibular disturbance [99, 101]. The tinnitus and hearing impairment are more likely to be permanent than with the other antimalarial drugs.

**Salicylates**

Acetylsalicylic acid (aspirin) was one of the first drugs to have come into common usage. Despite the introduction of new agents, it is still the analgesic, antipyretic, and anti-inflammatory drug most widely used in the world [102, 103]. Approximately 35,000 metric tones are produced and consumed annually, which is enough to make over 100 billion standard aspirin tablets every year [102, 104]. Besides its use as analgesic, antipyretic, and anti-inflammatory agent, aspirin is also extensively used in the prevention and treatment of various aspects of cardiovascular disease [105, 106], and it is under investigation in a number of other medical conditions including cancer [103, 107, 108].

Most pharmacological effects of salicylates are due to inhibition of prostaglandin formation via blockade of cyclooxygenase. Although there is no agreement about their molecular mechanisms of action, salicylates probably act because of their content in salicylic (orthohydroxybenzoic) acid [36, 102]. Aspirin also possesses distinct protein-acetylating capabilities, which may account for its unique pharmacological profile [109]. Salicylates are rapidly adsorbed from the gastrointestinal tract and well distributed in the body tissues and fluids. About 50% of orally administered aspirin is de-acetylated to salicylate in the liver immediately after absorption. Common metabolites are salicyluric acid, salicyl phenolic or acyl glucuronides, and gentisic acid. Salicylates are excreted in the urine. Plasma half-life of aspirin is about 15 min while the half-life of salicylate is between 2 and 12 h. Aspirin taken in low dose during pregnancy is generally considered safe. However, full-dose aspirin taken in the third trimester is considered to be in FDA pregnancy category D. Aspirin is excreted into human milk in small amounts and should be given to nursing mothers with caution [110].

Salicylates have been recognized as ototoxic longer than almost any other drug [111]. The main ototoxic
effects of salicylates are sensorineural hearing loss and tinnitus. Salicylate-induced hearing loss is typically mild to moderate, symmetrical, and flat or high frequency [112, 113]. The tinnitus is often described as a continuous high pitch sound of mild loudness. The neurobiological mechanism of salicylate-induced hearing loss and tinnitus remains obscure. However, several papers have shown that multiple actions of salicylates throughout the acoustic system may contribute. Salicylates administration profoundly affects cochlear function, possibly through downregulation of outer hair cells electromotile response with resultant decrease in cochlear neural output [114, 115]. Several other neurotransmitter systems are involved in salicylates ototoxicity at central level, including the glutamatergic and GABAergic system [112, 116–118]. Interestingly, sodium salicylate has been shown to partially protect against cisplatin ototoxicity and aspirin to partially protect against aminoglycoside ototoxicity, possibly because of their antioxidant properties [42, 119, 120].

**Approaches to Protection**

Salicylate-induced hearing loss is almost always reversible. Associated tinnitus usually subsides as hearing recovers, although this is not always the case. Quite large doses (6–8 g daily) are required to cause hearing loss and tinnitus [117]. The onset of tinnitus can be helpful as an early indicator of salicylate intoxication or salicylysm [121, 122]. Salicylysm is a potentially fatal poisoning that, partly because of the enormous amount of aspirin produced and consumed annually, remains a common cause for treatment in emergency departments, especially of children [123]. It is also noteworthy that salicylate intoxication is being reported increasingly often as a consequence of the use of herbal medicines [124–126].

**Miscellaneous Drugs that are not Considered Ototoxic**

Several different drugs may cause or aggravate tinnitus often without an effect on hearing. Some of these drugs may ease tinnitus in some patients, yet aggravate or cause it in others. Among these drugs are lidocaine, anticonvulsants, antidepressants, cannabinoids antihypertensives, β-adrenergic blocking agents, opioids (buprenorphine), caffeine, antihistamines, and several others. Unfortunately, the available evidence on the vast majority of these drugs is scarce and much of it anecdotal.

**Lidocaine**

Lidocaine is the prototypical amide-type local anesthetic, as well as one of the drugs most consistently reported as being efficacious in relieving subjective tinnitus. Available data consistently report that intravenous lidocaine is able to dose dependently inhibit tinnitus in approximately 60% of patients [127–130], although some authors report lower figures [131]. In some patients, tinnitus inhibition is complete, while in a small number of patients an exacerbation is perceived. Lidocaine is a voltage-gated sodium channel blocker able to reduce nerve cell responsiveness to stimuli in a time- and voltage-dependent fashion [132–134]. Lidocaine can also reversibly block voltage-gated potassium channels at concentrations compatible with plasma levels linked to tinnitus inhibition [135]. Since voltage-gated potassium channels are reported to play a key role in the encoding of auditory information, this effect of lidocaine may be relevant [136–139]. The site of action of lidocaine still remains unclear; earlier studies found a cochlear involvement [128, 140]; however, much evidence is now accumulating, which indicates a central site of action. In particular, auditory brainstem responses [141] and brain imaging techniques showed a central action of lidocaine and suggested that this drug may affect the functional linkage of several brain areas including auditory thalamus, auditory cortex, dorsolateral prefrontal cortex, and limbic system [142–144].

**Anticonvulsants**

Anticonvulsant drugs are increasingly used in the treatment of several nonepileptic conditions, including various psychiatric disorders, pain syndromes, and tinnitus [145]. Evidence of benefit from antiepileptic drugs in nonepileptic conditions varies among different drugs, but there is, in general, a lack of randomized double-blind trials in the literature [145, 146]. Diverse pharmacological mechanisms of action are responsible for the
therapeutic effects of antiepileptic drugs including effects on voltage-gated sodium and calcium channels, and neuronal inhibition mediated by γ-aminobutyric acid receptors. However, it may be hypothesized that the common final action is to reduce the tendency of neurons in sensory pathways to fire spontaneously or at inappropriately high frequencies. Carbamazepine, sodium valproate, and phenytoin are all incriminated as triggers and aggravators of tinnitus in some patients while they may help reduce it in others. Unfortunately, clear scientific evidence is unavailable at the moment.

**Antidepressants**

Antidepressants are widely used in many therapeutic protocols, including those for the management of tinnitus [147, 148]. This may be mainly because of the well-described comorbidity between major depressive disorders and tinnitus [147, 149, 150].

Among all antidepressants used for tinnitus, a particular interest has been paid to tricyclic drugs mainly because of the analgesic effect of this class of drugs [151, 152], in view of the proposed etiological correspondence between tinnitus and neuropathic pain [153, 154]. However, tricyclics may trigger or aggravate tinnitus in some patients. Amitriptyline has been reported as causing tinnitus in one case [155] and subsequently reported as being helpful in treating major depressive symptoms in tinnitus [156]. Recent evidence confirms the tinnitus-inducing effect of amitriptyline in some patients [157].

Selective Serotonin Reuptake Inhibitors (SSRIs) are the most widely prescribed antidepressants in many countries, mainly because of their clinical effectiveness and the reduced toxicity when compared to tricyclics. SSRIs are supposed to act by inhibiting the reuptake of serotonin into the presynaptic cell, thus causing a temporary increase in levels of 5-HT within the synaptic cleft. Despite their antidepressant effectiveness, SSRIs are frequently reported as inducing tinnitus either as a side effect of therapy or as a consequence of drug discontinuation syndrome [158–160]. Fluoxetine occasionally has a dramatic triggering effect, which may persist after the drug is stopped. The specific effectiveness of SSRIs in tinnitus has been recently questioned by several high-quality studies [148, 161, 162].

Among atypical antidepressants, the aminoketone bupropion acts as a norepinephrine and dopamine reuptake inhibitor and also as a nicotinic antagonist. Bupropion was originally marketed as an antidepressant but is now a fundamental drug in smoking cessation therapies along with nicotine replacement products [163, 164]. Bupropion is among the most frequently prescribed psychotropic drugs in the United States. It is not considered an ototoxic drug, but its association with tinnitus has been consistently reported in case reports as well as literature [165, 166]. Bupropion-induced tinnitus appears to be a temporary effect that disappears after the drug is discontinued. More research is needed to clarify the relationship between bupropion use and the development of tinnitus.

**Cannabinoids**

Cannabinoids (mainly tetrahydrocannabinol, cannabidiol, β-caryophyllene, and cannabigerol) are now being increasingly used in the treatment of several conditions including spasticity, multiple sclerosis, painful conditions (including neuropathic pain), asthma, and closed-angle glaucoma [167–169]. Natural and synthetic cannabinoids interact with the bodily endocannabinoid system by binding to specific G-protein–coupled cannabinoid receptors (CB1 and CB2). Agonists to CB receptors activate multiple intracellular signal transduction pathways, leading to a very complex picture involving inhibition of adenylate cyclase, activation of inwardly rectifying K channels, alteration of intracellular Ca levels, and influences on other ion channels and kinases [170–172]. Cannabinoid receptors are differentially expressed in the body tissues. CB1 is present in the brain and in the periphery it is present in adipose tissue, the gastrointestinal tract, skeletal muscles, heart, and in the reproductive system. CB2 is mainly expressed in the immune system [173].

As well as the chemically pure drug (such as Dronabinol and nabilone), cannabinoids are also available in some jurisdictions in the form of dried Cannabis indica leaves (marijuana). They are then generally self-administered by inhalation of marijuana smoke or through the gastrointestinal system. Despite consistent evidence of clinical efficacy and relative safety [174, 175], medical cannabis remains a controversial issue, mainly because marijuana is one of the most widely used recreational drugs in the world and remains illegal in many countries.

Cannabis smoke has been anecdotally reported to temporarily cause tinnitus in some patients, but
dramatically relieves it in some others. However, despite the reported occurrence of CB1 in the cochlear nucleus [176], there is no scientific evidence available of a direct role of cannabinoids in neurobiological basis of tinnitus [177]. However, more research on cannabinoids and tinnitus may be advisable, since a potential for clinical use may be obscured by other considerations. [175]

**Drug-Induced Ototoxicity: Final Considerations**

Ototoxicity is an adverse effect of several classes of drugs, such as the aminoglycosides, antineoplastic drugs, anti-inflammatory drugs, loop diuretics, antimalarial drugs, and others. Further, occasional cases of ototoxicity have been reported for a wide variety of other therapeutic compounds and chemicals.

Ototoxic agents can impair the sensory processing of sound at many cellular or subcellular sites. Much research has been performed to investigate the causes and the pathophysiology of ototoxicity to try to prevent this complication. However, the neurobiological mechanisms underlying ototoxicity have not been established for most of these drugs, and structure-toxicity relationships have not been determined. It is therefore quite difficult to predict the ototoxic potential of new drugs, and rational approaches to the prevention of ototoxicity are still lacking. In addition, the simultaneous administration of multiple agents, which are potentially ototoxic, can lead to synergistic loss of hearing. Exposure to loud noise may also potentiate hearing loss due to ototoxic drugs.

Drug-induced ototoxicity, although not life threatening, may induce considerable damage and cause severe disability. When increasing ototoxicity occurs, the ototoxic medication has to be discontinued if permanent hearing loss and/or tinnitus are to be minimized.

Although ototoxic injury is sometimes unavoidable, certain measures may reduce the risk. Prevention of drug-induced ototoxicity is generally based upon consideration and avoidance of relevant risk factors, as well as on monitoring renal function, serum drug concentrations, and cochlear and auditory functions before and during drug therapy.

**Conclusions**

- The treating physician should consider choosing a therapeutically equivalent nonototoxic drug whenever one is available, especially in patients with a heightened risk such as pre-existing cochlear hearing loss and renal insufficiency.
- During therapy with potentially ototoxic medications, the lowest dose compatible with therapeutic efficacy should be used.
- When indicated, periodically monitor serum peak and trough levels.
- Simultaneous use of multiple ototoxic medications (e.g., aminoglycosides and loop diuretics) should be avoided whenever clinical circumstances permit, as their concomitant use may increase the risk of permanent deficit.
- When early detection is important, audiological monitoring should include the very high frequencies as, generally, ototoxic drugs first destroy hearing in the very high frequencies, which are not normally tested (those above 8,000 Hz).
- Should a patient develop auditory (hearing loss and/or tinnitus) or vestibular (vertigo and/or disequilibrium) symptoms during therapy with a potentially ototoxic medication, audiometric testing and otological assessment should be arranged urgently especially if there is reluctance to stop the ototoxic medication.

**References**


**Keypoints**

1. It is now recognized that many forms of tinnitus-related neural activity are much more complex and multimodal than ever thought.
2. It has become evident that contribution of non-auditory pathways is involved in eliciting or modulating many forms of tinnitus.
3. Many forms of tinnitus can be modulated by different actions such as forceful muscle contractions of the head and neck as well as eye movements.
4. Somatosensory stimulation such as that from pressure of myofascial trigger points, cutaneous stimulation at specific locations, electrical stimulation of the median nerve and hand, finger movements, and orofacial movements can also modulate or cause tinnitus, as can pressure applied to the temporomandibular joint or lateral pterygoid muscle.
5. This chapter discusses the causes of somatosensory tinnitus and in particular the influence from both head and neck regions on the auditory pathways in individuals with tinnitus.

**Keywords** Tinnitus • Somatic • Somatosensory • Central nervous system • Muscle • Cervical spine • Temporomandibular joint.

**Abbreviations**

MTP = Myofascial trigger point
AMTP = Active myofascial trigger point
LMTP = Latent myofascial trigger point

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**Introduction**

For many years, tinnitus was thought to arise almost exclusively from abnormal neuronal activity within the auditory pathway. However, accumulated evidence suggested that tinnitus-related neural activity is much more complex and multimodal than previously anticipated.

More often than ever thought, tinnitus can be evoked or modulated by inputs from somatosensory, somatomotor, and visual-motor systems in some individuals. This means that the psychoacoustic attributes of tinnitus might be changed temporarily by different stimuli, such as:

- Forceful muscle contractions of head, neck, and limbs [1–3];
- Eye movements in horizontal or vertical axis [4–7];
- Pressure of myofascial trigger points [8];
- Cutaneous stimulation of the hand/fingertip region [4] and the face [9];
- Electrical stimulation of the median nerve and hand [10];
- Finger movements [11];
- Orofacial movements [12];
- Pressure applied to the temporomandibular joint or lateral pterygoid muscle [13, 14].

Such temporary changes are known as modulation of tinnitus. So, the contribution of non-auditory pathways has become more and more evident in eliciting or modulating existent tinnitus.

Although this phenomenon is yet to be fully understood, it seems to be clinical evidence of the existing neural connections between the somatosensory and auditory systems, whose “activation” may play a role in tinnitus. Anatomic and physiological findings in animal studies have shown that the trigeminal and
dorsal root ganglia relay some afferent somatosensory information from the periphery to secondary sensory neurons in the brainstem, specifically, the spinal trigeminal nucleus and dorsal column nuclei, respectively [15]. Each of these structures sends excitatory projections to the cochlear nucleus. Mossy fibers from the spinal trigeminal and dorsal column nuclei terminate in the granule cell domain while en passant boutons from the ganglia terminate in the granule cell domain and core region of the cochlear nucleus. Single unit and evoked potential recordings in the dorsal cochlear nucleus indicate that these pathways are physiologically active.

So, these clinical findings strongly suggest that those who are able to modulate their tinnitus should be considered as a specific subgroup of patients. Among all types of modulating factors that have been described, we are particularly interested in the influence of both head and neck regions on the auditory pathways.

Now, there is yet no consensus on the definition of “somatosensory tinnitus,” and this term has been used with different meanings. A group of researchers in the Tinnitus Research Initiative is presently working to define and differentiate “somatosensory tinnitus” (primary origin in head and neck trauma, dental or cervical manipulation, or even in unknown chronic pain) from “somatosensory modulation” (auditory origin with temporary somatosensory influence in loudness, pitch, or localization). Although many aspects still need clarification, we have already progressed in establishing some specific causes, methods of diagnosis, and treatment options to this subgroup, which will be described in this chapter.

Theories About Tinnitus Modulation

It is widely known that reorganization or re-mapping of specific central nervous areas occurs as a normal response of brain tissue to injury [16, 17]. However, as any double-edged sword, it is not possible to predict whether injury-induced plasticity will end up in limited or cross-modal effects, which in turn may result in compensatory or pathologic effects. Neuroplasticity is often implicated in tinnitus, and aberrant cross-modal plasticity seems to play a role in recently described cases of tinnitus evoked by somatosensory activation. This suggests that abnormal interaction between different sensory modalities, sensorimotor systems, neurocognitive, and neuroemotional networks may contribute to certain aspects of tinnitus [17].

Tinnitus modulation indicates that the psychoacoustic attributes of tinnitus change temporarily during some sort of stimuli [18]. Some of these modulation patterns (gaze-evoked, finger-evoked, and cutaneous-evoked tinnitus) were first described after acute unilateral total deafferentation of the auditory afferents, usually caused by the removal of skull base and posterior cranial fossa tumors. Some authors have hypothesized that in this form of modulation, important plastic changes occurred in the central nervous system after such deafferentation.

However, our own clinical experience showed that other types of modulation occur regardless of any surgical manipulation or degree of hearing loss [2, 3, 19]. An altered afferent input to the auditory pathway may be the initiator of a complex sequence of events finally resulting in the generation of tinnitus at the central level of the auditory nervous system. The effects of neural plasticity can generally be divided into early and later modifications, depending on the time of onset. Unmasking of dormant synapses, diminishing of (surround) inhibition, and generation of new connections through axonal sprouting are early manifestations of neural plasticity, resulting in lateral spread of neural activity and development of hyperexcitability regions in the central nervous system. The remodeling of tonotopic receptive fields within auditory structures (dorsal cochlear nucleus, inferior colliculus, and auditory cortex) seems to be a late manifestation of neural plasticity. The modulation of tinnitus by stimulating the somatosensory system might be explained by activating auditory regions through the non-classical pathway.

Tinnitus Modulation by Muscle Contractions

Sometimes tinnitus patients spontaneously report that contractions of head and neck muscle may change the loudness or pitch of their tinnitus. However, recent studies showed that a surprisingly large number of patients modulate tinnitus when they are specifically tested for it. Levine initially found that 68% of patients with tinnitus experienced some kind of modulation
when performing muscular contractions [1, 20]. Regardless of etiology or audiometric pattern, 71% could modify their tinnitus with a variety of cephalocervical isometric maneuvers or extremity contractions [21]. The head/neck isometric maneuvers were much more effective in modulating tinnitus than contractions of the limbs. Using a control group, Sanchez et al. pointed out that 65.3% of patients modulated loudness or pitch of their tinnitus during muscle contractions, while 14% of asymptomatic subjects could evoke tinnitus perception during the same maneuvers [2]. Later, other studies confirmed that the majority of tinnitus patients can modulate the phantom sound by stimulation of the somatosensory system [3, 19, 21, 22].

Considering the structure of the auditory pathway, it consists of several well-defined centers, although precise information about their interaction is still lacking. The cochlear nucleus is the first central nucleus of the auditory pathway, receiving information from the cochlear hair cells. In higher portions of the auditory pathway, the lemniscal system sends the received information to the primary cortical auditory areas, whereas the extralemniscal portion of the ascending pathways transmits auditory information to associated areas [10]. Many neurons of the extralemniscal system receive information from other sensorial tracts, such as the somatosensory system [23, 24].

The cuneate and gracile nuclei collectively form the dorsal medullary nucleus, whose position in the somatosensory system is analogous to that of the cochlear nucleus in the auditory system. It receives information directly from the dorsal roots, which in turn get information from the proprioceptive, tactile, and vibratory receptors of the body surface. The lateral cuneate nucleus is the end point of afferent fibers from the neck, ear, and suboccipital muscles, and carries information on head and ear position needed to process the acoustic information [25]. Because of reciprocal connections between the auditory and somatosensory systems, these authors postulated that projections from the cuneate to the cochlear nucleus may lead to excitation of the cochlear nucleus. Nevertheless, some electrophysiological studies in cats showed that the final effect of cuneate nucleus activation is the inhibition of the dorsal cochlear nucleus [26]. The exact mechanisms responsible for somatic modulation of tinnitus are currently unclear. If one considers that tinnitus results from aberrant neuronal activity within the auditory pathway, this could mean that somatosensory stimuli coming from head and neck muscle contractions might, through a multisynaptic pathway, disinhibit the ipsilateral cochlear nucleus, producing an excitatory neuronal activity within the auditory pathway that results in tinnitus.

As muscular contraction represents an activation of the somatosensory system, these anatomical connections between both systems might explain the influence of voluntary muscle contractions upon some types of tinnitus, thereby stimulating or inhibiting this symptom and presenting clinically as a modulation factor. In fact, we have seen patients with a typical history of acoustic trauma that could also clearly evoke tinnitus by several different stimuli, including during abdominal contraction.

Tinnitus Modulation Through Myofascial Trigger Points

Myofascial trigger points (MTP) are small hypersensitive spots located within the palpable taut bands of skeletal muscle fibers. Either spontaneously or under mechanical stimulation, they may cause local and referred pain [27].

MTP may be active (AMTP) when their stimulation causes a pattern of referred pain that is similar to the patient’s pre-existent pain complaint or may aggravate such pain [28]. They are frequently found on the neck, shoulders, pelvic girdle, and masticatory muscles [29], where they provoke spontaneous pain or movement-related pain.

MTP can also be latent (LMTP), which are located in symptom-free areas and provoke local and referred pain only when stimulated [28].

Although MTP may be detected in pain-free subjects, they are typical of patients with myofascial pain syndrome, who often complaint of an associated tinnitus [30].

Travell and Simons first reported that MTP palpation of the sternal division of the sternocleidomastoid evoked a sound perception in a tinnitus-free patient [27]. Later, Eriksson et al. described a patient who noticed differences in tinnitus when palpating a MTP in the sternocleidomastoid. Such association has also been verified in studies where tinnitus patients had their conditions improved through anesthesia-based MTP deactivation [31].
Recently, Rocha et al. (2007) [8] investigated whether myofascial trigger points could modulate tinnitus and examined the association between tinnitus and MTP. They evaluated 94 subjects with tinnitus and 94 without the disorder, who underwent bilateral digital pressure of nine muscles of the head, neck, and shoulder girdle usually tested in myofascial pain syndromes (infraspinatus, levator scapulae, superior trapezius, splenius capitis, scalenus medius, sternal portion of sternocleidomastoid, posterior digastric, superficial masseter, and anterior temporalis). Temporary tinnitus modulation was observed in 56% of the subjects during digital pressure, mainly in the masseter, splenius capitis, sternocleidomastoid, and temporalis muscles. The rate of tinnitus modulation was significantly higher on the same side of MTP tinnitus subjects to examination in six out of the nine muscles. A strong association between tinnitus and the presence of MTP was observed, as well as a laterality association between the ear with tinnitus and the side of the body with MTP [19].

We initially assumed that only AMTP (related to pain) would be able to modulate tinnitus. However, the compression of LMTP may also end up with modulation. One possible explanation is that both active and latent MTP evoke referred pain when stimulated. Another interesting discovery of this study was the fact that MTP located in head and neck muscles produced more tinnitus modulation than those located in the shoulder girdle, which supports previous study [2, 20] findings, in which head and neck muscular contraction maneuvers produced more modulation than those of the members. These results can be possibly explained by neuroanatomy, since connections between somatic and auditory pathways at the cephalic level would be richer.

One of the mechanisms that explains referred pain is transmission by autonomic pathways [32]. The autonomic phenomena referred to other areas besides the MTP region can be explained by increased sensitivity of sensory nerve endings (thin terminal axons) at the MTP region and consequent neural mechanisms to spread referred pain [25]. Whenever those LMTP remain in a given subject for lengthy periods, they give rise to sensitization of nervous fibers associated to vasoconstriction due to increased sympathetic neurovegetative activity [33]. According to Hubbard and Berkoff, sympathetic activity explains the autonomic symptoms associated with MTP and provides a mechanism through which local injury and nociception cause local tension. It is now accepted that there is direct sympathetic innervation to the intrafusal fibers of muscle spindles. In some tinnitus patients, the sympathetic nervous system apparently plays an important role. Studies have found that blocking the sympathetic input to the ear or a sympathectomy can alleviate tinnitus in some patients. Thus, the autonomic nervous system (sympathetic) may explain some of the findings regarding the effects of MTP stimulation on tinnitus.

Thus, the possible explanation for the relationship between tinnitus and MTP would be not only somatosensory–auditory system interactions but also the influence of the sympathetic system.

**Tinnitus Modulation During Tender Point Compression**

Tender points are discrete areas of pain in response to palpation on body surfaces and can be identified in many people, but those suffering from chronic pain disorders tend to be more affected. The difference between MTP and tender points is the location of pain and the point of maximum tenderness that causes the symptoms. MTP refers pain to a distant spot upon pressure; tender points do not [34]. Researchers have been debating whether trigger points are a subset of tender points.

Even with such similarities, there has been no report of tender points being able to modulate tinnitus. However, during the examination of 11 patients with tinnitus and frequent regional pain for at least 3 months in the head, neck, and shoulder girdle (ten with myofascial pain syndrome and one with only tender points), we surprisingly found that 5 of them modulated tinnitus upon digital pressure on some tender points, besides the modulation by trigger points. Moreover, two other patients only modulated tinnitus by tender points, including the subject who did not have myofascial pain syndrome.

As this finding appeared by chance during the development of a study focused to trigger points, new clinical studies with bigger samples are necessary in order to demonstrate a possible relationship between tender points and tinnitus, with or without an associated myofascial trigger point.
Tinnitus Associated with Cervical Whiplash

As a consequence of cervical whiplash, extensive injuries to the cervical joints, ligaments, and discs may occur [35]. These bony and soft tissue injuries may lead to a variety of clinical manifestations [36]. Neck pain is the most common symptom, reported in 88–100% of cases [37]. Surprisingly, tinnitus and other otological symptoms are found in approximately 10–15% of the patients [38–40]. However, among 109 patients evaluated, none reported otological symptoms in the acute phase following the whiplash injury [41]. In our opinion, a possible explanation might involve the secondary vicious muscular postures that patients adopt in order to avoid neck pain. Considering the relation previously described between tinnitus modulation and muscular tension, myofascial trigger and tender points [8, 19, 31, 42], it is possible that secondary findings in patients with whiplash injury may justify the later onset of tinnitus. As the relationship between whiplash itself and tinnitus is yet controversial, caution is recommended whenever attributing these symptoms to such an injury.

On the other hand, some studies have suggested a possible link between whiplash and temporomandibular joint dysfunction [43–45]. Whiplash might induce joint lesions and posttraumatic malocclusions, which would lead to dysfunction of the masticatory muscle, resulting in tinnitus [46]. However, other researchers claim that temporomandibular joint dysfunction is not associated with whiplash injuries [47–49].

In short, although whiplash is considered a cervical spinal disorder, its relation with tinnitus is controversial. Furthermore, evidence of somatosensory modulation of tinnitus in such patients is not yet supported by the literature.

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References

2. Sanchez, TG; Guerra, GCY; Lorenzi, MC; Brandão, AL; Bento RF. The influence of voluntary muscle contractions upon the onset and modulation of tinnitus. Audiol. Neurootol. 2002; 7:370–5.
9. Sanchez, TG; Marcondes, RA; Kii, MA; Lima, AS; Rocha, CAB; Ono, CR; Bushpiel, C. A different case of tinnitus modulation by tactile stimuli in a patient with pulsatile tinnitus. Presented at II Meeting of Tinnitus Research Initiative, Mônaco, 2007:21–3.


31. Estola-Partanen, M. Muscular tension and tinnitus: an experimental trial of trigger point injections on tinnitus [dissertation]. Tampere: Faculty of Medicine, University of Tampere; 2000.


Keypoints

1. Epidemiologic data indicate a frequent association between temporomandibular joint disorders and tinnitus.
2. It is extremely unlikely that tinnitus is directly caused by a mechanical relationship between the masticatory system and the middle ear.
3. Increased muscle tension of masticatory muscles may cause clonus of the palatine muscle via reflex muscle hypertension.
4. Increased muscle tension of masticatory muscles can influence tinnitus via somatosensory afferents.
5. In certain patients, occlusional appliances have been shown to normalize increased muscle tension and improve tinnitus, even if evidence for their efficacy is limited.

Keywords Tinnitus • Temporomandibular joint • Temporomandibular diseases • Masticatory muscles • Bruxism

Abbreviations

CNS Central nerve system
DCN Dorsal cochlear nucleus
TMD Temporomandibular disorder
TMJ Temporomandibular joint

Introduction

Patients with tinnitus frequently complain about temporomandibular disorders (TMD). Otological symptoms in connection with TMD have been widely described in the literature [1–4]. Some studies even reported that TMD treatment can successfully alleviate or cure tinnitus symptoms [5, 6], suggesting a potential causal relationship between TMD and tinnitus.

Disorders of the masticatory system may exert an influence on tinnitus via mechanical connections between the temporomandibular system and the ear or via neuronal influences. This chapter discusses the possible relationship between tinnitus and the masticatory system.

Temporomandibular Joint Disorders and Middle Ear Function

The structures of the middle ear and the temporomandibular joint derive from the first brachial arch. All muscle of the masticatory system and the tensor tympani muscle and the tensor veli palatine muscle are innervated by branches of the trigeminal nerve. In contrast, the M. stapedius, the stapes, the mimic muscles, and the sternocleidomastoid muscle derive from the second branchial arch. These muscles are innervated by the facial nerve.

The ontogenetic development of the masticatory system and the middle ear associates a close anatomical relationship (Fig. 44.1). There is a fibrous connection between the discal apparatus of the temporomandibular joint (TMJ) and the malleus of the middle ear [7, 8]. By this connection, pathologies of the TMJ may theoretically...
cause dysfunction of the middle ear, which in turn could cause tinnitus. However, recent anatomical studies have found no evidence for the hypothesis that traction of the discomalleolar ligament may trigger movement in the middle ear ossicles [7–9]. Also, a possible influence on middle ear ossicles via the sphenomandibular ligament has been discussed, but this may only be relevant in some rare cases of surgical manipulation [8] or traumatic mandibular dislocation [9, 10].

In summary, little evidence exists that an anterior dislocation of the articular disc (internal derangement) causes tinnitus via a mechanical influence on middle ear function, since such symptoms may only be provoked by an extensive protruded dislocation of the mandible.

**Masticatory Muscle and Muscles of the Middle Ear**

Muscles of the masticatory and the middle ear system (M. tensor veli palatini, M. tensor tympani) are both innervated by the trigeminus nerve. Neuromuscular dysfunction of the masticatory muscles may induce a “reflex hypertonia” of the tensor muscles of the middle ear [11]. The irregular tonus of the tensor veli palatini may result in a dysfunction of the Eustachian tube, which can result in aural congestion and tinnitus [12, 13] (Fig. 44.2). Even if there are conflicting views about the exact role of the tensor veli palatini, the levator veli palatini, and the tensor tympani muscle in Eustachian tube dysfunction, in the rare cases of tinnitus due to palatinal myoclonus, a potential influence of masticatory muscles should be considered.

Removable occlusal appliances made of resin have been recommended to relax masticatory muscles to eliminate or alleviate hearing symptoms triggered by TMJ disorders. However, up to now, occlusal appliances have not been proven to reduce or eliminate masticatory muscle dysfunction such as bruxism (grinding of the teeth). Nevertheless, some authors believe that a “perfect” reconstruction of the occlusal contacts of mandibular teeth, in association with maxillary teeth, may cure or at least alleviate masticatory muscle spasms [14–16]. Other authors argue that changing the occlusal relationship between the upper and lower jaw is not effective for reducing muscle hyperactivity [17–19]. A review of the Cochrane Collaboration concluded “the evidence is insufficient for affirming that the occlusal splint is effective for treating sleep bruxism” [20]. So, it is questionable whether occlusal appliances may have a beneficial influence on dysfunction of the tensor veli palatini or the tensor tympani muscles if there is no evidence for an effect on masticatory muscle function.

However, even if only a little evidence exists that the treatment of masticatory muscle hyperactivity has an

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**Fig. 44.1** Discomalleolar and sphenomandibular ligament. Cranial and lateral view

1 = Discus articularis  
2 = Ligamentum malleomandibulare  
3 = Ligamentum sphenomandibulare  
4 = Malleus  
5 = Os tympanon  
6 = Chorda tympani
impact on the tensor veli palatini, the use of occlusional appliances should be considered as a potential treatment approach in patients who suffer from tinnitus due to palatinal myoclonus.

**Somatosensory Influences on the Central Auditory System**

Many individuals can modulate their tinnitus by movements of the head and neck. The contraction of facial mimic muscles [21], as well as also clenching the jaw or moving the head or neck, can cause change in tinnitus. The most common change is an increase or decrease in loudness [22, 23].

Afferent nerves from head and neck muscles, particularly the second cervical nerve and the trigeminal nerve or ganglion, are known to interact with central auditory pathways in the dorsal cochlear nucleus (DCN) (see Chap. 9) [24–31]. Recent studies have attributed the DCN as a key role in the development of tinnitus [30, 31]. The trigeminal input to the DCN may suppress neural activity generated by sounds produced by chewing, self-vocalization, or respiration. Input to the DCN from the pinna area may serve localization of
sounds in animals that can move their pinna. It has been hypothesized that the DCN is also involved in movement programs, which orient the head and body toward the source of the perceived sound [27]. In studies on cats, pressure or stretching the head and neck structures evokes inhibitory and excitatory responses in neurons of the DCN with a predominance of inhibitory responses [27]. In this way, somatosensory input to the DCN may modulate excitability in central auditory pathways and it may cause aberrant neuronal activity in central auditory pathways, which may be perceived as tinnitus (see Chap. 10).

If tinnitus can be modulated by manipulation of somatosensory afferents from head and neck structures, dentists may contribute to the treatment of tinnitus. It has been emphasized that the position of the mandible and the hyoid may influence the posture of the head and the cervical spine, or vice versa [32]. Occlusal changes during dental treatment may alter the resting position of the hyoid and the mandible. If the occlusion contacts in the resting position shift the mandible into a protruded, retruded, or laterally displaced position, a muscular imbalance may occur, not only in the chewing muscles but, also in the muscles of the cervical spine (Fig. 44.3). Thus, imbalance of the occlusal system may result in abnormal somatosensory input to the DCN from the second cervical nerve and the trigeminal, which in turn modulates neuronal activity in other parts of the central auditory pathways. This mechanism has been studied in detail in animal experiments [25] and is assumed to account for the association between head and neck pathologies and tinnitus. However, it is controversial to which extent can occlusal equilibration or the use of occlusal appliances contribute to normalization of masticatory and neck muscle tension [18, 33].

Regarding indications for interventions such as those of occlusal appliances, it should also be considered that temporomandibular dysfunction may have existed for a long time before the start of tinnitus, and the condition may even be congenital. The association between craniofacial anomalies, malocclusion, and TMJ disorders has been widely investigated [34–38]. Despite the conflicting evidence in the literature, at

**Fig. 44.3** Association between the posture of the mandible and the surrounding structures of the hyoid and cervical spine area

1 = M. temporalis  
2 = Deep occipital muscles  
3 = Superficial occipital muscles  
4 = Infrahyoidal muscles  
5 = Suprahyoidal muslce  
6 = Cervical spine  
7 = Shoulder
least one tendency has been recognized, namely that
tenderness of the masticatory muscles may be associ-
ated with shorter posterior facial height and a shorter
mandible. Increased tenderness of shoulder muscles
was found in individuals with larger cranial base
angles, reduced mandibular prognathism, and larger
inclination and vertical jaw relationship [36]. The ten-
derness of these muscles may stimulate somatosensory
input through the second cervical spinal root and may
thus cause or affect tinnitus.

Despite the limited evidence about the exact nature
of the association between TMJ disorders and tinnitus,
patients who perceive their tinnitus differently by mov-
ing their mandible, head, or neck should be examined
by a dentist [39]. If a change in the mandibular resting
position by dental treatment has a beneficial effect on
a patient’s tinnitus, occlusal appliances or orthodontic
-treatment should be considered for alleviating the
-tinnitus.

References

Otological manifestations in temporomandibular joint dys-
development of the human lateral pterygoid muscle and its
relationships with the temporomandibular joint disc and
Recurrent tinnitus and associated ear symptoms in adults. Int
J Audiol 44:164–70
review: temporomandibular disorders in an integral otic
5. Gelb H, Calderone JP, Gross SM, Kantor ME (1967) The
role of the dentist and the otolaryngologist in evaluating
temporomandibular joint syndromes. J Prosthet Dent
18:497–503
associated with the temporomandibular joint syndrome. Eur
Arch Otorhinolaryngol 249:93–4
Discomalleolar ligament in the adult human. Cranio
4:299–305
and anterior malleolar ligaments: possible causes of middle
ear damage during temporomandibular joint surgery. Oral
9. Sencimen M, Yalcin B, Dogan N, Varol A, Okcu KM, Ozan
ments between the malleus and the temporomandibular
Discomallear and malleomandibular ligaments: anatomical
study and clinical applications. Surg Radiol Anat 25:152–7
Management of Head, Neck and TMJ Pain Dysfunction.
12. Malkin DP (1987) The role of TMJ dysfunction in the etiol-
ogy of middle ear disease. Int J Orthod 25:20–1
13. Zipfel TE, Kaza SR, Greene JS (2000) Middle-ear myo-
clonus. J Laryngol Otol 114:207–9
14. Ramfjord SP (1961) Bruxism, a clinical and electromyo-
15. Dawson PE (1973) Temporomandibular joint pain-dysfunction
problems can be solved. J Prosthet Dent 29:100–12
dysfunction syndrome with occlusal equilibration. J Prosthet
Dent 63:695–700
of biofeedback and occlusal adjustment on bruxism. J Periodontol
49:367–72
therapy: occlusal adjustment procedures. J Am Dent Assoc
110:743–50
tion splints for the management of patients with masticatory
Investig 8:179–95
GF (2007) Occlusal splints for treating sleep bruxism (tooth
grinding). Cochrane Database Syst Rev 4:CD005514
or hearing loss elicited by facial mimetic movement.
Laryngoscope 95:966–70
22. Levine RA (1999) Somatic (craniocervical) tinnitus and the
dorsal cochlear nucleus hypothesis. Am J Otolaryngol
20:351–62
N Engl J Med 347:904–10
ganglion innervates the auditory brainstem. J Comp Neurol
419:271–85
underlying somatic tinnitus. Prog Brain Res 166:107–23
cochlear nucleus: unit responses to acoustic and trigeminal
27. Kanold PO, Young ED (2001) Proprioceptive information
from the pinna provides somatosensory input to cat dorsal
Res 157:365–72
29. Kaltenbach JA (2000) Neurophysiologic mechanisms of tin-
30. Kaltenbach JA (2007) The dorsal cochlear nucleus as a con-
tributor to tinnitus: mechanisms underlying the induction of
hyperactivity. Prog Brain Res 166:89–106
31. Kaltenbach JA (2006) The dorsal cochlear nucleus as a par-
ticipant in the auditory, attentional and emotional compo-
32. Rocabado M (1983) Biomechanical relationship of the cranial,
cervical, and hyoid regions. J Craniomandibul Pract 1:61–6
Part IV
Differential Diagnosis of Tinnitus
Chapter 45  
Introduction

Berthold Langguth

Keypoints

1. Tinnitus is not a single clinical or pathophysiologic entity. There are many forms of tinnitus that differ in their pathophysiology.
2. Exact diagnosis is required in each patient in order to provide the best management of tinnitus.
3. It is especially important to identify those patients who can be treated by specific interventions and those in which tinnitus is a symptom of a severe underlying disease and those patients who require immediate therapeutic action.
4. Exact diagnosis is also of great importance in clinical trials.
5. In the future, new methods such as functional neuroimaging may be found to have additional diagnostic value.

Keywords  Tinnitus • Diagnosis • Heterogeneity • Pathophysiology • Diagnostic algorithm

Tinnitus can be experienced as a ringing, roaring, clicking, hissing, or buzzing. Tinnitus can start together with hearing loss but can also occur after neck trauma or during stressful live events. In some individuals, tinnitus is accompanied by insomnia, others have difficulty in concentrating, and some complaint about hyperacusis. Some individuals report that their tinnitus worsens by environmental sound; in others, the same sound may relieve their tinnitus. These clinical observations clearly show that tinnitus is not a single disease entity, but that there are many different forms of tinnitus that are likely to vary in their pathophysiology and in their response to treatment interventions. This, in turn, implies that an exact differential diagnosis is of utmost importance in the management of tinnitus.

This insightful view on tinnitus is not new. Already, more than 200 years ago (coupled with the systematic application of specific therapeutic interventions), diagnostic criteria for tinnitus were developed. The goal at that time was to identify patients who responded to galvanism, which was the then available therapy (Fig. 45.1, [1]).

It is assumed that the exact pathophysiological changes in an individual determine the efficacy of specific causally oriented therapies. In contrast, the mechanisms involved in generating the sensation of a sound when no sound reaches the ear may be less relevant for therapeutic methods that aim at habituation to the sound, such as tinnitus retraining therapy or cognitive behavior therapy. Hence, the increasing popularity of these methods in the last several decades has shifted the diagnostic focus. Clinical characteristics of the sound a person perceives with a potential reflection of its generating mechanism, such as sound characteristics, laterality, or duration, have been considered as less important. Instead, the interest has focused on detailed information about how the tinnitus impairs an individual’s life and its psychosocial consequences.

B. Langguth
Department of Psychiatry and Psychotherapy, University of Regensburg, Universitätsstraße 84, 93053 Regensburg, Germany
and
Interdisciplinary Tinnitus Clinic, University of Regensburg, Universitätsstraße 84, 93053 Regensburg, Germany
e-mail: Berthold.Langguth@medbo.de

the first symptom of potentially dangerous diseases, some of which may even become life threatening if left undiagnosed and untreated (e.g., carotid dissection and vestibular schwannoma). Therefore, each patient with tinnitus requires a careful and systematic diagnostic approach.

In this section (Part. III), a diagnostic algorithm (Chap. 46) will first be presented, which provides guidance for systematic diagnosis of clinically relevant and specific forms of tinnitus. The diagnostic steps, which are recommended in all patients, include a detailed case history (Chap. 47) and otological (Chap. 48) and audiological examinations (Chap. 49). Depending on the findings in these primary diagnostic procedures, further diagnostic steps for exactly diagnosing specific subforms of tinnitus may or may not be required. Indications for further diagnostic steps are, for example, acute tinnitus, pulsatile tinnitus, or severe general impairment of the individual.

Chapter 46, “Diagnostic Algorithm,” will give a synoptic overview about the diagnostic process and provide orientation of which diagnostic procedures are indicated in which case. These procedures are then described in detail in Chaps. (49) Neurotologic Assessment, (50) Neurologic Examination, (52) Diagnosis of Somatosensory Tinnitus, (53) TMJ Assessment, and (54) Psychological/Psychiatric Assessment. Parts II (causes of tinnitus) and IV (clinical characteristics of the different forms of tinnitus) concern these specific forms of tinnitus and their management. It should be noted that the proposed diagnostic approach refers mainly to the identification of currently known subforms of tinnitus with a well-understood pathophysiologic mechanism that also holds therapeutic relevance, such as tinnitus together with sudden hearing loss, or pulsatile tinnitus associated with a neurovascular conflict.

However, the frequently observed high variability in treatment outcome in clinical trials [4, 5] suggests the existence of further subforms of tinnitus, the specific clinical characteristics of which we do not yet know and for which our knowledge of the exact pathophysiologic underpinnings is still incomplete. This, in turn, may result in a vicious cycle: it is difficult to identify new promising treatments if we do not know according to which criteria tinnitus patients should be stratified. However, as long as no...
effective treatments are available, it is difficult to identify clinically relevant criteria for stratification. Different strategies may help overcome this problem. First, standardized assessment of clinical characteristics in clinical trials will provide the opportunity to identify clinical characteristics that predict responses to specific interventions. For this purpose, an effort has been made at the TRI meeting in Regensburg 2007 to arrive at a consensus about such a standard (http://www.tinnitusresearch.org; [6]). Also, the advent of new techniques such as functional neuroimaging or transcranial magnetic stimulation has shown some promise for better diagnosis of the different forms of tinnitus. Recent findings using these techniques suggest that clinical criteria such as tinnitus duration [7] or sound characteristics (pure tone vs. narrow band noise [8]) may have specific pathophysiologic reverberations and therefore seem to be relevant criteria for stratifying patients with tinnitus.

References

Chapter 46
Algorithm for the Diagnostic and Therapeutic Management of Tinnitus

Berthold Langguth, Ebergard Biesinger, Luca Del Bo, Dirk De Ridder, Ron Goodey, Carlos Herraiz, Tobias Kleinjung, Miguel J.A. Lainez, Michael Landgrebe, Michel Paolino, Benjamin Questier, Tanit G. Sanchez, and Grant D. Searchfield

Keypoints

1. Tinnitus can be a symptom of a wide range of different underlying pathologies and accompanied by many different comorbidities, indicating the need for comprehensive multidisciplinary diagnostic assessment.

2. Basic diagnostics should include a detailed case history, assessment of tinnitus severity, clinical ear examination, and audiological measurement of hearing function. For a considerable number of patients, these first diagnostic steps in combination with counseling will be sufficient.

3. Further diagnostic steps are indicated if the findings of basic diagnostics point to acute tinnitus onset, a potentially dangerous underlying condition (e.g., carotid dissection), a possible causal treatment option, or relevant subjective impairment.

4. Further diagnostic management should be guided by clinical features. There is increasing evidence that phenomenologic and etiologic aspects determine the pathophysiology and the clinical course of tinnitus. In a hierarchical diagnostic algorithm, the first differentiation should be between pulsatile vs. non-pulsatile tinnitus. In case of non-pulsatile tinnitus, differentiation between acute tinnitus with hearing loss, paroxysmal tinnitus, and chronic tinnitus is recommended. Further diagnostic procedures of constant non-pulsatile tinnitus will depend on concomitant symptoms and etiological conditions.

5. All diagnostic and therapeutic steps should be accompanied by empathic and insightful counseling.

6. The ultimate treatment goal is the complete relief from tinnitus. If causally oriented treatment options are available, these should be preferred. However, in many cases, only symptomatic therapies can be offered, and then the treatment goal in clinical practice will be defined as the best possible reduction of unpleasant hearing sensations and accompanying symptoms, that is, to improve quality of life.

Keywords Tinnitus • Pathology • Etiology • Comorbidity • Symptom • Diagnosis • Therapy

Abbreviations

AVM Arterio-venous malformation
ABR Auditory brainstem responses
BIH Benign intracranial hypertension
CBT Cognitive behavioral therapy
CHQ Case history questionnaire
CSF Cerebrospinal fluid
CT Computer tomography
ECoG Electrocochleography
EEG Electroencephalography
FDA US Food and Drug Administration
LP Lumbar puncture
MRI Magnetic resonance imaging
MVC Microvascular compression
OAЕ Otoacoustic emissions
PTSD Posttraumatic stress disorder
SOL Space occupying lesion
THI Tinnitus handicap inventory
TMJ Temporomandibular joint

B. Langguth (✉)
Department of Psychiatry and Psychotherapy, University of Regensburg,Universitätsstraße 84, 93053 Regensburg, Germany
and
Interdisciplinary Tinnitus Clinic, University of Regensburg, Regensburg, Germany
e-mail: Berthold.Langguth@medbo.de
Introduction

Diagnostic and therapeutic management of tinnitus is challenging for a variety of reasons. Multiple etiologies can result in the same phantom sound percept. Even though hearing disorders are the most important risk factors for the development of tinnitus, other diseases such as brain tumors, neck injuries, temporomandibular dysfunction, or emotional disorders generally covered by other disciplines (e.g., neurology, psychiatry, orthopaedics, dentistry, or neurosurgery) can be critically involved in the etiology or continuation of tinnitus. Therefore, the requirements of comprehensive tinnitus diagnosis and treatment can only be met by an integrated multidisciplinary approach.

Although tinnitus itself is not dangerous, it can be the first sign of potentially dangerous diseases that can even become life threatening if left undiagnosed and untreated. Furthermore, tinnitus by itself may become life threatening if accompanied by suicidal tendencies.

The authors1 of this chapter developed an algorithm in order to provide guidance for diagnosis and treatment of tinnitus based on currently available evidence (see Fig. 46.1).2 In particular, this algorithm is intended to assist clinicians who occasionally see tinnitus patients and may not be fully aware of the complexity of the condition. Subgroups of tinnitus require specific management or can benefit from specific treatment. Even if some of these conditions are relatively rare, considering the possibility of their occurrence is warranted because of the availability of specific treatment options.

A stepwise decision-tree approach for tinnitus management is proposed, starting with basic diagnostic steps, which are recommended in all patients [1] (Fig. 46.1, white boxes), and includes history taking for associated symptoms (Fig. 46.1, red boxes). Depending on the findings of the first step, further diagnostic or therapeutic measures may or may not be necessary (see also Table 46.1). The second step consists of tailored technical tests (Fig. 46.1, yellow boxes) for the diagnosis of specific tinnitus-related disorders (blue boxes), leading to a causal and therapeutic management. For cases in which a causally oriented treatment is not available, not possible, or not sufficiently successful, symptomatic treatment options can be offered (Fig. 46.1, grey boxes). It is emphasized that the entire diagnostic workup should be accompanied by empathic and insightful counseling (Chap. 70).

More detailed descriptions of the different diagnostic and therapeutic steps can be found in the specific book chapters in Part II (Causes of Tinnitus), Part III (Differential Diagnosis of Tinnitus), and Part IV (Clinical Characteristics of Different Forms of Tinnitus).

Basic Diagnostic Assessment

This first step, to be performed in every patient and not requiring any sophisticated instrumentation, will reveal enough clinical information about tinnitus, hearing, and comorbidities to decide whether further diagnostic steps are needed – if yes, the diagnostic assessment indicates which of them would be most appropriate. These basic diagnostics should include an in-depth case history including assessment of tinnitus severity (using screening tools or questionnaires) (for details see Chap. 47); clinical ear examination (for details, see Chap. 48); and audiological measurement [1] (for details, see Chap. 49).

As previously discussed, for a considerable number of patients, these first diagnostic steps in combination with counseling will be sufficient (see Table 46.1). For example, further diagnostic steps are not necessary if there is no hint of a dangerous underlying disease and

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1The authors are clinicians from different specialties who all have extensive experience in the management of tinnitus patients. The algorithm for the diagnostic and therapeutic management of tinnitus patients has been developed in the framework of the Tinnitus Research Initiative (http://www.tinnituresearch.org). In order to avoid bias due to specific disciplines or specific health care systems, the group was multidisciplinary (consisting of otolaryngologists, audiologists, neurologists, psychiatrists, psychotherapists, and a neurosurgeon) and multinational (Belgium, Brazil, France, Germany, Italy, New Zealand, and Spain).

2In the original powerpoint version of the flowchart (available at http://www.tinnituresearch.org), a mouse click on any item opens a separate slide, which provides more detailed additional information. In order to avoid redundancy, only the synopsis is presented in this chapter.
Algorithm for the Diagnostic and Therapeutic Management of Tinnitus

If the patient is not impaired by the tinnitus. However, the basic assessment will also identify those cases where further diagnostic and/or therapeutic steps are required and those where further steps should be performed immediately.

An example for a life-threatening emergency is concomitant suicidal tendencies, which require immediate therapeutic action (see Chap. 54). Other conditions, such as acute tinnitus in combination with sudden hearing loss of recent onset, should also be treated as soon as possible (see Chap. 56). In conditions where a severe underlying disease cannot be excluded or when a patient is bothered by his tinnitus, further diagnostic steps are clearly indicated.

Further diagnostic procedures should be guided by clinical features (Fig. 46.1, red boxes). In a hierarchical diagnostic algorithm, the first differentiation should be between pulsatile vs. non-pulsatile tinnitus. This differentiation acknowledges the fundamental pathophysiological difference between these two forms of tinnitus.
Pulsatile Tinnitus

It is important to note that pulsatile refers to heart synchronous or respiration synchronous (venous hum) pulsatile tinnitus. Pulsatile tinnitus with a rhythm different from the heart rate or respiration rate should be classified here as pseudopulsatile or non-pulsatile tinnitus (see Chap. 59). Further diagnostic assessment of heart synchronous pulsatile tinnitus requires a neurovascular examination (yellow box). Diseases such as arteriovenous malformation, sinus venous thrombosis, benign intracranial hypertension, or high jugular bulb may be identified as potential causes of pulsatile tinnitus. A detailed description of the diagnostic management of pulsatile tinnitus is found in the chapter on pulsatile tinnitus.

Non-pulsatile Tinnitus

Non-pulsatile tinnitus is much more common than pulsatile tinnitus and requires further differentiation according to chronicity, concomitant symptoms, and etiologic factors. As a first step, a differentiation between acute tinnitus with sudden hearing loss, paroxysmal tinnitus, and chronic tinnitus can be useful. In case of acute tinnitus accompanied by sudden hearing loss, diagnostic and therapeutic procedures should not be postponed in order to identify possible causes for the hearing loss and start appropriate treatment. This is described in detail in the chapter on sudden hearing loss and tinnitus.

Paroxysmal or intermittent tinnitus can be a symptom of auditory nerve compression, superior canal dehiscence syndrome, Ménière’s disease, palatal myoclonus, or even epilepsy (blue boxes). For a differential diagnosis, magnetic resonance imaging (MRI), auditory-evoked potentials, and electroencephalography (EEG) are indicated (yellow boxes) (for more details, see Chap. 58).

Further diagnostic procedures of constant non-pulsatile tinnitus will depend on concomitant symptoms and etiologic conditions (Fig. 46.1, red boxes). Constant non-pulsatile tinnitus can be accompanied by conductive or sensorineural hearing loss. Conductive hearing loss can be caused by otosclerosis, different forms of otitis, or Eustachian tube dysfunction. More information about tinnitus with conductive hearing loss is given in Chap. 34.

In case of sensorineural hearing loss, further diagnostic procedures are indicated for identifying the exact etiology. These can include magnetic resonance imaging (MRI) and auditory brainstem responses (ABR) (e.g., for excluding vestibular schwannoma) and also otoacoustic emissions for assessment of outer hair cell function (detailed description in Chaps. 35 and 36).

Diagnostic assessment and therapeutic management of tinnitus occurring together with vertigo is indicative of specific pathologies such as Ménière’s disease, superior canal dehiscence, or damage to the vestibulocochlear system. More details are found in Chaps. 36, 38–40, 60, and 84.

If tinnitus presents with associated headache, increased intracranial pressure has to be excluded. Potential underlying pathologies such as space occupying lesions (SOL), benign intracranial hypertension (BIH), disorders of cerebrospinal fluid (CSF) circulation, or craniocervical anomalies can be diagnosed by MRI. In specific cases, lumbar puncture and furosemide tests may help determine whether reduced CSF pressure also alleviates tinnitus (more details are found in Chap. 61).

The co-occurrence of depression, anxiety, and insomnia with severe tinnitus has been frequently described. Immediate action is required when a patient reports acute suicidal thoughts. A detailed explanation of diagnostic procedures in case of psychiatric comorbidity is provided in Chap. 58, 63–65.

When tinnitus is associated with neck or temporomandibular dysfunction or pain, a more detailed examination of these systems should be considered. Radiologic tests are indicated if structural alterations are suspected, whereas functional impairments can be best detected by physical examination performed by experienced dentists and physiotherapists. More details are presented in Chap. 43, 44, 52, 53, and 95.

Specific diagnostic tests are indicated if tinnitus begins or worsens within 3 months after a traumatic event. It is important to note that a delay of several weeks between trauma and tinnitus onset does not exclude a potential etiologic relationship. Traumatic events may cause tinnitus in different ways. The indication for further diagnostic procedures depends on the trauma mechanism. In particular, noise, ear, head, neck, and even emotional trauma should be considered. In case of posttraumatic pulsatile tinnitus, immediate
Algorithm for the Diagnostic and Therapeutic Management of Tinnitus

Diagnostic workup for vascular pathologies (especially carotid dissection) is mandatory. A detailed description of pathologies, which can occur as a consequence of trauma and which may be involved in the generation of tinnitus, is given in Chap. 66. A separate chapter is devoted to blast injuries (Chap. 67) since tinnitus has become one of the most relevant warfare-related health problems in the last years. Blast injuries are a particular diagnostic challenge since the tinnitus-inducing mechanisms may include noise, ear, head, neck, and emotional trauma.

Hyperacusis and phonophobia occur frequently together with tinnitus and require specific management, which is described in detail in Chaps. 3 and 4.

**Symptomatic Treatment**

Symptomatic treatment should be considered in every patient who feels impaired by his tinnitus if specific causally oriented treatments are not available, not sufficiently effective, or not indicated for any other reason. Cognitive behavioral therapy (CBT) (see Chap. 71) and auditory stimulation with counseling are the most established treatment options. Auditory stimulation can be essentially differentiated in the use of sound for masking or partially masking tinnitus (see Chaps. 74 and 75) and in attempts to compensate for hearing deficits, for example, by hearing aids (see Chaps. 74, 76, and 77). Also, specific forms of sound stimulation with a frequency composition according to the individual audiogram have been proposed (see Chap. 75). Tinnitus retraining therapy (TRT), a specific combination of auditory stimulation and counseling, is widely used and described in Chap. 73.

Pharmacotherapeutic options for the treatment of tinnitus are limited. However, even if there is currently no drug, which is approved by the US Food and Drug Administration (FDA) for the treatment of tinnitus, there are some promising results from clinical studies indicating beneficial effects of specific drugs for subgroups of patients (Chap. 78). Neuromodulatory approaches have been proposed very recently. Most evidence is available for transcranial magnetic stimulation (TMS); neuromodulation, transcranial direct current stimulation (tDCS), cutaneous stimulation, and cortical electrical stimulation have also demonstrated promising results. A description of how these techniques are performed and which results have been obtained is given in the respective Chaps. 86, 88–90.

In summary, a wide range of different pathologies can underlie tinnitus. The diagnostic challenge can best be met by a stepwise approach consisting of basic assessment procedures followed by more detailed diagnostic tests in selected patients. Here, important indications for further diagnostic steps and immediate treatment are summarized. If these diagnostic procedures do not reveal causally oriented treatment options or if results from such therapies are not satisfying, the available symptomatic treatment possibilities should be considered. It should also be mentioned that this algorithm is based on currently available knowledge and is expected to evolve and be refined with time and criticism.

**Reference**

Chapter 47
History and Questionnaires

Berthold Langguth, Grant D. Searchfield, Eberhard Biesinger, and Karoline V. Greimel

Keywords
Tinnitus • Case history • Questionnaire
• Structured interview • Quantitative assessment
• Clinical management • research applications • Visual analogue scales

Introduction

Tinnitus has many forms and many characteristics. However, tinnitus is not readily apparent to others, and currently no objective procedures are yet established for diagnosis of tinnitus. The assessment of the perceptual aspects of tinnitus is difficult. Only by listening to the patient can one find out whether a patient has tinnitus and what form of tinnitus he/she has. The case history is of high importance for correct diagnosis in all areas of medicine; this is especially true for tinnitus, since it is fundamentally a self-report phenomenon. Moreover, the subjective nature of tinnitus is a challenge not only in the clinical management of the individual with tinnitus but also for research applications.

Therefore, in addition to an otologic (see Chap. 48) and audiologic assessment (see Chap. 49), a detailed case history is required in all tinnitus patients in order to obtain the necessary information for deciding about the therapeutic management (see Chap. 46). In many patients, a comprehensive diagnostic assessment including a detailed case history can be sufficient for tinnitus management. If a severe disorder (e.g., tumor or carotid dissection) can be excluded and the tinnitus is not perceived as a problem, no treatment is necessary. In all other cases, the detailed case history represents the first therapeutic step, since the patients can make their experience known, they see that their complaints are taken seriously and that the clinician is competent, caring, and understands the effects of tinnitus.
For the clinical management of the individual patient information about the perceptual characteristics of tinnitus (e.g., pulsatile or non-pulsatile), its time course (e.g., recent onset or chronic), influencing factors (e.g., reduction by environmental sound), and associated symptoms (e.g., reduced sound tolerance) are important. These qualitative data can be best obtained by case history questionnaires or a structured interview.

Loudness of tinnitus can be evaluated quantitatively either by rating or by matching methods (see Chap. 49). In addition to details about the tinnitus percept, information about the perceived severity and the impact on an individual’s life also have to be assessed. Personality, comorbidities, or environmental circumstances contribute more to tinnitus-related distress, impairment, disability, and handicap than the perceptual characteristics of tinnitus [1, 2]. Therefore, the evaluation of tinnitus consequences on a person’s life needs to be multidimensional, taking into account psychological and social factors.

Screening tools allow an estimation of tinnitus severity based on a few questions, whereas for quantitative assessment of tinnitus severity, many psychometrically validated questionnaires are available. These questionnaires are helpful tools for quantifying disabling and handicapping effects of tinnitus, providing insight into how the tinnitus sensation generates a disability at a personal level and a handicap on the societal level. Responses on these questionnaires can be summed resulting in a total score or subscale scores. Based on the score, the tinnitus severity of an individual patient can be determined (e.g., in low, medium, moderate, or severe).

If a patient is moderately or severely impaired, additional assessment by a psychologist or a psychiatrist is frequently necessary. Psychological and psychiatric assessment involves the integration of information from multiple sources and tools, including the clinical interview, rating scales, questionnaires, and the observation of the patient’s behavior during the interview. It is not only what the patients say but also how they say it that is of relevance. Sometimes, interviews with significant others or reports from previous therapists or physicians provide further important information.

The clinician also needs to be aware that when people complaint about tinnitus, other problems may be contributing to any negative emotional state. For example, a coexistent hearing impairment or hyperacusis, balance problems, pain, anxiety, or depression may contribute to the person’s difficulties. Daily stressors or major life events may also have an impact on the person’s ability to cope with the tinnitus, and patients may attribute their feelings of depression and anxiety incorrectly to the tinnitus. An aim of the initial assessment may also be to disentangle causal connections between tinnitus distress, other stressors, and negative emotional states.

Another important area of investigation concerns the risk of suicide. Rather than avoiding asking questions about suicide, the clinician should address this issue directly. Patients may consider suicide as a means to escape from tinnitus or it may be concurrent to a depressive disorder (for more detailed information see Chap. 54). If results indicate the potential for self-harm, the clinician should manage for this or refer the patient to another specialist.

In addition to clinical applications, quantitative assessment by tinnitus questionnaires is an important tool for all kinds of different research applications.

This chapter reviews methods for obtaining qualitative and quantitative information about a condition, which is purely subjective in its nature, namely a patient’s tinnitus and its disabling and handicapping effects. It is our intent to provide a useful and practical reference for both clinicians and researchers seeking information about the availability of different methods. Also, the limitations of the different methods will be discussed in order to allow the readers to select the most appropriate method for their specific clinical or research application (see Table 47.1).

### Case History

A detailed history and primary source of descriptive data of the patient’s tinnitus or tinnitus-related conditions can be obtained through the initial intake, either

<table>
<thead>
<tr>
<th>Table 47.1</th>
<th>Questions to consider when performing an assessment</th>
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<tbody>
<tr>
<td>(1)</td>
<td>What do I want to know? &gt; Kind of information.</td>
</tr>
<tr>
<td></td>
<td>(e.g., tinnitus characteristics, tinnitus related impairments, comorbidity)</td>
</tr>
<tr>
<td>(2)</td>
<td>Why do I want to know? &gt; Reason for evaluation.</td>
</tr>
<tr>
<td></td>
<td>(e.g., for screening, treatment planning, measuring treatment outcome)</td>
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<tr>
<td>(3)</td>
<td>How can I get the information? &gt; Choice of appropriate assessment tools.</td>
</tr>
<tr>
<td></td>
<td>(e.g., interview, rating scales, questionnaires, protocols)</td>
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</table>
by a questionnaire or by structured interviews. The goal of the intake interview is to arrive at a thorough understanding of the nature of the tinnitus by exploring a broad range of inquiry, including causal, descriptive, and diagnostic variables. This information, together with otologic and audiological assessment, is the basis for further diagnostic and therapeutic management. In detail, the following areas should be explored comprehensively:

1. The history and descriptive characteristics of the tinnitus;
2. Specific behavioral, social, interpersonal, and emotional consequences of tinnitus;
3. Factors that may either exacerbate or reduce tinnitus severity;
4. Previous tinnitus treatments;
5. Relevant comorbidities.

Many practitioners prefer questionnaires. Case history questionnaires offer advantages of standardized questions to provide reliable and complete information; furthermore, they require less clinician time than interviews. Detailed patient information can be especially important in medicolegal cases.

Several case history questionnaires have been published [3–5], but many clinicians and researchers have developed their own questionnaires in which they include those questions that they consider important and relevant. In the context of a consensus workshop on tinnitus assessment in Regensburg in July 2006, an “items list” for tinnitus case history questionnaires (see Table 47.2) has been compiled including items that are common to most questionnaires in current use and are considered important by experts in the field. This list consists of 14 essential (level A) items and 21 highly desirable (level B) items. Also, a case history questionnaire has been developed, which can be used as an example of how these items might be expressed (Tinnitus Sample Case History Questionnaire (TSCHQ), available in English, French, Spanish, Italian, German, Portuguese, Flemish, and Czech languages at http://www.tinnitusresearch.org [4]).

Depending on their individual background, some clinicians will consider additional items as relevant (e.g., a clinician with physiotherapeutic experience will be interested in more detailed information about postural complaints). The item list should therefore only be considered as a core list to which individual specializations should be added. In the following description, we want to give some examples about the relevance of the proposed items for further diagnostic or therapeutic procedures. A comprehensive description of the clinical characteristics of the different forms of tinnitus is found in Part V.

### Table 47.2 “Items list” for tinnitus case history questionnaires [4]

| Items are ordered according to their level of significance: |
| Category “A” (= essential) in bold type. |

| Background |
| 1. Age. |
| 2. Gender. |
| 3. Handedness. |

| Tinnitus history |
| 5. Initial onset. Time? |
| 6. Initial onset. Mode? Gradual or abrupt? |
| 7. Initial onset. Associated events? Hearing change, Acoustic trauma, Otitis media, Head trauma, Whiplash, Dental Treatment, Stress, Other. |
| 9. Site. Right ear? Left ear? Both ears? (symmetrical?) Inside head? |
| 10. Intermittent or constant? |
| 11. Fluctuant or non-fluctuant? |
| 12. Loudness. Scale 1-100. At worst & at best? |
| 13. Quality. Own words/Give a list of choices. |
| 14. Pure tone or Noise? Uncertain/polyphonic? |
| 16. Percentage of awake time aware of tinnitus? |
| 17. Percentage of awake time annoyed by tinnitus? |
| 18. Previous tinnitus treatments (no, some, many)? |

| Modifying influences |
| 19. Natural masking? Music, everyday sounds, other sounds? |
| 20. Aggravated by loud noise? |
| 21. Altered by head and neck movement or touching of head or upper limbs (specification of the respective movements)? |
| 23. Effect of nocturnal sleep on daytime tinnitus? |
| 24. Effect of stress? |
| 25. Effect of medications? Which? |

| Related conditions |
| 26. Hearing impairment? |
| 27. Hearing aids (No, left ear, right ear, both ears; effect on tinnitus)? |
| 28. Noise annoyance or intolerance? |
| 29. Noise induced pain? |
| 30. Headaches? |
| 31. Vertigo/dizziness? |
| 32. Temporomandibular disorder? |
| 33. Neck pain? |
| 34. Other pain syndromes? |
| 35. Under treatment for psychiatric problems? |
Demographic data, such as age, are of relevance since the causes of tinnitus are different in younger and older people. In elderly people, tinnitus is frequently associated with presbycusis [6]; other causal factors such as noise exposure may be more prominent in younger patients [7]. A positive family history of tinnitus complaints can point to a genetic form of hearing loss as an underlying disorder. There is also some suggestion that genetic factors may play a role for individual susceptibility to tinnitus [8] (see Chap. 7).

The duration of tinnitus is of high relevance for further diagnostic and therapeutic management. Whereas acute tinnitus, especially with abrupt onset, may be a sign of an acute dangerous disease, this is only very rarely the case in chronic tinnitus. Also, acute tinnitus requires an entirely different therapeutic management than chronic tinnitus. The circumstances under which tinnitus started are also important (e.g., onset of tinnitus related to neck trauma needs a different diagnostic work-up than tinnitus that started during a stressful life event).

Concerning the sound characteristics that patients report, the differentiation between pulsatile and non-pulsatile tinnitus is of greatest importance. In patients who describe pulsatile sounds, particularly if synchronous with the heartbeat, vascular origin should be suspected. Pulsatile tinnitus requires specific diagnostic procedures (see Chap. 46). Low-pitched tinnitus with intermittent occurrence may be a cue for the diagnosis of Ménière’s disease. Neurophysiologic differences have been suggested for tinnitus resembling “a pure tone” and “noise,” and response to specific therapeutic procedures may depend on this distinction [9, 10].

Tinnitus loudness can be assessed with numeric rating scales or visual analogue scales and gives an estimate of the subjectively perceived loudness of the patient’s tinnitus. The percentage of time patients are aware of their tinnitus varies enormously between “sometimes in quiet environments” and “always.” Also, there is a difference between the time patients are aware and the time patients are annoyed by their tinnitus. These factors are important for determining how intrusive the tinnitus may be in a specific patient.

Factors that improve or worsen tinnitus can be important predictors for treatment success (e.g., use of a sound generator if environmental sounds reduce tinnitus). Determination of therapies that have been trialed, successfully or not, can also provide useful information as to a future treatment choice. When therapies in the past have failed, it should be asked exactly how the therapy had been performed. Possible reasons for failing could be an inadequate performance or insufficient duration of a given treatment.

There are several health disorders, which are frequently associated with tinnitus, such as hearing loss, hyperacusis, neck or temporomandibular joint disorders, vertigo, insomnia, headache, anxiety, or depression. These comorbidities may be a cause or a consequence of tinnitus. In all cases, the co-occurrence of these disorders is of relevance for the therapeutic management. Irrespective of whether there is a causal relationship or not, successful treatment of tinnitus comorbidities can improve the patient’s quality of life enormously. This, in turn, may also improve the patient’s ability to cope with tinnitus, even if perceptual characteristics remain largely unchanged (see chapters in Part V for more details).

Although case history questionnaires are useful tools for obtaining information, they should not replace a thorough clinical intake interview. However, the use of a case history questionnaire can make the intake interview more efficient by providing an opportunity to discuss relevant items in detail. Patients should be encouraged to clarify questions when they are uncertain how to answer. The discussion allows patients to also describe in their own words aspects of special importance to them. The discussion of the different items helps establish rapport between the clinician and the patient. In this context, it is always helpful to ask patients what bothers them the most about their tinnitus. This varies from patient to patient and has implications for the therapeutic management. If, for example, a patient suffers mainly from the lack of control, this can be addressed by cognitive–behavioral therapy; if the main complaint is difficulty in sleeping, the treatment of the sleeping problem should also be the main focus. Furthermore, the impact of tinnitus on the person’s work, sleep (falling asleep and staying asleep), participation in enjoyable activities, social interaction (with friends, family, and partner), and the general lifestyle has to be examined. Reactions to tinnitus can be very different, and it is the patient’s reaction to tinnitus that causes problems rather than the sound by itself. If this message reaches the patient during the intake interview, a very important first step toward treatment has been achieved.

Quantitative Assessment of Tinnitus

Many people with tinnitus are neither bothered nor concerned about their tinnitus. There is also a group of patients who see a physician only because they are
History and Questionnaires

Concerned that their tinnitus may be a sign of a serious ear or brain disease. Apart from those, all other people with tinnitus who seek medical attention are to some extent bothered by their tinnitus. However, there is a large variability in distress, ranging from those who have learned to cope but would welcome some relief from the sound, to those who have severe problems with tinnitus in their daily lives. It has been repeatedly shown that the loudness or the pitch of the tinnitus sensation does not predict suffering [1, 2, 11]. Methods that directly quantify tinnitus distress, disability, and handicap are more appropriate for assessing the amount of suffering. Screening tools allow an estimation of tinnitus severity based on a few questions, whereas for quantitative assessment of tinnitus severity several questionnaires are available.

Psychometric and Methodological Aspects

Different methodological aspects have to be considered in the use of quantitative measurement techniques.

Validity

Is there a specific questionnaire assessing disability, handicap, or coping styles? In general, the validity of an instrument is reflected by its ability to yield “truthful,” “correct,” or “real” information (see also Fig. 47.1). Validation strategies include content validity, criterion-related validity, and construct validity. Content validity demonstrates to which extent the items of the scale reflect the characteristics to be measured; criterion-related validity measures how well the instrument correlates with a “gold standard”; and construct validity reflects the degree to which an instrument purports to measure a theoretic construct of the characteristics to be assessed [12, 13].

Standardization and Norming

Can data assessed at place X at time X be compared to those at place Y and time Y?
Is there a specific score high or low as compared to most other patients?
Standardization means that data are always assessed and performed in the same standardized way. Relevant issues can be whether a questionnaire is completed as an electronic version or as a paper version, or whether it is completed before or after the first consultation. Only a standardized way of assessment allows comparison across individuals, time, and clinical settings. Norming means obtaining information about the distribution of measures in a target population in the form of means, standard deviations, or percentiles. Normative data allow placement of the score of an individual in context of a target population.

Reliability

Does the tinnitus questionnaire have high test–retest reliability and stability?
Reliability describes the precision of the instrument and includes internal consistency but also reproducibility. Internal consistency reflects the inter-item consistency of a scale or subscales. It is expected that several items

![Fig. 47.1 Validity and reliability](image)
that assess the same construct (e.g., tinnitus handicap) correlate with each other. The statistical measure of this internal consistency is Cronbach’s $\alpha$.

Reproducibility can be differentiated in short- and long-term reproducibility. Short-term reproducibility may reflect effects of day-to-day fluctuations; long-term reproducibility describes stability over longer time intervals. This is of relevance when a questionnaire is used for evaluating effects of a specific intervention. If there is a lack of knowledge about the changes in a questionnaire score over time occurring without any treatment intervention, one cannot rely on uncontrolled observations of treatment effects. Documented changes in tinnitus scores may not be due to the treatment, per se, but rather due to measurement error of the questionnaire used for assessing treatment outcome.

**Responsiveness**

Is the questionnaire sensitive for treatment-induced changes?

Responsiveness reflects the ability of a questionnaire to register changes following an individual’s response to a treatment intervention. This is especially required when an instrument is used as treatment outcome measure. This aspect of measurement instruments has also been characterized as evaluative [14, 15]. Variables that are stable over time and reflect, for example, the individual’s personality are called trait parameters, whereas variables that reflect mainly the actual condition are called state variables.

From an evaluative questionnaire, one would expect that it samples mainly state variables that are likely to change under treatment. A large amount of change-insensitive trait variables are useless for detecting treatment effects and may even obscure them. In contrast, the inclusion of trait parameters can be useful for an instrument designed for diagnostic use (e.g., for discriminating between individuals with severe vs. mild tinnitus, see Table 47.5).

Another factor related to the responsiveness of a questionnaire is the number of response options for each item. A questionnaire, which consists of items that can only be answered with two or three levels (e.g., yes and no), is, in general, less sensitive to changes than a questionnaire with five or more answer options per item.

It should be noted that the currently available tinnitus questionnaires have not been specifically designed for evaluating treatment-related changes, but most of them have been used as outcome measurers in clinical trials. New questionnaires specifically designed to evaluate treatment-related changes will emerge in the near future [15].

**Feasability**

Is the questionnaire easily applicable?

Feasibility reflects the property of an instrument to be practically applicable in a real-world context. As an example, in order to be applicable in a busy clinical practice, tinnitus questionnaires should be brief and easy to administer, understand, score, and interpret.

**Cultural and Language Bias**

Questionnaires designed and tested in one population and language are not necessarily equally applicable in another. Questionnaires developed in one culture do not necessarily measure the same factors in another, even if the language is the same [16]. Likewise, translation from one language to another can introduce changes in meaning. One way of addressing these variations is to validate the questionnaire in each language and setting. This may lead to some items from the original questionnaire being moved into a different factor or rejected as invalid. While this approach has merit in optimizing the questionnaire for a particular population, there are at least two significant downsides: (1) considerable time is required to validate the questionnaire in each setting and (2) cross-population comparisons become difficult. The latter of these two issues is most troublesome for researchers who might want to compare outcomes from two populations using the “same” questionnaire. For example, a questionnaire developed in the US but optimized for New Zealand might omit questions [16]. If two treatments are compared between these countries and found to have the same questionnaire scores, it cannot be assumed that the treatments are equally effective because they actually do not ask the same questions. On the other hand, if the original questionnaire is used in its original form in both countries, cultural idiosyncrasies mean that they still measure different factors.
This paradox is a limitation of questionnaires. Any “worldwide” standard should retain as many of its original items and factor structure when validated in different populations. Researchers should recognize the potential for population differences when using questionnaires.

**Screening of Tinnitus Severity**

In the daily routine of an audiological or otolaryngologic clinic, there is a high need for fast and reliable classification of tinnitus patients according to their severity. Those who suffer from tinnitus require an entirely different management than those who simply experience tinnitus, but are only slightly impaired by it. Here, a screening tool is presented that consists of three questions and allows screening for tinnitus severity in an objective and economic way (B-Scale; [17]; Table 47.3).

Another possibility of a single, global measure of the impact of tinnitus on individuals is the following global item [15]:

How much of a problem is your tinnitus?
- Not a problem 0
- A small problem 1
- A moderate problem 2
- A big problem 3
- A very big problem 4

<table>
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<tr>
<th>Table 47.3</th>
<th>B-scale for screening of tinnitus severity [17]</th>
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<tr>
<td><strong>Grading of tinnitus impairment by asking the following three questions</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Grade I</strong></td>
<td>No impairment, Compensated Tinnitus</td>
</tr>
<tr>
<td><strong>Grade II</strong></td>
<td>Slight impairment Sometimes annoying in defined conditions, e.g. in quiet environment or in stressful situations</td>
</tr>
<tr>
<td><strong>Grade III</strong></td>
<td>Permanent annoyance with disturbances in private and professional areas</td>
</tr>
<tr>
<td><strong>Grade IV</strong></td>
<td>Severe impairment Severe disturbances in private and working life, unable to work</td>
</tr>
</tbody>
</table>
Preliminary data from the tinnitus clinic in Regensburg show that this five-level response scale correlates highly with the score of the tinnitus questionnaire (see Fig. 47.2), indicating that it is a reliable, well-functioning global item for screening patients.

**Tinnitus Questionnaires**

Different questionnaires are available to assess specific aspects of patients with tinnitus (see Table 47.4). These questionnaires are score driven, which means that the responses to single items are scored and then summed or averaged. Thus, total scores or subscale scores can be calculated.

There are specific questionnaires for assessing tinnitus-related cognitions (TCQ) or coping styles (TCSQ). The other questionnaires mainly aim to quantify tinnitus distress, disability, or handicap. Items of the different scales largely overlap. Accordingly, there is a relatively high correlation between the total scores of the different tinnitus severity and handicap questionnaires.

In addition to clinical applications, quantitative assessment by tinnitus questionnaires allows research applications. For example, tinnitus questionnaires have been successfully used for investigating the relationship between personality and psychopathology and their impact on tinnitus severity [18]. Furthermore, tinnitus questionnaires provide a method for researchers to quantify tinnitus severity as a criterion for subject selection. That is, self-report measures allow investigators to select only those patients indicating a certain degree of tinnitus severity to be included in a particular study. Minimum and maximum values of the individual’s tinnitus score are defined in the research protocol as inclusion criterion, if it is expected that the treatment under study shows best effects in patients with specific tinnitus severity (see Chap. 22). Furthermore, self-report measures can be used to evaluate the effectiveness of a particular experimental treatment. Even if none of the currently available questionnaires has been specifically designed to be sensitive to treatment-related changes, there is general consensus that questionnaire scores are the best available measures of tinnitus consequence and should be used as primary outcome variables in randomized clinical trials for tinnitus treatment [4, 19].

Table 47.4 gives an overview about the most widely used quantitative tinnitus questionnaires in the English language. In the following discussion, each of these questionnaires is presented in detail. A description of the questionnaire is followed by a short explanation of how the questionnaire is scored and interpreted. Psychometric characteristics of the questionnaire are presented, strengths and weaknesses of the instrument are discussed, and finally it is indicated whether validated translations for the questionnaire are available.

Even if self-report questionnaires have been proven to be useful tools both in the clinical management of
History and Questionnaires

Tinnitus and in research applications, some caution is advised in their use and interpretation. First, completing a questionnaire might not only measure specific aspects of a patient’s tinnitus but also influence the patient’s tinnitus. Especially, catastrophizing statements may induce or reinforce maladaptive coping strategies (e.g., statements about suicide). Also, statements such as, “I cannot sleep because of my tinnitus,” may induce incorrect attributions. A patient, who repeatedly reads such a statement, may become convinced that his insomnia may be caused by his tinnitus, which is not necessarily true and might result in incorrect beliefs such as, “as long as I have my tinnitus I will never be able to sleep well.”

Second, it should be considered that self-report questionnaires can be subject to dissimulation or aggravation. Thus, just as a low score cannot exclude a significant impact of the tinnitus on a patient’s life, a high score is not proof of severe suffering. Therefore, questionnaire results always have to be evaluated in the context of the clinical impression and the patient’s tinnitus-related behavior.

### Tinnitus Severity Scale (TSS) [20]

The tinnitus severity scale (TSS) aims at quantifying individuals’ cognitive and behavioral responses to tinnitus. The 15 items are categorized under the factors intrusiveness (four items), distress (six items), hearing loss (three items), sleep disturbance (one item), and medication (one item). Responses refer to the past week and range in score from 1 (no impact) to 4 (most impact). Each item is weighted from 1 to 3 points (total weight score = 39 points). The total score is calculated by multiplying each item’s score by its weight and summing these products, resulting in a range between

<table>
<thead>
<tr>
<th>Questionnaire</th>
<th>Number of items</th>
<th>Response options for each item</th>
<th>Factors</th>
</tr>
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<tbody>
<tr>
<td>Tinnitus Severity Scale [20]</td>
<td>15</td>
<td>Four levels: wording of response options varies between item</td>
<td>One-dimensional</td>
</tr>
<tr>
<td>Subjective Tinnitus Severity Scale [21]</td>
<td>16</td>
<td>Two levels: yes and no</td>
<td>One-dimensional</td>
</tr>
<tr>
<td>Tinnitus Questionnaire [24, 25]</td>
<td>52</td>
<td>Three levels: true, partly true, not true</td>
<td>Five factors (1 – emotional distress, 2 – auditory perceptual difficulties, 3 – intrusiveness, 4 – sleep disturbances, 5 – somatic complaints)</td>
</tr>
<tr>
<td>Tinnitus Handicap/Support Scale [36]</td>
<td>28</td>
<td>Five levels: 1 = strongly disagree; 5 = strongly agree</td>
<td>Three factors (1 – perceived attitudes, 2 – social support, 3 – disability/handicap)</td>
</tr>
<tr>
<td>Tinnitus Handicap Questionnaire [27]</td>
<td>27</td>
<td>100 levels: 0 = strongly disagree; 100 = strongly agree</td>
<td>Three factors (1 – physical, emotional, social consequences of tinnitus, 2 – interfering effects of tinnitus on the hearing ability of the patient, 3 – the patient’s view of tinnitus)</td>
</tr>
<tr>
<td>Tinnitus Handicap Inventory [42]</td>
<td>25</td>
<td>Three levels: 4 = yes, 2 = sometimes, 0 = no</td>
<td>Three factors (1 – functional, 2 – emotional, 3 – catastrophic)</td>
</tr>
<tr>
<td>Tinnitus Reaction Questionnaire [29]</td>
<td>26</td>
<td>Five levels: 0 = not at all; 4 = almost all of the time</td>
<td>Four factors (1 – general distress, 2 – interference, 3 – severity, 4 – avoidance)</td>
</tr>
<tr>
<td>Tinnitus Cognitions Questionnaire [61]</td>
<td>26</td>
<td>Five levels; negative items: never = 0, frequently = 4, positive items: never = 4, frequently = 0</td>
<td>Three factors (1 – positive evaluation of tinnitus, 2 – hoplessness/despair, 3 – helplessness/victimization)</td>
</tr>
<tr>
<td>Tinnitus Coping Style Questionnaire [62]</td>
<td>33</td>
<td>Seven levels: never = 1; always = 7</td>
<td>Two factors (1 – effective coping, 2 – maladaptive coping)</td>
</tr>
<tr>
<td>Tinnitus Severity Index [54]</td>
<td>12</td>
<td>Version 1: three and four levels Version 2: three to five levels</td>
<td>One-dimensional</td>
</tr>
<tr>
<td>Tinnitus Beeinträchtigungs Fragebogen (TBF-12) [60]</td>
<td>12</td>
<td>Three levels: 4 = yes, 2 = sometimes, 0 = no</td>
<td>Two factors (1 – emotional-cognitive impairment, 2 – functional-communicative impairment)</td>
</tr>
</tbody>
</table>
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39 points (39 weighting points × 1-point item score) and 156 points (39 weighting points × 4-point item score). The TSS has acceptable test–retest reliability ($r = 0.86$). No other psychometric data are available, limiting clinical and research applications. Validated translations of the English scale into other languages are not published.

**Subjective Tinnitus Severity Scale (STSS) [21]**

The 16-item subjective tinnitus severity scale (STSS) was developed to provide a simple questionnaire to assess tinnitus severity. Each question is answered with either a “yes” or a “no” response. Ten of the 16 items earn a point if the response is “yes” (e.g., “Are you almost always aware of your tinnitus?”), whereas the other six items earn a point if the response is “no” (e.g., “When you are busy, do you quite often forget about your tinnitus?”), summing up to scores between 0 and 16 with higher scores reflecting greater overall severity. A Cronbach’s $\alpha$ of 0.84 indicates high consistency reliability. The validity was established in a sample of 30 patients, where mean STSS scores were found to correlate highly with two independent clinical ratings of severity.

The STSS is extremely simple to administer and score. The lack of a classification scheme for the total score limits its diagnostic use. Furthermore, no data about test–retest reliability are available, limiting its applicability for measuring treatment outcome.

The original questionnaire is in the English language, validated translations in Dutch [22] and French [23] have been published.

**Tinnitus Questionnaire (TQ) [24]**

The 52-item Tinnitus Questionnaire (TQ) developed by Hallam and colleagues has been designed to measure several dimensions of patients’ tinnitus complaints, namely emotional distress, auditory perceptual difficulties, and sleep disturbance. Questions either relate to the “noises” in the ear as the major cause of distress or reflect lack of coping skills.

Individuals indicate their level of agreement to each statement using one of the three response alternatives: true (2 points), partly true (1 point), or not true (0 points).

![Table 47.5 Important aspects of questionnaire construction](image-url)
Affirmative responses to an item (indicated by true) are identified as complaints about tinnitus, with the exception of the items 1, 7, 32, 40, 44, and 49, which are reverse scored because they are considered positive statements. Possible scores range from 0 to 104 points, with higher scores reflecting greater tinnitus complaints.

The TQ instrument has been found to have high internal consistency reliability (Cronbach’s $\alpha = 0.91–0.95$) and high test–retest reliability ($r = 0.91–0.94$) [25, 26]. The high test–retest reliability suggests good stability over time.

High correlations were also found between the TQ and measures of tinnitus handicap (Tinnitus Handicap Questionnaire (THQ) [27], tinnitus handicap inventory (THI) [28], and tinnitus distress (TRQ) [29]. Factor analyses conducted in separate populations were consistent with the factors originally identified by Hallam and colleagues in the United Kingdom supporting the instrument’s validity.

The TQ has been found to measure a number of different dimensions of tinnitus complaints and is a stable measure over time. In this connection, the TQ would be useful as an outcome measure in determining the effectiveness of treatment. However, the responsiveness of the TQ to changes has not been evaluated, and no data are available to assist the clinician in determining what is considered a statistically significant or clinically relevant change in scores following intervention for a given patient.

The TQ has been translated into the German language and extensively validated [30–32]. Factor analysis of the German translation of the TQ revealed that the dimensions of emotional and cognitive distress, intrusiveness, auditory perceptual difficulties, sleep disturbances, and somatic complaints can be differentiated [32]. This validation also resulted in a different scoring system, where some items were not used at all and others loaded in two factors, resulting in a maximum score of 84 points. Further translations in Dutch and French are based on the German version [33]. Recently, a Chinese version of the TQ has been validated [34]. Also, a short version of the TQ has been presented in the German language (Mini TQ, [31]), which has also been validated in Portuguese [35]. Furthermore, official translations of the Mini TQ in most European languages are available at http://www.eutinnitus.com/country-selection.php.

### Tinnitus Handicap/Support Scale (TH/SS) [36]

The 28-item tinnitus handicap/support scale (TH/SS) assesses the attitudes of significant others toward the person with tinnitus. Three factors were identified, including perceived attitudes or reactions of others (factor 1; 9 items), social support (factor 2; 10 items), and personal and social handicaps (factor 3; 9 items). Each statement on the scale is scored from 1 (strongly disagree) to 5 (strongly agree). The reliability of the TH/SS has not been examined. Construct validity of the TH/SS was assessed using a 10-item Tinnitus Severity Questionnaire (TSQ). This is the only questionnaire that has been designed to assess the influence of significant others in the overall management process, which can be helpful for counseling. The lack of retest reliability data limits both its clinical and its scientific use.

The questionnaire is in the English language and has not been validated in any other language.

### Tinnitus Handicap Questionnaire (THQ) [27]

The Tinnitus Handicap Questionnaire (THQ) description has been developed to be broad in scope but sensitive to patients’ perceived degree of tinnitus handicap. By factor analysis, three factors have been differentiated. Factor 1 (15 items) reflects the physical, emotional, and social consequences of tinnitus; factor 2 (8 items) assesses the effects of tinnitus and hearing; and factor 3 (4 items) explores the patient’s view on tinnitus.

For each item, the individual responds with a number between 0 and 100 indicating how much he or she disagrees (0 = strongly disagrees) or agrees (100 = strongly agrees) with the statement. After inverting scores obtained on items 25 and 26 by subtracting them from 100, mean scores can be calculated for the total or for each of the three factors. Higher scores indicate greater handicap.

The THQ demonstrated high internal consistency and reliability for the total scale (Cronbach’s $\alpha = 0.95$), factor 1 (0.95), and factor 2 (0.88). Factor 3 yielded a
low alpha (0.47), which may be due to the small number of items comprising this factor. A similar factor structure has been obtained in Australian [26] and New Zealand [37] samples.

Adequate construct validity of the THQ was documented by relative high correlations ($r > 0.50$) with perceived tinnitus loudness, life satisfaction, hearing threshold, depression, and general health status. High test–retest correlations have been obtained assessed over a 6-week period for the total score ($r = 0.89$), factor 1 ($r = 0.89$), and factor 2 ($r = 0.90$), whereas factor 3 yielded inadequate retest reliability ($r = 0.50$) [38]. Normative data for the THQ are available [27]. The percentile ranking allows determining severity for an individual patient relative to other patients with tinnitus. Comparison of the scores for Factor 1 (emotional and social effects) and Factor 2 (hearing) has been used to guide clinicians in treatment selection (high Factor 1, greater psychological management; high Factor 2, hearing aids [39]).

The 100-point response scale may be relatively sensitive for changes [40], but it may be somewhat problematic, especially for items, dealing with subjective strength of belief.

According to their authors, the THQ is among the most widely used questionnaires [40]. A French translation of the THQ has been validated [41]; official (unvalidated) translations in various languages are available at http://www.uihealthcare.com/depts/med/otolaryngology/clinics/tinnitus/questionnaires/index.html.

**Tinnitus Handicap Inventory (THI) [42]**

The 25-item Tinnitus Handicap Inventory consists of three subscales. The functional subscale (11 items) evaluates role limitations, the emotional subscale (nine items) reflects affective responses to tinnitus, and the catastrophic subscale (five items) probes the most severe reactions to tinnitus. However, the distinctness of the subscales has been questioned, and the use of only the total score was recommended [43].

For each item of the inventory, the patient responds with “yes” (4 points), “sometimes” (2 points), or “no” (0 points). The responses are summed, with a total score ranging from 0 to 100 points. Higher scores represent greater perceived handicap. Handicap severity categories (0–16: no; 18–36: mild; 38–56: moderate; 58–100: severe) have been developed based on quartiles calculated for the total THI score [44].

The THI has very good internal consistency reliability (Cronbach’s $\alpha = 0.93$) and high test–retest reliability for the total score ($r = 0.92$), as well as the subscales (ranging from 0.84 to 0.94). Test–retest reliability assessed on average 20 days after the initial administration was also high for the total score and the three subscales. A 95% confidence interval of 20 points for the total scale suggests that in an individual, a difference of 20 points or more between pre- and post-treatment administration can be considered statistically significant. Convergent validity was assessed using the THQ, whereas construct validity was assessed using the Beck Depression Inventory, Modified Somatic Perception Questionnaire, symptom rating scales (e.g., sleep disturbance, annoyance), and perceived tinnitus pitch and loudness. High convergent validity with the TQ has been demonstrated recently [28].

The THI is briefly and easily administered and scored. It assesses the domains of function that are addressed by many available treatment interventions.

The test–retest data allow clinicians to judge effects of treatment interventions. Further data about retest stability over longer time intervals are desirable in order to evaluate changes in perceived handicap over the medium and long term.

The THI is the most widely used tinnitus questionnaire, as evidenced by the number of citations. Validated translations are published in Danish [45], Spanish [46], Korean [47], Portuguese [48, 49], German [50], Italian [51], and Chinese [52].

In a consensus meeting, the (additional) use of the THI has been recommended for clinical studies in order to facilitate comparability between studies [4].

**Tinnitus Reaction Questionnaire (TRQ) [29]**

The Tinnitus Reaction Questionnaire has been developed for quantifying the psychological distress associated with tinnitus [29]. The 26 items of the TRQ relate to distress consequences such as anger, confusion, annoyance, helplessness, activity avoidance, and panic.
Each item on the TRQ is scored on a 5-point scale, ranging from 0 to 4 points. The scores are summed with the total score ranging from 0 to 104 points, with higher scores reflecting greater distress.

The TRQ has high internal consistency reliability (Cronbach’s $\alpha = .96$), as well as test–retest reliability ($r = .88$). Concerning construct validity, there are moderate to high correlations between the TRQ and clinician ratings and self-reported measures of anxiety and depression. A factor analysis revealed the factor’s general distress, interference, severity, and avoidance.

The TRQ represents an easy clinical tool for assessing tinnitus distress. However, no cut-off values for severity categories are available. High test–retest reliability over a period ranging from 3 days to 3 weeks indicates short-term stability of the TRQ and its usefulness in quantifying treatment outcome, at least for short interventions. However, no data are available about what is considered a statistically significant or a clinically relevant change of the score. A French translation of the TRQ has been validated [23] and compared with the English version, demonstrating only minor effects of language [53].

The Tinnitus Severity Index

The Tinnitus Severity Index is a 12-item questionnaire that measures the effect of tinnitus on work and social activities and overall quality of life [54]. The 12 items of the TSI are totaled for a single severity index. This is one of the shorter tinnitus questionnaires that has been published. There have been two versions of the TSI, the original [54] using 3- and 4-point scales and a modified version using primarily a 5-point scale, with two 4-point questions and one 3-point question [55]. The TSI has had limited use outside of the US, but the original version has been normed in New Zealand as well [37]. The TSI has good internal consistency in both US and NZ (Cronbach’s $\alpha > 0.87$) populations. The TSI has been found to correlate to the subjective rating of tinnitus loudness but not hearing loss [37]. The TSI and THQ are correlated ($r = 0.77$, $p < 0.05$), suggesting that each questionnaire is measuring similar, but not exactly the same, elements of tinnitus [37]. The TSI scores have been shown to improve following comprehensive audiology-based tinnitus management programs [56, 57], and use of SSRIs has improved scores [58]. Persons with tinnitus following head injuries have greater TSI scores than those whose tinnitus develops from other injuries [59].

The Tinnitus Handicap Questionnaire

Based on the Tinnitus Handicap Inventory (THI) [42], a short version in German language has been developed [60]. The number of items was reduced based on rigorous psychometrical testings. The final German version encompasses 12 items and distinguishes between the factors emotional cognitive (items 3, 4, 6, 8, 10, 11, and 12) and functional communicative impairments (items 1, 2, 5, 7, 9).

The internal consistency reliability of the TBF is high (Cronbach’s $\alpha = 0.90$). The TBF-12 is easy to understand and administer, psychometrically robust, and well suitable as a screening instrument in primary care. The TBF-12 is currently used as the primary outcome measure for evaluation of the efficacy of a pharmacologic compound in phase III trials. In case the trials will be positive and the compound will be approved by the Food and Drug Administration (FDA) or the European Agency for the Evaluation of Medical Products (EMEA), it will set a standard for further drugs to be approved. In the context of the phase III trial, the questionnaire has been translated and linguistically evaluated in Spanish, Dutch, French, Portuguese, Czech, Spanish, Polish, and English, as well as in African languages.

Tinnitus Cognitions Questionnaire

In contrast to the scales assessing distress, disability, or handicap, the Tinnitus Cognitions Questionnaire (TCQ) focuses on the patient’s reaction to tinnitus from a cognitive perspective. The 26 items assess positive and negative thoughts associated with tinnitus [61], especially important in the context of the psychological
management of tinnitus. The TCQ consists of 13 negative items and 13 positive items, which are clearly separated. Each of the items is rated on a five-point scale (0–4). The negative items (1–13) are scored 0–4, whereas the positive items (14–26) are reverse scored, 4–0. The addition of the item scores reveals the total TCQ score, which can range from 0 to 104, with higher scores reflecting a tendency toward more negative and less positive thoughts in response to tinnitus.

The TCQ yielded both good test–retest reliability ($r = .88$) and internal consistency reliability (Cronbach’s $\alpha = .91$). A factor analysis revealed that the negative and positive cognitions represent independent factors. Construct and convergent validity was assessed between the TCQ-total, TCQ-positive, and TCQ-negative scores, as well as other measures of tinnitus-specific symptomatology (e.g., distress, handicap, complaint behavior), depression, automatic thoughts, and loss of control. The TCQ showed moderate correlations with other tinnitus-related measures (i.e., TRQ, THQ, and TQ), with the TCQ-negative subscale demonstrating higher correlations with each of the tinnitus- and non-tinnitus measures.

The TCQ is different from other questionnaires by focusing on cognitive responses in individuals with tinnitus. The information gleaned from the TCQ responses is especially useful in the context of cognitive–behavioral therapy for screening or stratifying patients, but also for outcome measurement. However, the latter requires data about test–retest reliability. It has to be considered that reporting about cognitions or thoughts may not be identical to engaging in these thoughts. No validated translations of the instrument have been published.

**Tinnitus Coping Style Questionnaire (TCSQ) [62]**

The Tinnitus Coping Style Questionnaire (TCSQ) is a 33-item scale developed to assess adaptive and coping strategies and consists of two factors [62]. Eighteen items comprise the maladaptive coping factor; the other fifteen items comprise the effective coping subscale.

For each item, the patient indicates how frequently he/she employs each of the coping strategies on a 1 (“never”) to 7 (“always”) scale. Higher scores on the maladaptive coping subscale reflect poorer coping skills, whereas higher scores on the effective coping dimension are characterized by better acceptance of the tinnitus and use of a broad range of adaptive coping skills.

The internal consistency reliability values for the maladaptive coping and effectiveness coping subscales were 0.90 and 0.89, respectively. The two subscales were not significantly correlated ($r = 0.13$). Maladaptive coping strategies were significantly associated with measures of tinnitus severity, depression, and anxiety. In contrast, effective coping was not correlated with any of the tinnitus adjustment measures [62].

The TCSQ is specifically focused on coping strategies used by tinnitus sufferers. Information obtained with the TCSQ is fundamental in developing a cognitive–behavioral therapy program. After probing test–retest reliability of the instrument, the TSCQ might also be suitable for monitoring changes during cognitive–behavioral therapy.

**Other Questionnaires**

Beside tinnitus-related questionnaires, several other instruments referring to different comorbid conditions may be useful as part of a broad assessment of the patient and their problems. A large variety of self-report questionnaires are available for assessing depression, anxiety, sleep disorders, or health-related quality of life. A detailed description of these questionnaires is beyond the scope of this chapter. In general, these instruments are not necessary for basis assessment in every patient but may be helpful in specific cases.

**VAS Scales**

Rating scales (visual analogue scales or Likert-type scales/numerical rating scales) can be used for assessing different characteristics of tinnitus, such as loudness or annoyance. Examples for such scales are given in Fig. 47.3. Rating scales are easy to understand, but sometimes patients report difficulties, because generally the maximum end of the scale is only very vaguely defined.
Apart from loudness and annoyance, other qualitative features of tinnitus can also be easily assessed with rating scales (e.g., intrusiveness or ability to ignore tinnitus). This can provide an opportunity to understand what the most important problem is for a given patient and may lead to the use of more individually tailored assessment and monitoring tool, such as tinnitus diaries or tinnitus protocols.

A big advantage of rating scales is that they are fast to perform and can be repeated easily, e.g., in the form of tinnitus protocols. Unfortunately, there is limited psychometric data for visual analogue and numeric rating scales. One recent study shows that in individuals with tinnitus who do not seek medical attention, loudness rating scores are much lower than in those who seek help for their tinnitus [63]. It has also been shown that results of visual analogue loudness scale correlate with the THI scores ($r=0.56$) [64].

**Tinnitus Protocols**

Tinnitus protocols are self-report instruments for assessing different aspects of tinnitus over time. As an example, tinnitus loudness, annoyance, mood, and stress can be assessed daily, and results can be displayed in a diagram (see Fig. 47.4).

A tinnitus protocol can be an appropriate tool to examine changes of different tinnitus aspects over time, and correlations between intensity of tinnitus (e.g., loudness, annoyance) and different psychobehavioral factors such as mood or stress. This allows, for example, the detection of triggers or rhythmic changes over time. As shown in the example in Fig. 47.3, such a protocol can reveal that mood and stress are correlated closer to tinnitus annoyance than loudness. By monitoring different parameters over a certain period of time, the patient can learn that it is not just the noise that borders them but other emotional, cognitive, and behavioral factors that influence tinnitus perception and reaction. This can be helpful to motivate patients for cognitive–behavioral therapy.

**Conclusion**

There are different forms of tinnitus that require specific management. The intake interview is of highest importance for obtaining comprehensive information about the patient’s tinnitus in order to be able to make an exact diagnosis. Collected information from the interview, observation of the patient, and the various self-report scales should enable the clinician to formulate a view about the nature of the tinnitus, its time course, its perceptual characteristics, its comorbidities, the difficulties experienced by the patient, the person’s coping strategies, loudness the consequences...
of the tinnitus for the person’s life. Based on all these information, a specific treatment program may be developed, which is likely to be effective for an individual patient.

**References**


17. Searchfield GD, Goodey R, editors. Tinnitus Discovery: Asia Pacific Tinnitus Symposium, 2010


30. Goebel, G, Hiller, W. The tinnitus questionnaire A standard instrument for grading the degree of tinnitus. Results of a multicenter study with the tinnitus questionnaire. HNO, 1994;42:166–72


56. Folmer, RL. Long-term reductions in tinnitus severity. BMC Ear Nose Throat Disord, 2002;2(1):3
58. Folmer, RL, Shi, YB. SSRI use by tinnitus patients: interactions between depression and tinnitus severity. Ear Nose Throat J, 2004;83(2):107–8, 110, 112
59. Folmer, RL, Grist, SE. Chronic tinnitus resulting from head or neck injuries. Laryngoscope, 2003;113(5):821–7
64. Figueiredo, RR, Azevedo, AA, Oliveira, PM. Correlation analysis of the visual-analogue scale and the Tinnitus Handicap Inventory in tinnitus patients. Braz J Otorhinolaryngol, 2009; 75(1):76–9
Chapter 48
Clinical Otologic Assessment

Tobias Kleinjung

Keypoints

1. This chapter describes the diagnostic procedures of the otorhinolaryngologist, which can be performed in an office setting.
2. The otologic assessment is important for the identification of underlying causes that might be accessible to medical or surgical intervention.
3. Special attention should be paid to all kinds of objective tinnitus, which is often caused by an organic pathology of the ear or the neck.
4. The combination of otological, radiological, and audiological findings will help to make the correct diagnosis.

Keywords Tinnitus • Otoscopy • Endoscopy • Auscultation • Doppler ultrasound

Introduction

Some disorders of the conductive apparatus of the ear can cause both objective and subjective tinnitus. As already described in Chap. 23, the otorhinolaryngologist is often the first port of call for patients with new-onset tinnitus. Otologic diagnosis in patients with tinnitus should therefore concentrate on conditions that might cause tinnitus. When taking the history of a patient with tinnitus, it is essential to enquire about ear pain, sensation of aural fullness, otorrhea, sinusasal problems, hearing loss, and a history of previous infection or surgery involving the ear. Methods used in otological examination of a patient with tinnitus include tympanic microscopy, endoscopy of the nasopharynx, auscultation and Doppler ultrasound examination of the neck vessels, and auscultation of the aural region.

Inspection of the external ear can reveal developmental anomalies that may be important for diagnosis of tinnitus. Surgical scars detected during examination of the retro-auricular region may be an indication of previous middle ear surgery, another possible cause of tinnitus.

OtoscopY

Examination of the external auditory canal and tympanic membrane is performed ideally using a microscope with up to 10× magnification. Normally, the external auditory canal is wide and the tympanic membrane is transparent. Attention should be paid to bony exostoses¹ and tumors on the skin lining the ear canal. The long process of the incus is often visible through the tympanic membrane in many cases, as well as the manubrium of the malleus with the umbo. Reddening of the tympanic membrane or increased vascular markings is indicative of acute otitis media. Otoscopy can detect fluid build-up in the tympanic cavity. Negative air pressure in the tympanic cavity causes retraction of the tympanic membrane and suggests Eustachian tube dysfunction. Perforations of the tympanic membrane indicate chronic otitis media.

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¹Exostosis: A cartilage-capped bony projection arising from any bone that develops from cartilage. From: Stedman’s Electronic Medical Dictionary.
The presence of cholesteatoma may cause the epitympanic perforation to be filled with keratin debris or granulation tissue. Stinking otorrhea is also characteristic of the presence of a cholesteatoma. In many cases, large tympanic membrane perforations permit inspection of the auditory ossicular chain and assessment of the tympanic mucosa. Otoscopy findings are usually normal in otosclerosis. Schwartze’s sign is a rare finding, consisting of a pink blush of the mucosa near the promontory that can be seen through a translucent tympanic membrane. It is a sign of proliferation and dilatation of blood vessels at the promontory and should be regarded as indication of cochlear otosclerosis.

Movement of the tympanic membrane in time with the patient’s respiration on forced nasal breathing is indicative of a patulous Eustachian tube. Pulsatile movement of the tympanic membrane may indicate the presence of a glomus tumor. This bluish-red tumor growing in a grape-like cluster may cause the tympanic membrane to bulge outward. In advanced cases, the tumor may also impinge into the external auditory canal. Rhythmic movements of the tympanic membrane are also noted in patients with myoclonus of the tensor tympani muscle.

Valsalva’s maneuver and Toynbee’s maneuver are clinical tests of Eustachian tube function. In Valsalva’s maneuver, patients pinch their nose tightly between thumb and forefinger and then attempt to breathe out forcibly while also keeping their mouth closed. An outward bulging tympanic membrane during a Valsalva’s maneuver is an indication that the Eustachian tube functions normally and that air can reach the middle ear via the Eustachian tube. In Toynbee’s maneuver, the patient is instructed to swallow, again with nose pinched closed. This normally results in a negative air pressure in the tympanic cavity that can be detected by otoscopy as an inward retraction of the tympanic membrane. Objective measure of Eustachian tube function can be obtained by tympanometry (see Chap. 49).

Endoscopy

Further examination of possible causes of dysfunction of the Eustachian tube can be done by examination of the nasal and pharyngeal structures using rigid or flexible endoscopes. Polypoid mucosal changes in the nasal cavity are indicative of chronic rhinosinusitis that may affect the function of the Eustachian tube. Attention should also be paid to the possible presence of space-occupying lesions in the nasopharynx that may obstruct the tubal orifice (hypertrophied lymphatic tissue, nasopharyngeal carcinomas). Many patients with a patulous Eustachian tube have a widening of the opening of the Eustachian tube in the nasopharynx. Biopsy of suspected space-occupying lesions should be done for histopathological examination. Objective tinnitus may be caused by myoclonus of the palatine muscles (the tensor and levator veli palatini). Twitching of these muscles can be observed by endoscopy.

Auscultation

Auscultation of the ear and neck vessels can be important in the diagnosis of objective tinnitus. A stethoscope can be used to detect signs of carotid artery stenosis, such as what may occur from atherosclerosis, vascular compression, and arteriovenous malformations. Dural AV fistulas can be detected by auscultation of the upper neck and in the post-auricular region. According to, the use of an electronic stethoscope is more sensitive than classic auscultation techniques. If a patient’s tinnitus is affected by compression of the neck or from turning the head, it is a sign that the cause is of venous origin. Such maneuvers would have no effect on pulsatile tinnitus of arterial origin. Abnormal auscultatory findings require further clarification using Doppler ultrasound or angiography. Direct auscultation of the middle ear performed with a Toynbee tube inserted into the external auditory canal makes it possible to hear the tinnitus that is caused by contractions of the stapedius or tensor tympani muscles of the middle ear.
Supplementary Radiological Diagnosis

Radiological examinations may be justified when otologic examinations are inconclusive regarding pathologies that may be involved in causing tinnitus [7]. High-resolution computed tomography (CT) of the petrous portion of the temporal bones can be used to detect and examine structural bony changes of the external ear and its surroundings, the middle ear, and inner ear.

MRI can be used to visualize a fluid-filled cochlea. Finally, this technique is used to diagnose intra- or extrameatal acoustic neurinomas. Detailed information regarding these techniques is provided in Chap. 19.

Doppler studies of vessels on the neck are helpful in the diagnosis of pulsatile tinnitus (see Chap. 59). In some countries (e.g., Germany), ultrasound techniques are part of an otologic examination. Doppler ultrasound allows visualization of carotid stenoses, arterial dissections, and arteriovenous malformations. Sismanis and Smoker [8] also recommend extending the use of this modality to include the subclavian arteries. Digital subtraction angiography is employed for the preoperative assessment of a glomus tumor, permitting identification of the main supplying vessel on the basis of the tumor blush. Interventional embolization of the supplying vessels during the same session prepares for efficient control of bleeding during subsequent surgical resection.

References

Chapter 49
Audiologic Clinical Assessment

Umberto Ambrosetti and Luca Del Bo

Keypoints

1. Tinnitus may be the symptom of many different disorders.
   An accurate assessment of a patient’s history, symptoms, and signs is important to establish a correct diagnosis.

   The tinnitus handicap inventory (THI) and the visual analog scale (VAS) are very useful tests to evaluate the handicap caused by tinnitus and the entity of tinnitus, respectively.

2. An objective assessment of ear, head, neck, and temporomandibular articulation should be performed.

3. Pure-tone audiometry (the frequency range from 125 to 16 KHz), tympanometry, acoustic middle-ear reflex testing, speech recognition threshold testing, and speech discrimination tests help determine the type of hearing loss and the status of the middle ear.

4. Otoacoustic emission (OAEs) testing allows for precise evaluation of the outer hair cell function.

5. Acufenometry is performed to determine pitch and loudness of tinnitus by defining minimum masking levels (MMLs), loudness discomfort levels (LDLs), and the residual inhibition.

6. Auditory brainstem responses (ABR) are used in selected patients for further evaluation and exclusion of disorders such as vestibular schwannoma.

   Electrocochleography (ECochG) is used in order to evaluate the electric phenomena of the inner ear.

Keywords Tinnitus • Assessment • Pure-tone audiometry • Speech audiometry • Impedance • Otoacoustic emissions • Acufenometry • Brainstem-evoked potentials.

Abbreviations

ABR Auditory brainstem response
AEP Auditory evoked potential
dB Decibel
DPOAE Distortion product otoacoustic emissions
ECochG Electrocochleography
fMRI Functional Magnetic Resonance Imaging
HL Hearing level
Hz Hertz
IHCs Inner hair cells
LDL Loudness discomfort level
MEG Magneto Encephalographic
MLR Middle latency response
MML Minimum masking level
MRI Magnetic resonance imaging
OAE Otoacoustic emissions
OHCs Other hair cells
PET Positron emission tomography
RI Residual inhibition
NMR Nuclear magnetic resonance
CT Computed tomography
TEOAE Transiently evoked otoacoustic emissions
THI Tinnitus handicap inventory
VAS Visual analog scale
WN White noise

L. Del Bo (*)
Fondazione Ascolta e Vivi, via Foppa 15, 20144 Milan, Italy
e-mail: delbo@sordita.it

Introduction

An audiologic diagnosis is crucial to identify auditory system pathologies. In particular, an accurate assessment must precede any further treatment. A detailed assessment may even be therapeutic for patients with tinnitus, as it often reduces many of the patient’s concerns and reactions to their disease. Audiologic assessments can rule out severe diseases, which may have been of great concern to the patient, and the real cause of tinnitus is identified.

Tinnitus may be the symptom of many different disorders of the human auditory system that must be accurately investigated in order to reach a diagnosis that allows the best treatment. These treatments allow for the cure or management of the patient’s tinnitus by means of medical or surgical therapies, as well as specific dietary regimens or other available treatments.

Audiological History

The medical history of patients with subjective tinnitus should include information about infectious diseases during childhood, previous surgical interventions, endocrine and metabolic disorders, and hypertension. Middle-ear diseases (stenosis and insufficiency of the Eustachian tube, secretive otitis media, acute and chronic otitis media, tympanosclerosis with a close or open tympanum, or otosclerosis) and inner ear diseases (toxic drug–induced damage, exposure to acoustic traumas, Ménière’s disease, or presbyacusis) should also be included in the history.

Also, alterations in the skeleton, either congenital or induced by injuries and traumas, should be investigated, as well as posture variations due to pathologies of lumbar-sacral and cervical vertebral column, pelvic girdle, and, in particular, the stomatognatic apparatus [1, 2].

The characteristic of tinnitus should be determined, including duration, intensity, loudness, continuity, intermittence, pulsatility, variations produced by physical effort, and psychological effects [3, 4].

The tinnitus handicap inventory (THI) questionnaire [5] is a very useful instrument to evaluate the handicap caused by tinnitus; it is simple, short, and immediate. McCombe and colleagues [6] reduced the THI questionnaire scorings into a grading scale to estimate the development of tinnitus: Grade 1 (0–16), slight tinnitus; Grade 2 (18–36), mild tinnitus; Grade 3 (38–56), moderate tinnitus; Grade 4 (58–76), severe tinnitus; and Grade 5 (78–100), catastrophic tinnitus.

The visual analog scale (VAS) is a scale ranging from 0 to 10, which is used to quantify the entity of the disease as tinnitus, hyperacusis, or deafness by directly evaluating the symptoms reported by the patient [7, 8].

Objective Examination

Once all the significant anamnestic data have been collected, the objective examination of the outer and middle ear along with all the ear, nose, and throat area is to be performed.

The areas relating to the auricle, mastoid, and temporomandibular articulation must be investigated in order to identify possible malformations, stenosis, atresia in the external auditory canal, or any asymmetry between the two auricles, which is indicative of malformations or associated syndrome. Palpation of the whole auricular area must then be performed and, in particular, it must be investigated whether the patient experiences pain on both sides of the temporomandibular joint (TMJ) either with a closed mouth or during mastication.

The cartilaginous and bone portions of the auditory canal and the tympanic membrane are then examined: an operating microscope should be used, which allows a stereoscopic visualization. The examination of the external auditory canal may reveal build-up of wax or epidermal residues that must be removed in order to allow the examination of the tympanic membrane. Earwax must be totally removed, even when it is “spread” on the tympanic membrane or located in the anterior tympano-meatal corner. In fact, in these cases, earwax can often be difficult to remove with a simple remove wash without causing a sense of ear occlusion, hearing loss, or tinnitus. Hairs or hair fragments in the ear canal may also be accurately removed, as they may cause tinnitus, which resolves spontaneously after removal. Dermatologic diseases in the skin of the ear canal or anatomic stenosis, caused by single or multiple osteomas, may also be looked for.

The color of the tympanic membrane, which normally appears pearly, must be carefully examined for its brightness and the possible presence of air blisters,
scars, dermo-epidermal blisters, single retraction pockets, or partial and total perforations.

In case of tympanic perforation, its extension and localization must be carefully assessed, as well as the possible presence of otorrhea, which should be removed by means of an aspirator if it is abundant. Small granulations, if present, must also be removed in order to proceed with a proper assessment of the tympanic membrane.

Once the external auditory canal has been duly cleaned, secretion-induced crusts must be removed from the tympanic membrane; after removing any residue that can mask tympanic perforations or retraction pockets, the presence of whitish tissue must be investigated, in particular, in the epitympanum, pathognomonic of cholesteatomatous chronic otitis.

Besides the otologic clinical assessment, an objective examination of both nose and throat must be performed, possibly with the aid of a fiber-optic endoscopy.

In case of pulsating tinnitus, the large vessels of the ear and neck must be ausculted in order to identify objective tinnitus [9, 10].

Diagnostic imaging is required if the clinical assessment reveals a suspicion of expansive growth pathologies in the middle ear or pontocerebellar corner. Computed tomography (CT) and gadolinium magnetic resonance imaging (MRI) of the cerebellopontine angle is warranted to exclude a vestibular schwannoma in case of audiologic suspicion (asymmetric hearing loss, no middle-ear reflexes). Further diagnostic work-up may include echo-doppler of supra-aortic trunks or the cochlear labyrinth, as warranted. Angio-MRI may be done if vascular diseases are suspected. Additional diagnostic imaging techniques, such as positron emission tomography (PET), functional magnetic resonance imaging (fMRI), magneto encephalographic (MEG) – all used in clinical research – have not yet been included in clinical diagnostic protocols.

Hematochemical analyses provide information on possible metabolic disorders (dyslipidemia, altered blood viscosity), flogistic diseases (immunocomplexes, antibodies), endocrine diseases (dysthyroidism, diabetes), and others.

In case of isolated tinnitus, causing or favoring factors should be investigated within the vascular system (hypertension, vertebrobasilar insufficiency) or within the neurologic system (epilepsy).

Psychiatric etiology should also always be considered (hysteria, psychosis, schizophrenia), as well as any manifestation of anxiety or depression [4].

## Audiometric Evaluation

Several audiometric tests are required to complete a clinical examination of patients with tinnitus. This includes both subjective and objective tests (Table 49.1), which should be carried out in a succession to allow a comprehensive evaluation of the auditory system, from the periphery to the central auditory nervous system.

Audiometric evaluation should include the following tests:
- Pure-tone thresholds
- Speech recognition thresholds
- Acoustic middle-ear reflex testing (reflex tone decay only if comfortably tolerated by patient) and tympanometry

### Table 49.1 List of subjective and objective audiologic examinations and relating aims in tinnitus diagnosis

<table>
<thead>
<tr>
<th>Subjective examination</th>
<th>Objective examination</th>
<th>Aim</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pure-tone thresholds</td>
<td>Tympanometry and acoustic reflex testing</td>
<td>Auditory threshold assessment and damage site localization</td>
</tr>
<tr>
<td>Speech recognition thresholds and discrimination</td>
<td>Otoacoustic emissions</td>
<td>Middle ear and Eustachian tube function, cochlear and retrocochlear dysfunctions</td>
</tr>
<tr>
<td>Acufenometry</td>
<td></td>
<td>Damage site definition and confirmation, communication capabilities assessment</td>
</tr>
<tr>
<td>Loudness disconfort level (LDL)</td>
<td>Acoustic evoked potentials: ECochG, ABR, MLR</td>
<td>Tinnitus pitch and loudness, MML and RI definition</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Doscomfort threshold definition</td>
</tr>
</tbody>
</table>
Pure-Tone Thresholds

The first test to be performed is a plain pure-tone audiogram [11] with the patient sitting in a soundproof booth; pure tones of known intensity and frequency are delivered through insert earphones, or through supraaural headphones, and an audiometer, which is controlled by the examiner.

Clinical audiometers usually have a frequency range from 125 to 8,000 Hz, at an intensity range between 10 and 120 dB hearing level (HL). However, for use in patients with tinnitus, the range must be extended up to 16,000 Hz [12, 13].

Audiometer calibration must be performed at least once a year, since it is important to be able to detect even minor threshold variations from normal conditions [14]. This test, despite being a relatively simple one, actually requires an experienced examiner with excellent clinician–patient communication skills, so that the most precise threshold can be obtained.

In case of large degree of asymmetry in hearing threshold between the two sides, “masking” noise should be delivered to the healthier ear, so that this latter does not perceive the acoustic stimulus via bone conduction [15]. The “cross-talk” effect is different for different earphone types, being much greater for supraaural head transmission than for insert earphones (see [16], page 299).

Bone conduction audiometry that allows direct stimulation of the inner ear (hair cells), “bypassing” the ear canal and middle-ear transmission system, makes it possible to separately evaluate the hearing threshold without the influence of the conductive apparatus.

For that, a small vibrator is placed on the patient’s mastoid bone. The vibrator allows delivery of stimuli frequency ranging from 250 to 4,000 Hz, and the highest deliverable intensity varies according to the tested frequency [17].

When the air conduction threshold is worse than the bone conduction one, it is an indication of conductive hearing loss. When both thresholds are equal and elevated, it is a sign of sensorineural hearing loss.

Elevation of the hearing threshold, even a mild one, within the frequency range 8,000–16,000 Hz may be the cause of acute tinnitus.

Elevation of the hearing threshold in the frequency range between 3,000 and 6,000 Hz (determined in steps of half octaves) must also be investigated; such hearing loss may be an indication of isolated selective damages to the auditory system that could lead to frequency-specific tinnitus in this frequency range. Some individuals with tinnitus have several dips in their audiograms, which may be indications of contact between a blood vessel and the auditory nerve root (see Chap. 40).

Tympanometry and Acoustic Middle-Ear Reflex Testing

Tympanometry measures the change in the ear’s acoustic impedance when the air pressure in the ear canal is varied. The results appear as a curve of the acoustic impedance (or more often its inverse, known as the acoustic admittance). The term “immittance” is often used to describe the acoustic compliance of the ear. Tympanometry cannot be carried out when the external auditory duct is occluded or the tympanic membrane is perforated.

This method gives objective clinical information; it is a non-invasive exam and a quick one.

The examination is performed by introducing into the external auditory canal a probe equipped with a soft end cap, ensuring pneumatic capacity, besides the following three additional functions: (1) it generates a sound called “probe tone” (220 Hz–65 dB HL); (2) by means of a pump, it varies the pressure from positive values (+200 mm/H2O) to negative values (−400 mm/H2O); and (3) a microphone records the sound pressure in the ear canal. The sound pressure in the ear canal is a measure of the ear’s acoustic impedance (or admittance).

One single exam allows, in a very short time, three different tests to be performed: tympanometry, assessment of acoustic stapedial reflexes, and assessment of the Eustachian tube function.

The test has two parts: one is tympanometry, which evaluates the compliance (mobility) of the middle ear. The second test uses similar techniques to record
contractions of the middle-ear muscles (mainly the stapedius muscle) elicited by a loud tone presented to the same or the opposite ear. A normally functioning middle ear has a tympanogram that is bell shaped with a peak center on zero pressure (Type A); a tympanogram showing a flat line is, instead, consistent with an increase in rigidity of the tympanic ossicular system (Type B); a bell-shaped tympanogram with a peak shifted to the negative pressure values on the graph (Type C) is indicative of pressure in the middle-ear cavity that can be caused by secretive otitis media or Eustachian tube stenosis [18] (Table 49.2).

Tympanometry allows the evaluation of the function of the Eustachian tube, such as its ability to open during deglutition. Once the tympanogram has been obtained, external auditory canal pressure is increased to +200 mm H2O, and the patient is asked to swallow; under normal conditions, the tympanogram changes as a sign of air coming out of the middle ear and returns to normal after deglutition. This test is crucial for tinnitus evaluation, since altered tubal function is associated to tinnitus appearance.

The acoustic middle-ear reflex (stapedial reflex) is elicited with tones in the frequency range 500–1,000 and in the range of 2,000–4,000, applied to the contralateral or the ipsilateral stimulus one at a time. Contraction of the middle-ear muscles (stapedius and tensor tympani) causes the ear’s acoustic impedance to change, and that makes a non-invasive way of recording the response of the acoustic middle-ear reflex. If the middle-ear pressure is different from the ambient pressure, as indicated by the tympanogram, it has to be equalized before testing the acoustic middle-ear reflex. Under normal conditions, the threshold for the acoustic middle-ear reflexes for tones is 90 dB HL in a normally hearing individual. It is slightly (2–10 dB) lower when elicited from the ipsilateral ear.

Cochlear hearing loss is associated with a distortion phenomenon called “recruitment of loudness”, so that a sound is perceived louder than normal. This should be distinguished from hyperacusis, which is a lowered tolerance for sounds (see Chap. 3). The stapedial reflex threshold, as recorded for different frequencies, allows the physician to objectively determine recruitment of loudness.

Assessment of the decay of the response of the acoustic middle-ear reflex at prolonged stimulation may be an indication of auditory nerve diseases (Anderson’s test) [19]. However, the reliability of this test has been disputed.

It has been estimated that 40% of individuals with tinnitus also have hyperacusis. Such patients may experience discomfort from acoustic overstimulation, such as in testing the acoustic middle-ear reflex.

### Speech Recognition Thresholds and Speech Discrimination

In order to properly assess an individual’s hearing capacity and quantify its impact on speech recognition, threshold, and speech discrimination scores, speech audiometry is used, by means of words and sentences as test sounds. Recorded speech material is preferred to live voice presentation. Standardized lists of words and sentences are delivered at different intensities, to one ear at a time, recording the number of correct answers given by the patient. The healthier ear must be masked in patients with a marked tone threshold asymmetry when the ear with hearing loss is tested.

Under normal conditions, the articulation curve is S shaped; in cases of conductive hearing loss, the curve is shifted to the right. In patients with cochlear or retrocochlear hearing loss, the shape of the curves, in addition to being shifted to the right, are changed, and 100% intelligibility will not be achieved.

### Otoacoustic Emissions

The damage to other hair cells (OHCs) is believed to be involved in the development of peripheral tinnitus in individuals with a normal hearing sensibility [20, 21]. Recording of otoacoustic emissions (OAEs) is
important because it allows to identify changes in the function of the OHC.

OAEs are weak sounds generated by the electromotility of the OHC, which pass through the middle-ear ossicles and tympanic membrane and can be measured in the external ear canal [22].

Recordings of OAEs are objective tests of cochlear (OHC) function and their presence, in general, is indicative of normal hearing. This test is very sensitive to cochlear damage that involves OHCs and can often be detected before clinical evidence of hearing loss is present. Successful recording of OAEs depends on normal function of the middle ear.

In the clinical practice, particularly in the study of tinnitus, the subclasses of OAEs, which are most useful and utilized, are transiently evoked otoacoustic emissions (TEOAEs) and distortion-product otoacoustic emissions (DPOAEs).

TEOAEs are less affective by contralateral white-noise suppression in individuals with tinnitus than in individuals with similar hearing impairments and without tinnitus [23]. One explanation of this observation may be a hyperactivity of the OHCs resulting from pathological cochlear activity [24].

Recording of distortion-product otoacoustic emissions (DPOAEs) can provide a detailed and tonotopic OHC test that can identify small areas of cochlear damage; DPOAEs are recorded at frequencies as high as 8–10 kHz with up to 10 points/octave.

The value of recording DPOAEs in patients with tinnitus has been controversial [25–29].

**Tinnitus Loudness and Pitch Matching**

Acufenometry is the technique used to determine the frequency range (pitch) of tinnitus, its subjective intensity (loudness), the ability of sounds to mask the tinnitus of an individual person, and the residual inhibition of tinnitus. The test becomes difficult to perform when the tinnitus has the character of complex noise.

It is well known that one tone may be masked by another pure tone with enough intensity. Narrowband noises also have masking characteristics similar to those of pure tones while tinnitus that sounds like wideband noises do not have similar masking features. A tone may easily be masked by a wideband noise, while a wideband noise is hardly ever masked by a pure tone.

Tinnitus that is referred to one ear may be masked by sounds applied to the ipsilateral as well as the contralateral ear. Contralateral masking requires higher sound levels than ipsilateral masking [30].

Masking or suppressing tests make it possible to determine the tone intensity or noise that completely suppresses the perception of tinnitus. Once the type of sound that effectively masks the tinnitus has been found, the stimulus intensity is then increased 2 dB steps, up to a level where the tinnitus can no longer be heard.

The intensity difference, measured in dB, between the hearing threshold for the masking tone and the level of minimum masking intensity is the “minimum masking level” (MML).

In most patients, the MML plotted as a function of frequency intersects with the hearing threshold, indicating the specific frequency of tinnitus [31].

Hyperacusis is studied by determining the level of acoustic stimulation threshold that produces discomfort (loudness discomfort level – LDL) to the patient.

Hyperacusis is defined as a lower than normal discomfort level (see Chap. 3). Hyperacusis is different from recruitment of loudness, which is a perception that sounds are abnormally loud, and the threshold of the acoustic middle-ear reflex is lower than it would be for the same hearing loss that occurs without recruitment. Masking noise may often induce temporary tinnitus suppression or a temporary relief from tinnitus, known as “residual inhibition” (RI) or “residual suppression” [31–34].

To perform the test, the noise level established for MML is raised by 10 dB and presented for exactly 1 min.

The examiner waits for a few seconds and then asks, “Does your tinnitus sound the same as before, or is there any difference?” The residual inhibition is present if the patient reports a lower level of tinnitus. The total duration of residual inhibition is recorded and measured in seconds [35]. As a masker intensity is increased to +20 dB, the depth and duration of RI increase.

RI is one of the few procedures that may reduce or eliminate tinnitus for a brief period.

The presence of RI, in individuals with hearing loss, allows efficient control of tinnitus by means of hearing aid, which provides sufficient amplification at the frequencies of the tinnitus [36].
**Auditory-Evoked Potentials**

Essentially, four kinds of auditory-evoked potentials are in clinical use (Table 49.3). These tests cannot be done in patients with hearing loss exceeding 80–90 dB HL.

When elicited by a transient sound, click, or short tone burst, electrocochleography (ECochG) records the sound-evoked potentials generated in the cochlea (cochlear microphonics and summating potentials) and in the distal portion of the auditory nerve (compound action potential). ECochG is performed by placing a recording electrode either deep in the ear canal or using a needle that is inserted through the tympanic membrane to come in contact with the cochlea capsule. It allows a detailed evaluation of the electric phenomena occurring in the cochlea and in the distal portion of the auditory nerve [37].

Recording of the auditory brainstem response (ABR) allows evaluation of the auditory threshold when elicited by high-intensity clicks or tone bursts (approximately 65 dB HL). ABR recordings provide information about the integrity of the ascending auditory pathways up to the midbrain level (inferior colliculus).

ABR recordings provide diagnostic information about auditory nerve injuries, presence of vestibular schwannoma, and other pathologies of the lower auditory pathways. ABR test requires the application of three surface electrodes placed on the patient’s skin after degreasing the skin and spreading a conductive paste [to forehead center (or better on the top of the head of Cz), ear lobes]. The response to 2,000 click sounds delivered through a headset is normally averaged to get an interpretable record.

By increasing the frequency of the acoustic stimulation (high rate potential), further information may be obtained concerning the latencies of the components of the ABR recordings; Selters and Brackman [38] reported that the ABR had a high sensitivity for deflecting the presence of vestibular schwannoma when the recorders are compensated for age-related changes in hearing threshold.

Godey and colleagues [39] reported that the ABR alone detected vestibular schwannoma in 92% of patients, while together with recordings of acoustic middle-ear reflex and caloric vestibular responses, the sensitivity was 98%.

<table>
<thead>
<tr>
<th>Table 49.3</th>
<th>Different types of auditory evoked potential and investigated site</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Auditory evoked potential</strong></td>
<td><strong>Investigated site</strong></td>
</tr>
</tbody>
</table>
| ECochG | • Cochlea  
| | • Auditory nerve |
| ABR | • Auditory nerve  
| | • Nuclei of the ascending auditory pathway |
| MLR | • Thalamus  
| | • Primary auditory areas |

**References**

Chapter 50
Clinical Otoneurological Examination

Carlos Herráiz

Keypoints

1. The relevance of the anterior labyrinth (cochlea) in tinnitus generation opens the possibility that some patients will show damage in the posterior labyrinth as a whole inner ear disease.
2. The connections between the central vestibular pathways and the auditory, visual, and somatosensory systems could be also implicated in the mechanisms of tinnitus.
3. An exhaustive otoneurological examination is recommended in those patients we suspect a vestibular affection. The medical history will give us the most important information for the etiology and the severity of the symptoms.
4. The examination is driven toward three main systems: cranial pairs and cerebellum, the vestibulo-ocular reflex, and the vestibulo-spinal reflex.
5. A basic office exam can give us much information about the affected side and the compensation stage.
6. The instrumental examination, mostly performed in the chronic stages, will objectify and measure some aspects of the vestibular impairment. The oculomotor system examination and vestibulo-ocular reflex (rotatory and caloric stimulation) will be tested with the videonystagmograph.
7. The vestibulo-spinal reflex will be checked using the dynamic posturography. This instrument will test the balance and lateropulsion from a dizzy patient, as well as the strategies for keeping the equilibrium right over specific sensorineural afferent disruption (visual, somatosensory, and vestibular).
8. Understanding the mechanisms and pathophysiology of the vestibular system will help us to connect the multiple sensory pathways involved in some forms of tinnitus.

Keywords  Tinnitus • Dizziness • Vertigo • Videonystagmography • Dynamic posturography

Introduction

The cochlea is the location of the pathology that causes some forms of tinnitus, but there is now evidence that the central nervous system (CNS) is the anatomical location of the abnormal neural activity that causes many forms of tinnitus. There is also evidence that these changes are caused by activation of neural plasticity and that acoustic deprivation is the most important cause and chronic maintenance of these changes [1]. The abnormal neural activity that causes tinnitus may be generated along the peripheral and central auditory pathways (CAP), or in other systems connected to these pathways such as the somatosensory or the limbic–amygdala complex. While there is evidence that connections between the central vestibular pathways and the auditory, visual, and somatosensory systems might be implicated in causing some forms of tinnitus, there is no evidence of similar interaction with the vestibular system. It is therefore only when vestibular disorders accompany tinnitus that it is important to perform a complete otoneurological examination.
Medical History

Dizziness can represent a group of highly heterogeneous symptoms and sensations. Vertigo is the sensation of motion or spinning from oneself or from the environment and is often accompanied by vegetative symptoms. It is secondary to vestibular disorders in the majority of cases (central or peripheral). Imbalance or unsteadiness describes a loss of equilibrium on movement, or in situations in which there are conflicting sensory cues. It can represent the normal pattern of compensation after an acute vestibular lesion or a poor compensation if the symptoms are persistent after a few months. Multisensory disorders (visual, somatosensory, motor, or cerebellar) can manifest imbalance. Dizziness is any vague sensation of discomfort in the head: light-headedness, disorientation, floating, etc. Poor vestibular compensation or psychiatric disorders such as depression, anxiety, or hyperventilation can cause it. The syncope is secondary to cardiovascular disorders with reduced cerebral blood flow. Severe bilateral peripheral vestibular disorders or central diseases can produce a lack of stabilization of the visual field with passive whole-body movement called oscillopsia [2, 3]. Duration of vertigo, trigger factors, characteristics and frequency of spells, and presence of vegetative symptoms will help us in the diagnosis of the disease (Tables 50.1 and 50.2).

Otoneurologic Examination

The otoneurological examination is driven toward three main systems: cranial nerves and the cerebellum, the vestibulo-ocular reflex, and the vestibulo-spinal reflex.

Table 50.1 Causes of vertigo and imbalance according to duration of the symptom

<table>
<thead>
<tr>
<th>Seconds</th>
<th>Minutes/hours</th>
<th>Days</th>
<th>Months</th>
</tr>
</thead>
<tbody>
<tr>
<td>BPPV</td>
<td>Ménière’s disease</td>
<td>Vestibular neuritis</td>
<td>Vestibular neumora</td>
</tr>
<tr>
<td>Perilymphatic Fistula</td>
<td>Endolymphatic hydrops</td>
<td>Sudden deafness</td>
<td>Ototoxicity</td>
</tr>
<tr>
<td>DSSC Sdr.</td>
<td>Vestibular migraine</td>
<td>Labyrinthis</td>
<td>Bilateral vestibular</td>
</tr>
<tr>
<td>Cerebellar disorders</td>
<td>Perilymphatic fistula</td>
<td>Ramsay–Hunt syndrome</td>
<td>disorders: genetic,</td>
</tr>
<tr>
<td>Falls</td>
<td>Vertebrobasilar stroke</td>
<td>Cerebrovascular stroke</td>
<td>bilateral MD, idiopathic</td>
</tr>
<tr>
<td></td>
<td>Seizures</td>
<td>Multiple sclerosis</td>
<td>Psychiatric disorders</td>
</tr>
<tr>
<td></td>
<td>VIII nerve vascular compresion syndrome</td>
<td>Viral infections of the brain stem</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Central paroxysmal vertigo of the cerebral stem</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Panic attack</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>Postural phobic vertigo</td>
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<td></td>
</tr>
</tbody>
</table>

Table 50.2 Differences between peripheral and central vertigo

<table>
<thead>
<tr>
<th></th>
<th>Peripheral vertigo</th>
<th>Central vertigo</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical symptoms</td>
<td>Harmonic*</td>
<td>Dys-harmonic</td>
</tr>
<tr>
<td>Optocinetic/pursuit test</td>
<td>Normal</td>
<td>Altered</td>
</tr>
<tr>
<td>Nystagmus</td>
<td>Direction-fixed</td>
<td>Direction-changing, dissociated</td>
</tr>
<tr>
<td>Compensation process</td>
<td>Usual</td>
<td>Unusual</td>
</tr>
<tr>
<td>Hearing</td>
<td>Altered</td>
<td>Normal</td>
</tr>
</tbody>
</table>

*Nystagmus

Nystagmus is an involuntary rhythmic oscillation of the eyes, usually occurring simultaneously in both eyes. Ocular nystagmus is a pendular eye movement with similar speed toward both sides. It is related to abnormal visual fixation associated to loss of central vision. The vestibular nystagmus has a slow phase followed by a fast phase (saccadic movement) toward the opposite side. Nystagmus may be caused by any peripheral or central vestibular disorders of brainstem origin affecting control of the eye muscles. Nystagmus classification is given in Table 50.3. When the fast phase of nystagmus beats toward the healthy ear, it is a sign of peripheral vestibular problems. The intensity of the nystagmus increases when the patient looks toward the side to which the fast phase beats. Nystagmus can be spontaneous, gaze evoked, or voluntarily induced. Vestibular nystagmus is reduced when the patient fixes the gaze. A few individuals have nystagmus without
any known disease. These kinds of nystagmus involve eye movements in the horizontal plane.

The examination of nystagmus involves having the patient follow the examiner’s finger in both directions of gaze, but no more than 30 grades. The use of Frenzel glasses abolishes the gaze fixation making nystagmus appear more intense [2, 3]. Vertical nystagmus is rare and a sign of central nervous system diseases.

There are computerized tests of the vestibular oculomotor reflex (VOR) that measure gain and phase of the VOR. The gain is a measure of how accurately the eyes move in the direction opposite of head movement and the phase angle is a measure of the timing of the movement of the eyes. The videonistagmography tests the oculomotor movements (saccadic, pursuit, optokinetic) and the VOR through a double stimulation: rotatory and caloric (Figs. 50.1 and 50.2). The examination of postural vertigo through Dix-Hallpike maneuvers will diagnose benign positional paroxysmic vertigo. A fistula test is performed with the patient sitting with their head 60° backward, so the horizontal canal of the vestibular apparatus is in the vertical position. With a snugly fitting pneumatic otoscope, alternate positive and negative pressure is rapidly applied to the external ear canal. If the patient becomes dizzy and gets nystagmus, it is a sign of the presence of a perilymphatic fistula.

### Table 50.3 Characteristics of nystagmus

<table>
<thead>
<tr>
<th>Direction</th>
<th>Direction-changing</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Direction-fixed</td>
</tr>
<tr>
<td>Plane</td>
<td>Horizontal, horizonto-rotatory, rotatory, vertical, oblique</td>
</tr>
<tr>
<td>Intensity</td>
<td>Three degrees (Alexander’s law)</td>
</tr>
<tr>
<td>Mode of occurrence</td>
<td>Spontaneous, gaze-evoked, induced (optokinetic, head shaking, fistula test, drugs)</td>
</tr>
<tr>
<td>Specific forms</td>
<td>Endpoint N, congenital N, periodic alternate N</td>
</tr>
</tbody>
</table>

### Tullio Phenomenon

Vertigo that is induced by high intensity sound is known as the Tullio phenomenon. It is positive in dehiscent superior semicircular canal syndrome and in the perilymphatic fistula. Making the Valsalva maneuver will typically induce vertigo in a patient who has a perilymphatic fistula or patients with an Arnold–Chiari malformation [4].

Unsteadiness and lateropulsion can be a consequence of vestibulo-spinal reflex (VSR) disorders.
Romberg, Unterberger, and the test of the gait with the Babinsky–Weil test should be performed in all tinnitus patients with vestibular symptoms. Dynamic posturography is a computerized method to evaluate the VSR and the equilibrium strategies: visual, vestibular, and proprioceptive (Fig. 5.3). Recording of the vestibularevoked myogenic potentials (VEMP) is a novel technique to test the saccula function.
Conclusions

Otoneurological examination in patients with tinnitus can be useful in the diagnosis of only some forms of tinnitus where the vestibular system is suspected to be involved, such as in connection with vestibular schwannoma. If vertigo or any kind of dizziness occurs together with tinnitus, an otoneurologic examination may be justified. Such examination must be done by a person who understands the mechanisms and pathophysiology of the vestibular system. Tests of the vestibulo-ocular and vestibulo-spinal reflexes are important in the examination of the vestibular disorders.

References


Fig. 50.3  Dynamic posturography for vestibular rehabilitation
Chapter 51
Diagnosis of Tinnitus: Neurological Examination

Miguel J.A. Láinez, Anna Piera, and Alejandro Ponz

Keypoints

1. There is an urgent need for a set of assessment methods to be agreed upon and utilized by the international tinnitus research community.
2. Neurological examination of tinnitus patients is essential to achieve a good diagnostic approach to the different forms of objective and subjective tinnitus.
3. This chapter summarizes the neurological examination in tinnitus, including the protocol used in the authors’ tinnitus clinic, which is based on the consensus of the Tinnitus Research Initiative (TRI).

Keywords

Tinnitus • Questionnaire (TSCHQ) • Tinnitus handicap inventory (THI) • Neurological examination • Doppler sonography

Abbreviations

MR Magnetic resonance
MRA Magnetic resonance angiography
SCM Sternocleidomastoid
TCCS Transcranial color coded sonography doppler
HAM-D Hamilton rating scale for depression
TCD Transcranial doppler
TMJ Temporo-mandibular joint
TRI Tinnitus Research Initiative
TSCHQ Tinnitus sample case history questionnaire

Introduction

Chronic tinnitus, the phantom perception of sound, can be a debilitating and life-altering experience. It affects millions of people in western countries. Despite the enormous social and economic burden tinnitus causes, no well-established treatment for this specific disorder is available. Among the reasons for this unsatisfactory situation are the difficulties in assessing tinnitus, as it is a purely self-reported phenomenon.

There is an urgent need for a set of assessment methods to be agreed upon and utilized by the international tinnitus research community. This includes assessment of patients with tinnitus and subsequent measurement of outcomes following intervention [1].

There is a need for standardization of the ways patients with tinnitus are assessed and the way outcomes of interventions are measured to facilitate more effective cooperation and more meaningful evaluations and comparisons of the outcomes of treatment. So far, three meetings organized by the TRI were held in 2006, 2008, and 2009 to develop a consensus for patient assessments and outcome measurements. This has already contributed to better cooperation between research centers in finding and evaluating treatments for tinnitus by making it possible to better compare the results of studies and treatments [2].

During these workshops, the participants reviewed provisional consensus summaries, and after receiving feedback from all authors, a final consensus was created, giving consideration to the possibility of further modifications [2].

This chapter summarizes the neurological examination in tinnitus, making a brief introduction of protocol used in the authors’ tinnitus clinic, based on the consensus arrived at by workshops arranged by the TRI.
Evaluation of Patients with Tinnitus

Case History

Information about the history and descriptive characteristics of the patient’s tinnitus or tinnitus-related conditions could be obtained by questionnaires and related interviews. The best example is a tinnitus sample case history questionnaire (TSCHQ), in which demographic data and other clinical data are compiled:

- Name, date of birth
- Age, gender, handedness, family tinnitus history
- Time of onset of symptoms
- Was the beginning perception gradual or abrupt
- Was the onset of tinnitus related to loud blast of sound, whiplash, change in hearing, stress, or others
- Is the tinnitus pulsatile or not pulsatile
- Is the tinnitus specific to the right or left ear, the head, or is it similar in both ears
- Constant or intermittent
- Loudness (scale 0–100)
- Sounds like noise, tone(s), sounds of crickets, other
- High pitch, medium, or low frequency (hum)
- Percent of total awake time with tinnitus (1–100%) – Treatments received
- Can the tinnitus be masked by sound
- Is the tinnitus worse in a noisy environment
- Somatic modulation: can head and neck maneuvers, TMJ movement change the tinnitus
- Stress influence
- Hearing problem and hearing aids
- Hyperacusis
- Physical discomfort
- Neck pain
- TMJ disorder
- Psychiatric problems [1]

Psychophysical measures of perceived tinnitus intensity and severity are important for proper diagnosis. Even though these instruments are not specifically designed to be sensitive to treatment-related changes, they have been used as outcome measures in clinical trials. Tinnitus handicap inventory (THI) and Hamilton rating scale for depression (HAM-D) are used if there is psychiatric comorbidity like anxiety and depression [3–6]. Psychopathological aspects of patients with tinnitus and psychiatric comorbidity are discussed extensively in Chaps. 54, 62–64.

Neurological Examination

Neurological examination is essential in tinnitus patients to achieve a good diagnostic approach to the etiology in secondary tinnitus or in objective and subjective tinnitus. Below are the algorithms used by our clinic according to tinnitus workshops and publications. The patient’s case history should always be taken into account when using these suggested assessments (see also Chap. 50).

Vital Signs

1. Blood pressure, pulse rate and character
2. Inspection:
   a. General appearance: anxiety, sadness, attention
   b. Head: irregularities or other deformities in cranial scars and signs of previous trauma; anomalies in temporal arteries
   c. Eyes: Ophthalmologic abnormalities can provide many clues to the etiology of tinnitus – bilateral exophthalmos in thyroid disease, unilateral proptosis in carotid-cavernous fistula, recent skew at basal inspection.

Higher Cortical Functions

A basic neurological examination of higher cortical functions like language and speech are needed to determine if a patient has neurologic, as opposed to psychiatric, disease.

Cranial Nerves

- Optic nerve CN II. Visual acuity and visual fields should be obtained in addition to ophthalmoscopic examination. These tests are indicated if intracranial hypertension or changes in vision caused by any vascular disorders or brain lesion along the optic pathway are suspected.
- The extraocular motor nerves CN III, IV, and VI control the eyelids, pupils, pupillary reflexes, eye movements, and nystagmus. Examination of the eyes begins with inspection – looking for any obvious ocular malalignment or skew, abnormal lid position, or abnormalities of the position of the globe within
the orbit. In routine cases where there are no eye complaints and the likelihood of abnormality is low, the ocular motility examination may be limited to assessing versional pursuit movements in the six cardinal positions of gaze, including full lateral gaze to each side, as well as upgaze and downgaze when looking to either side. If an abnormality of one or more extraocular motor nerves is found in a patient with tinnitus, a few causes should be considered:

1. Carotid-cavernous fistula
2. Ischemic or hemorrhagic ictus in the brainstem area
3. Carcinomatous meningitis – the term carcinomatous meningitis, also called leptomeningeal metastasis, leptomeningeal carcinomatosis, or leptomeningeal dissemination is correct in that case (see footnote)
4. Inflammatory/autoimmunity diseases
5. Horner syndrome in carotid dissection and head trauma
6. Space-occupying lesions

Evaluating nystagmus and other ocular oscillations is essential. Nystagmus is a complex topic. When faced with a patient with nystagmus or similar abnormal eye movements, the usual clinical action includes two steps: deciding if the nystagmus is an indication of neurologic pathology and if so, whether the pathology is central or peripheral. Depending on the etiology [7], nystagmus in patients with tinnitus may be a sign of a neurological disorder or may be the sign of a vestibular disorder as described below.

- Trigeminal nerve CN V. Only a basic examination evaluating the sensitivity in the face of the three nerve branches needs to be checked in the tinnitus neurologist clinic.
- Facial nerve CN VII. Examination of the motor aspect of the facial nerve is essential in patients with tinnitus when the cause may be a vestibular schwannoma. Tinnitus rarely occurs together with Bell’s palsy.
- Vestibular auditory nerve CN VIII. This nerve is mainly examined by the otologist and the audiologist (see Chap. 48 and 49).
- Glossopharyngeal and vagus nerves CN IX and X. The voice, ability to swallow, and dysphasia are functions affected. Nuclear and infranuclear processes that may affect CN IX and X include intramedullary and extramedullary neoplasm, all of which may cause tinnitus. Tinnitus may occur together with glomus jugulare tumors, skull base fracture, surgical trauma, demyelinating disease, or brainstem ischemia. Some of the examinations needed are done by otologists. Palatal myoclonus hyperactivity of muscles innervated by CN IX, in a few cases neuropathic, cause objective tinnitus and are almost always associated with reduced voluntary contraction.
- Spinal accessory nerve CN XI. The examination is limited to evaluation of the functions of the spinal portions. One sternocleidomastoid (SCM) muscle acts to turn the head to the opposite side or to tilt it to the same side. Acting together, the SCMs thrust the head forward and flex the neck. Hyperactivity of the SCM and trapezius may cause spasmodic torticollis in cervical dystonia and in a few patients, it can modulate or cause tinnitus.
- Hypoglossal nerve CN XII. A purely motor nerve, associated with the tongue. Hypoglossal palsy in patients with tinnitus may indicate the presence of infections or neoplastic meningitis, trauma of the skull base, or cervical surgical trauma.

Motor and Sensory Systems

A basic neurological motor and sensitive examination is needed in order to rule out focal neurologic disorders, which are never specific in patients with tinnitus, but necessary to rule out. Some ischemic lesions, brain tumors, and infiltrative disorders may cause tinnitus as a symptom [8, 9].

Cerebellar Function, Gait, and Posture

The cerebellum refines motor commands and is necessary for normal control and regulation of muscle contraction, but it is not involved in the generation of motor commands. It is important to assess cerebellar functions in patients with tinnitus when a secondary cause of the

1 Leptomeningeal dissemination is a condition in which a solid tumor diffusely spreads to the leptomeninges. Lung tumors, breast tumors, and malignant melanoma comprise the majority of solid tumors spreading to the leptomeninges. Alternative definition: an infiltration of carcinoma cells in the arachnoid and subarachnoid space may be primary or secondary.

Synonyms: leptomeningeal carcinoma, leptomeningeal carcinomatosis, meningeal carcinomatosis, leptomeningeal metastasis.
tinnitus is suspected to be in the posterior fossa, near to the cerebellopontine angle. Examples of such causes are vestibular schwannoma, arteriovenous malformations of the posterior fossa, cholesteatoma, and ischemic lesions of vertebrobasilar territory. Depending on the parts of the cerebellum and its annexes involved, patients may suffer from various combinations of tremor, incoordination, difficulty walking, dysarthria, and nystagmus. Thus, examination of coordination in multi-joint movements, the finger–nose maneuver to examine appendicular coordination, tremor, muscle resistance to passive movement, eye movements, equilibratory coordination, gait, and articulation of speech are indicated.

**Neurovascular Examination**

A neurovascular examination is essential to a clinical neurovascular examination in patients with tinnitus because pulsatile and intermittent tinnitus can be secondary to many aspects of diseases such as atherosclerotic disease [10], dural arteriovenous fistulas, aneurysms, and other vascular disorders [11] [benign intracranial hypertension (pseudotumor cerebri), etc.].

**Inspection**

Hardening and tenderness of temporal arteries may reflect giant cell arthritis or may be an indirect sign of an arteriovenous fistula.

**Auscultation**

Auscultation of the head is sometimes useful. Bruits may be heard best over the temporal regions of the skull, the eyeballs, and the mastoids. Cephalic bruits may occur with angiomas, aneurysms, arteriovenous malformations, neoplasms that compress large arteries, and in the presence of atherosclerotic plaques that partially occlude cerebral or carotid arteries. Ocular bruits usually signify occlusive intracranial cerebrovascular disease. A carotid bruit may be transmitted to the mastoid, resulting in objective tinnitus. An ocular bruit in a patient with an arteriovenous aneurysm may disappear when carotid compression is applied.

**Neurovascular Examination with Supra-Aortic and Transcranial Doppler Test**

Transcranial Doppler (TCD) is an imaging test that measures blood flow velocity using ultrasound. It is also known as Transcranial Doppler sonography. Used more recently, transcranial color coded sonography Doppler (TCCS), eco-duplex, and power imaging are tests that can measure the velocity of blood flow through the brain’s blood vessels [12, 13]. The tests are relatively quick and inexpensive ways to aid in the diagnosis of emboli, stenosis, and vasospasm from a subarachnoid hemorrhage, as well as other vascular problems. TCD is often used in conjunction with other tests, such as magnetic resonance (MR), magnetic resonance angiography (MRA), carotid duplex ultrasound, and CT scans.

Two methods of recording may be used in TCD studies. The first uses “B-mode” imaging, which displays a two-dimensional image as seen by the ultrasound probe. Once the desired blood vessel is found, blood flow velocities may be measured with a pulsed Doppler probe, which provides graphical information about blood flow velocities over time. Together, these make a duplex test. The second method uses only the second probe function, relying instead on the training and experience of the clinician in finding the correct vessels.

The equipment used for these tests is becoming increasingly portable, making it possible to use them at hospital bedsides, a doctor’s office, or a nursing home for both inpatient and outpatient studies, and TCD can be used routinely in daily neurovascular examinations by the office of a neurologist.

**Applications of Carotid and Transcranial Doppler Sonography in Patients with Tinnitus**

**Arterial Stenosis**

Doppler sonography is very sensitive to detect carotid and vertebrobasilar stenosis and allows an accurate measure of the size of the stenosis (degree of occlusion).
**Arteriovenous Malformations**

Doppler sonography can detect indirect signs of arteriovenous malformations and dural fistula, revealing coincidental blood supply from other intracranial or extracranial vessels. TCD with echo enhancement is very sensitive to detect these malformations directly, although it is slightly less sensitive than MRA [14].

**Benign Intracranial Hypertension**

TCD provides useful information on cerebral circulation even under raised intracranial pressure. The systolic spike in blood flow as measured by TCD and the size of the arterial pulse (pulsatility index) are useful diagnostic parameters for both acute intracranial hypertension and benign intracranial hypertension. In benign intracranial hypertension, which is a known cause of tinnitus, the pulsatility index measured by TCD is highly correlated with the values of intracranial pressure and cerebral blood flow, thus making it possible to assess and monitor the therapeutic response in patients with such pathologies.

**Lumbar Puncture**

Lumbar puncture is a diagnostic and, at times, therapeutic procedure that is performed in order to collect a sample of cerebrospinal fluid for biochemical, microbiological, and cytological analysis. Very rarely, this procedure is used as a treatment to relieve increased intracranial pressure. Lumbar puncture is performed in patients with tinnitus if an infiltrative or inflammatory cause is suspected, for example, carcinomatous meningitis, subacute encephalitis, cerebral sarcoidosis, or subacute/chronic infections of the central nervous system in general. These conditions rarely cause tinnitus. It is important to recognize that they may be important exceptions.

The value of neuroimaging as a complementary tool in the diagnosis of tinnitus patients is discussed in Chap. 18.

**References**

Chapter 52
Diagnosis of Somatosensory Tinnitus

Tanit Ganz Sanchez and Carina Bezerra Rocha

Keypoints

1. The contribution of non-auditory pathways to the pathology of tinnitus has become more and more evident.
2. Because many different stimuli can modulate tinnitus (forceful muscle contractions of the head and neck, eye movements, pressure of myofascial trigger points, cutaneous stimulation of the face, orofacial movements, etc.), it is important to diagnose somatosensory tinnitus and somatosensory modulation of tinnitus.
3. This chapter discusses how somatosensory tinnitus and somatosensory modulation of tinnitus can be diagnosed, mostly by means of anamnesis and physical evaluation. The chapter provides practical information to the health care professionals regarding such diagnosis.

Keywords Tinnitus • Somatosensory • Central nervous system • Muscle • Cervical spine • Temporomandibular joint

Abbreviations

MTP Myofascial trigger points
PA Pressure algometry
TMJ Temporomandibular joint

Introduction

It is now generally accepted that many incidences of tinnitus can be evoked or modulated by inputs from the somatosensory system, the somatomotor, and the visual-motor systems. Some individual’s tinnitus can be modulated by stimulation of parts of the somatosensory system. Such tinnitus is known as somatosensory tinnitus (other names have been used and the name somatosensory tinnitus has been used for other forms of tinnitus). Somatosensory tinnitus is different from tinnitus that is not affected by activations of non-auditory systems. We will call such tinnitus auditory tinnitus. The effect of somatosensory stimulation on tinnitus can be demonstrated by inducing forceful muscle contractions of the head, neck, and limbs [1–3]; by orofacial movements [4–8]; by applying pressure to myofascial trigger points (MTP) [9]; or by stimulating the skin of the face and hands [4, 10]. Eye movements can often induce tinnitus or modulate existing tinnitus [5]. Among these and other types of modulating factors that have been described (see Chap. 43), the influence of stimulating head and neck regions on the auditory pathways are particularly interesting.

Definition of Somatosensory Tinnitus

The terms “somatosensory tinnitus” and “somatic tinnitus” have been used with different meanings. Efforts are now made to differentiate between somatosensory tinnitus (primary origin in head and neck disorders) and somatosensory modulation of tinnitus.
Somatosensory Tinnitus

Somatosensory tinnitus is suspected when a patient’s history shows at least one of the following events has occurred before the onset of tinnitus:

- Head or neck trauma
- Manipulation of teeth or jaw or cervical spine
- Recurrent pain episodes in head, neck, or shoulder girdle
- Increase of both pain and tinnitus at the same time
- Inadequate postures during rest, walking, working, or sleeping
- Intense periods of bruxism (grinding of the teeth) during day or night

Other forms of stimulation of structures of the head and neck may or may not cause the loudness, pitch, or localization of somatosensory tinnitus to change.

The most important characteristic of somatosensory tinnitus is that it is related to problems of the head and neck, rather than to problems of the ear.

The complexity of somatosensory tinnitus requires that patients with this disorder be evaluated by an integrated team including an experienced dentist and physiotherapist (or similar professions, depending on the organization of local health care structures) to evaluate possible bone and muscular disorders of the face and neck as well as dental problems. Prompt diagnosis is important because treatment must be started as early as possible to obtain the best results (see Chap. 80).

Somatosensory Modulation of Tinnitus

Somatosensory modulation of tinnitus may be perceived as transient changes in loudness, pitch, or localization of the tinnitus. Such modulation of tinnitus can be induced in individuals with either auditory or somatosensory tinnitus. Since the tinnitus of many individuals can be elicited or modulated by somatosensory stimulation (65–80%) [1, 2], all patients who seek help for their tinnitus should be tested for somatosensory modulation. If a patient spontaneously reports that his/her tinnitus changes temporarily during common daily movements of the jaw or neck (opening mouth, clenching teeth, or turning head) or by applying pressure on the temples, mandible, cheek, mastoid, or neck with a fingertip, it is a strong sign that the patient has somatosensory modulation of tinnitus. Other signs of somatosensory modulation may become evident when a professional examines the patient and actively searches for modulation of the tinnitus by applying different kinds of stimuli to different locations on the patient’s body. The patient may mention an immediate change in the loudness of the tinnitus (increase or decrease assessed by a visual analogue scale) or changes in the pitch or the localization of the tinnitus. If this occurs during at least one maneuver involving the somatosensory, somatomotor, or visual-motor systems, it is a strong sign that the patient has somatosensory tinnitus. The effect of many kinds of modulation is short lasting and it is difficult to use a questionnaire to evaluate tinnitus before and after such maneuvers. However, a simple instrument such as a visual analogue scale can be used to quickly evaluate the magnitude of the induced changes in tinnitus.

Different stimuli can be used to detect somatosensory modulation of a patient’s tinnitus, such as active jaw movements (with and without resistance by the examiner), opening and closing the mouth, moving chin forward and backward, or lateralizing chin right and left. Passive muscular palpation can be used to find MTPs or tender points in the masseter, temporalis, and lateral pterygoid muscles. The fatigue test (teeth closed with spatula between them in anterior, right, and left positions for a duration of 1 min) is another useful test that can reveal somatosensory tinnitus. Such movements increase the signals elicited by the tense muscles in the area innervated by the sensory part of the trigeminal nerve, which is anatomically and physiologically connected to the acoustic pathways [11].

Active neck movements (with and without resistance by the examiner) such as moving neck forward and backward, rotation right and left, and lateralization right and left can be used to test if signals from the neck can modulate the patient’s tinnitus. Passive muscular palpation searching for MTPs or tender points in the trapezius (upper fibers), sternocleidomastoid (sternal division), splenius capitis (near the mastoid process), and splenius cervicis are other important tests.

The jaw and upper cervical spine are considered to be a part of an integrated motor system; therefore, observing the posture of the patient is also important for diagnosis and treatment of tinnitus that can be modulated with somatosensory stimulation. For example, if a
patient has the mandible and/or neck protruded forward, this might suggest an attempt to compensate for wrong dental occlusion.

**Gaze-Evoked Tinnitus or Gaze-Modulated Tinnitus**

Eye movements can both cause tinnitus and modulate tinnitus (gaze-evoked tinnitus or gaze-modulated tinnitus) [5]. Influence of gaze on tinnitus can be tested by having the patient start looking straight forward (a neutral position) and then gaze first to the maximal right and then left; after that, looking upwards and downwards. Each position should be maintained for 5–10 s. With the patient placed in a silent environment, changes in tinnitus may occur during each eye movement.

Methods for measuring modulation of tinnitus are not yet standardized. Some centers only describe it as “present” or “absent,” and some use a visual analogue scale (from 0 to 10 or from 0 to 100). Toward standardization, we propose a scale for modulation of tinnitus that is centered at 0 (rest state of tinnitus) and ranging from minus 5 (disappearance) to plus 5 (biggest increase ever thought).

**Other Indications of Somatosensory Tinnitus**

Bone problems of the temporomandibular joint (TMJ) and the neck (osteophits, arthrosis, spondylosis, etc.) may justify the presence of pain and management of accompanying muscular problems (which also can occur in isolation). Such bone problems can seldom be reversed completely, but an approach directed to treat the muscular tension may also lead to control of somatosensory tinnitus and the associated pain (Chap. 80). It may therefore be recommended that the TMJ and neck disorders may be diagnosed and treated to allow better tinnitus control.

Although modulation of tinnitus can occur regardless of the presence of pain, some extra clues to diagnosing somatosensory tinnitus may be added if pain is also included in the following rationale:

- Does the patient have frequent regional pain?
- If so, is it in the head, neck, and shoulder girdle?
- If so, has it a similar duration as the patient’s tinnitus?
- If so, does the patient’s tinnitus become worse when the pain increases?

In general, a patient’s history, together with a clinical examination by a physician might be sufficient for diagnosing temporomandibular and neck disorders in most patients, but complimentary exams (X-ray, computed tomography, or magnetic resonance imaging) may be helpful in reaching a firm diagnosis.

**Myofascial Trigger Points**

MTP are hyperirritable spots in skeletal muscles, which are associated with hypersensitive palpable nodules in taut bands. Basically, there are two kinds of MTPs: active or latent. Active MTP cause clinical pain complaint and refer patient-recognized pain during palpation. A latent MTP may have all the other clinical characteristics of an active MTP and always has a taut band that increases muscle tension and restricts range of motion. It does not provoke spontaneous pain and it is painful only when palpated (upon palpation, the latent MTP can provoke pain and altered sensation in its distribution expected from a MTP in that muscle) [12].

The diagnosis of active MTP is very important for the treatment of myofascial pain syndrome, for pain release, and possibly even for tinnitus relief.

The relationship between MTP, myofascial pain syndrome, and tinnitus has been studied during the preceding years (see Chaps. 9, 43, and 80). Tender points should also be identified and their possible modulation of the patient’s tinnitus must likewise be determined.

Palpation should be performed with sustained deep single-finger pressure during up to 10 s with a spade-like pad at the end of the distal phalanx of the index finger or through pincer palpation (thumb and finger) moving across the muscle band at the hypersensitive area (Figs. 52.1 and 52.2).

When a palpable taut band and spot tenderness is detected, the patient should be asked (Fig. 52.3) if he/she feels any sensation in other area besides the one being pressed upon; if the sensation is like the one that
is a problem to the patient; and finally, if the loudness or pitch of the tinnitus changed.

The ten muscles in the head, neck, and shoulder girdle (infraspinatus, levator scapulae, trapezius, splenius capitis, splenius cervicis, scalenus medius, sternocleidomastoid, digastric, masseter, and temporalis) should all be examined for the presence of MTPs [12]. The examination is expected to reveal whether MTPs were present or not, and if so, in which muscles, in which side of the body relative to the tinnitus, and if palpation of one or more MTPs modulated the patient’s tinnitus. If tender points were present, the patient...
Manual palpation of the muscles is the easiest way of examining patients for somatosensory tinnitus. However, a more objective measurement may be obtained using a hand-held force gauge with a rubber tip to measure the pressure required for eliciting MTP activity [pressure algometry (PA)]. PA has been used to document the tenderness of MTP and is also applied to measure the referred pain threshold and pain tolerance. The reliability and validity of measurements using PA have been established [13, 14]. Pre- and posttherapeutic effectiveness of various procedures on MTP have been assessed by measurement of pressure threshold with PA (Fig. 52.4).

Conclusions

The ability to correctly diagnose somatosensory tinnitus and somatosensory modulation of tinnitus relies mainly on the patient’s history and a thorough physical examination. However, these forms of tinnitus have only recently been studied in detail. Knowledge about the signs of these forms of tinnitus is therefore not generally known. Health care professionals may need to be informed about how to diagnose these forms of tinnitus in their daily routine of treating patients with tinnitus.

Acknowledgments

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References

2. Sanchez, TG; Guerra, GCY; Lorenzi, MC; Brandão, AL; Bento, RF. The influence of voluntary muscle contractions upon the onset and modulation of tinnitus. Audiol. Neurootol. 2002; 7: 370–5.
10. Sanchez, TG; Marcondes, RA; Kii, MA; Lima, AS; Rocha, CAB; Ono, CR; Bushpigel, C. A different case of tinnitus modulation by tactile stimuli in a patient with pulsatile tinnitus. Presented at II Meeting of Tinnitus Research Initiative, Mônaco, 2007; 21–3.
Chapter 53
Differential Diagnosis of Temporomandibular Joint and Masticatory Muscle Disorders in Patients with Tinnitus

Ralf Bürgers, Martin Gosau, Sebastian Hahnel, and Michael Behr

Keypoints

1. This chapter aims at providing non-dental healthcare specialists engaged in tinnitus treatment with a description of a short screening of individuals with tinnitus to clarify the involvement of temporomandibular disorders (TMD) in such patients. A screening test for TMD seems to be reasonable for all tinnitus patients. Patients with a positive TMD screening should be referred to an experienced TMD specialist.

2. TMD short screening consists of an anamnesis, an examination of the temporomandibular joint (TMJ) (jaw motion and TMJ sounds), and an examination of the masticatory muscles (palpation, isometric contraction, and parafunction).

3. Individuals with TMD-related tinnitus suffer more frequently from masticatory muscle pain than from joint syndromes, whereby the majority of individuals with TMD-related tinnitus – in contrast to patients with tinnitus only – describe their tinnitus as fluctuating.

Keywords  TMD • Tinnitus • Short screening

Abbreviations

N  Newton
TMD  Temporomandibular disorder(s)
TMJ  Temporomandibular joint

Introduction

Diagnosis of temporomandibular disorders (TMD) should be made on the basis of the medical history and clinical examination of a patient. It is the opinion of this author that diagnosis of TMD requires a detailed evaluation by a dentist or physician with advanced experience in treating temporomandibular joint (TMJ) and masticatory muscle disorders. This chapter, however, cannot provide a detailed and comprehensive tutorial, neither for diagnosing TMJ and masticatory muscle disorders nor for differentiating between the various forms of TMD; such information can only be obtained from textbooks such as “Temporomandibular Joint and Masticatory Muscle Disorders” by Zarb et al. [1]. The present chapter aims at providing health care specialists of different fields who are engaged in tinnitus treatment a brief guide regarding how to best clarify possible TMD involvement in patients. Patients who have tested positive for TMD and tinnitus should be referred to an experienced TMD specialist for further diagnosis and therapy. The differential diagnosis should rule out pain resulting from other causes but with similar symptoms, such as trigeminal neuralgia and atypical facial pain [2].

It is well documented that individuals with both tinnitus and TMD have more pain and higher dysfunction index scores than individuals with only TMD [3–7]. Therefore, screenings of patients with tinnitus for related TMD can be brief. Patients with a suspicion of having TMD and those without a clear diagnosis may be referred to a TMD specialist for further diagnosis. A short screening can be conducted in approximately 5 min, because it is known that TMD is often accompanied by tinnitus; it may therefore be reasonable to screen all patients with tinnitus for TMD complaints.
TMJ Short Screening Procedure

Often, patients with tinnitus will not relate their “ear symptoms” to possible stomatognathic1 or TMD. Furthermore, many patients with chronic TMD, such as joint clicking or grinding of their teeth, hesitate to consult a dentist or report their symptoms to an otorhinolaryngologist because they regard them (mostly free of pain) as “normal” and not pathogenic. Therefore, screening for TMD should be generally included in examinations of patients with tinnitus [1, 5]. Short screenings to evaluate the incidence of TMD in patients with tinnitus have been described in the literature [8, 9]. The screening described below is adapted to the specific conditions in TMD-related tinnitus. When TMJ involvement is found (positive screening), the patient should be referred to an experienced dentist or TMD specialist for a more detailed diagnosis and TMD therapy [10].

Anamnesis

Ask the patients about pain in the face, jaw, temple, in front of the ear, in the neck, or in the shoulders in the past month and let them point to the area the pain is felt. All patients with tinnitus should be asked if they have had treatments for TMD in the past (such as splint therapy, physiotherapy, medications, etc.), if they have had pain in their temple and tinnitus from mental pressure or medication [2, 11–13].

Patients who have pain in the TMJ or the masticatory muscles (myofascial pain) should have detailed diagnostic tests for TMD.

Jaw Motion

The vertical range and opening pattern of the mandible [10, 14] should be tested (Fig. 53.1). Ask the patient to close their mouth with teeth lightly touching and then slowly open their mouth as wide as possible, even if it is painful.

1. Note if the patient has an initial deviation to one side but corrects to the midline before reaching the maximum mandibular opening or an uncorrected deviation of the jaw to one side.
2. Measure the maximum unassisted opening from the incisal edge of the maxillary central incisor to the opposing mandibular incisor.
3. Ask the patient to do largest possible movements of the mandible: left lateral excursion and right lateral excursion, protrusion, and retrusion.

Patients who have reproducible opening deviations or limited vertical range (<40mm) or with painful mandible movements should have detailed diagnostic tests for TMD.

TMJ Sounds

Ask the patient if they hear any sounds when opening and closing their mouth. Place left index finger over the patient’s right TMJ and the right index finger over the left TMJ; ask the patient to slowly open the mouth as wide as possible, even if it is painful (Fig. 53.2). Palpation has to be done bilaterally.

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1Stomatognatic system: mouth and jaws and closely associated structures.
Record clicking (short) or continuous sounds, like a stone grinding against another stone (crepitus). Ask the patient if the palpation was painful.

More detailed diagnostic tests for TMD are needed in patients who have reproducible TMJ sounds or joint pain during palpation.

In addition to the lateral palpation (preauricular), the TMJ palpation may also be performed from dorsal (intraauricular) direction, with the examiner’s fingers in the right and left acoustic meatus, finger pads orientated forward.

**Masticatory Muscle Tenderness**

Palpation and isometric contraction of the muscles of mastication (Fig. 53.3) may be useful for detecting muscle tenderness. Ask the patient to open their mouth and take the cheek between index finger and thumb. Have the patient lightly clench to identify the masseter muscle and then palpate the whole muscle in a passive state (approximately 2 lb/10 N of pressure). Ask the patient to lightly clench and move the mandible forward and backward to identify the Temporalis muscle and palpate the entire muscle in a passive state (approximately 2 lb/10 N of pressure).

The examiner holds up the mandible with both hands below the chin while asking the patient to open their mouth and hold the position for 1 min (abduction isometric contraction).

Deposit two cotton rolls or swabs between the upper and lower jaw in the region of the premolars and ask the patient to clench and hold with constant pressure for 1 min (adduction isometric contraction).

More detailed diagnostic tests for TMD are needed in patients with masticatory muscle palpation pain or muscle pain during isometric contraction.

Palpation of the remaining masticatory muscles (medial pterygoid, lateral pterygoid, stylohyoid, suprahyoid, and digastricus) may also be done, but localizing these muscles may be difficult even for experienced TMD specialists. Myogelosis and hypertrophies should also be noticed. Movements of the head or cervical spine can cause changes in tinnitus perception [15] (see Chap. 9). Disorders of the neck or cervical spine may influence TMD-related tinnitus and should therefore—if existent—be further examined by a specialist.
Parafunction

Ask the patient for grinding of their teeth (bruxism), clenching and rocking of teeth. Examine the oral cavity

Fig. 53.3 Masticatory muscle palpation. (a) Digital palpation of masseter muscle between index finger and thumb. (b) Palpation of temporalis muscle in toto. (c) Abduction isometric contraction during mouth opening. (d) Adduction isometric contraction through clenching two swabs between the upper and lower jaw in the premolar region

Parafunction\textsuperscript{2}

Ask the patient for grinding of their teeth (bruxism), clenching and rocking of teeth. Examine the oral cavity for hard tissue attritions (not age-based) or soft tissue impressions (of the tongue or inside of the cheek). Hypertrophies of the masticatory muscles (masseter) and asymmetries of the face should be recorded associated with occlusal trauma. Also called parafunctional habits or oral habits. (From Mosby’s Dental Dictionary, 2nd edition. © 2008 Elsevier, Inc. All rights reserved.)

\textsuperscript{2} Parafunction: the habitual movements (e.g., bruxism, clenching, and rocking of teeth using teeth for tools) that are normal motions associated with mastication, speech, or respiratory movements and that result in worn facets and other problems
because they are indicators for parafunctions. More detailed diagnostic tests for TMD are needed in patients with signs of parafunction. Patients should be asked if their tinnitus changes during mandible movements or palpation of joint and masticatory muscles.

**Special Considerations in Patients with Tinnitus**

One of the difficulties in diagnosing patients with both TMD and tinnitus is to distinguish patients who have tinnitus because of TMD from patients who hear tinnitus independently of their TMD. In patients with TMD-related tinnitus, therapy should primarily focus on TMD. Often the tinnitus will abate after successful TMD treatment. In patients whose TMD and tinnitus are independent of each other, TMD therapy is unlikely to affect the tinnitus. Such patients should therefore be referred to a tinnitus specialist. Patients with both symptoms are often classified as having tinnitus or TMD.

Individuals with tinnitus have been described to suffer more frequently from masticatory muscle pain (and especially from myofascial pain) than from joint symptoms [3, 6, 16]. However, Henderson et al. reported that tinnitus does not occur more frequently in patients with TMD involving disc displacement than in patients with physiological disc position [17]. Therefore, TMD diagnosis and related short screenings in patients with tinnitus should particularly focus on examining the masticatory muscle system and muscular disorders. Most individuals with TMD-related tinnitus describe their tinnitus as fluctuating, and TMD occlusal splint therapy has been found significantly more effective in patients with fluctuating tinnitus than in patients with continuous and severe tinnitus [16, 18]. Patients with TMD might be diagnostically separated from patients with tinnitus-related TMD because of the character of their disorder (joint disorder vs. muscle pain) (Fig. 53.4). So far, such unequivocal signs that should allow distinction between the pathology in these two groups have not been described. The quality of the tinnitus might be a predictable indicator for the involvement of the TMJ and masticatory muscle system, which should therefore be examined even more thoroughly in patients with tinnitus of a specific quality (fluctuating tinnitus).

**Conclusions**

There are several reasons why testing for TMD would be beneficial to patients with tinnitus, especially for patients with TMD who do not have a known cause of tinnitus. It is known that patients with tinnitus benefit from efficient treatment of their TMD and, therefore, patients with tinnitus should be screened for TMJ problems, as every patient with TMD should be asked if they have tinnitus [1, 19].

**References**

Keypoints

1. Psychiatric comorbidity occurs frequently in patients with tinnitus, especially in moderate to severe forms.
2. Depression and anxiety are the most frequently found comorbid conditions.
3. Tinnitus severity and impairment in quality of life can be linked to psychiatric symptoms.
4. For every professional who treats tinnitus patients, it is important to recognize signs of potential psychiatric comorbidity.
5. Potential warning signs are high scores in tinnitus questionnaires. Screening instruments that are easy to use may help to identify comorbid psychiatric disorders such as depression or anxiety.
6. Further diagnosis and treatment should be done by specialists such as psychiatrists or psychologists.
7. Patients who appear suicidal should be promptly referred to a psychiatrist.

Keywords Psychiatric comorbidity • Quality of life • Suicidality • Diagnostic screening • Depression • Anxiety disorder

Introduction

Chronic tinnitus represents a frequent condition experienced by about 10–20% of the general population (see Chaps. 5 and 6). One to two percent of individuals with tinnitus have reduced quality of life [1]. Psychiatric comorbidity occurs especially in individuals with severe tinnitus [2]. Major depression, anxiety, and somatoform disorders are frequently reported as comorbid conditions to tinnitus [3–7]. However, psychosis and personality disorders may also be associated with tinnitus [8]. Major depressive disorder and anxiety disorder occur most frequently in individuals with chronic disabling tinnitus; a prevalence rate of 60% or more has been reported [5, 9, 10]. From a clinical point of view, it is important to note that chronic disability and suffering in tinnitus patients is frequently linked to concomitant depressive symptoms [11] and improvement of depression is paralleled by an improvement of functional disability [12]. Several studies have shown that tinnitus severity and tinnitus-related distress is correlated with depression [4, 6, 7]. Psychiatric comorbidity, especially depression and anxiety disorders, is a common phenomenon in tinnitus patients and adds considerably to the suffering and impairment in quality of life. It is, therefore, important that clinicians who treat tinnitus patients are observant to any comorbid psychiatric symptoms, especially depression and anxiety, and provide treatment of tinnitus that takes any affective symptoms into account. Effective treatment regimes for tinnitus aimed at the cause of the patient’s symptoms are still missing, and treatment of comorbid psychiatric disorders can substantially reduce the burden of the disease and improve the quality of life of individuals with tinnitus. The difference between a severely suffering tinnitus patient and a well-compensated individual is sometimes adequate treatment of a psychiatric comorbidity. Thus, it is important that comorbid psychiatric disorders are diagnosed and efficiently treated.
Management of Psychiatric Comorbidity

In this chapter, we will focus on the detection of comorbid psychiatric disorders; for the exact clinical description and the clinical management, we refer to Chaps. 62–64.

Detection of Psychiatric Comorbidity in a Nonpsychiatric Setting

The majority of tinnitus patients do not show any signs of psychiatric comorbidity, especially those who have mild forms of tinnitus, which are well managed. However, if tinnitus patients suffer from depression or anxiety, it is important to recognize and treat these disorders.

In clinical practice, patients primarily seek medical help for their tinnitus, not of their depression or anxiety. Depending on the health care system, patients with tinnitus seek help from a general practitioner, an otolaryngologist, or an audiologist, but rarely from a psychiatrist. Patients seek help because of their tinnitus; additional symptoms, which may be present are only mentioned in passing. Most patients are reluctant to talk about affective symptoms such as mood disturbances or anxiety in a nonpsychiatric setting and that increases the likelihood that these symptoms are overlooked, as has been shown in large survey studies. In practitioner offices, correct diagnosis of a major depression represents a substantial problem; patients with mild to moderate depression are particularly at risk of being overlooked [13]. Clinicians who are not specialized in psychiatry yet caring for tinnitus patients should ask the questions: (a) What are the signs that a psychiatric comorbidity is likely when treating a patient with tinnitus? (b) What are reasonable screening instruments for psychiatric disorders? (c) What should be done if comorbid psychiatric disorders are suspected? (d) How patients with risk of suicide should be managed?

Warning Signs of Potential Psychiatric Comorbidity

Most individuals with tinnitus do not suffer substantially and have little or no impairment of their quality of life. They are also typically able to work and are not restricted by their tinnitus in their everyday life. Some patients with tinnitus seek medical help because they are concerned their tinnitus may be a sign of a dangerous disease such as a brain tumor. Information about the pathophysiology of tinnitus, counseling (see Chap. 70), and a suitable test to rule out a vestibular schwannoma is sufficient, and any psychiatric comorbidity appears to be unlikely in these cases. Severe and disabling tinnitus, however, is often accompanied by symptoms such as depressed mood and anxiety. Tinnitus severity can be assessed either by scales, which are easy to perform [14], or by a validated questionnaire (see Chap. 42). Patients with grade III and IV of Biesinger (Table 54.1) or high scores in tinnitus questionnaires (i.e., a total score of more than 47 in the tinnitus questionnaire of Goebel and Hiller [15] or more than 37 in the Tinnitus Handicap Inventory) should be examined with focus on signs of depression or anxiety [16].

What are Reasonable Screening Instruments for Psychiatric Symptoms?

The most frequent symptoms of depression are depressed mood, loss of interest, and sleep disorders. However, in a nonpsychiatric setting, circumstances such as time limitations often do not allow an extensive interview to specifically explore all potential symptoms of depression or anxiety. Also, specific training and experience is required for assessing affective signs and should be done by a psychiatrist or psychologist. However, everybody who treats tinnitus patients should be familiar with screening instruments for frequent psychiatric disorders, which are based on a few key questions and are easy to perform.

Table 54.1 Tinnitus grading according to Biesinger et al. [14]

<table>
<thead>
<tr>
<th>Grade</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Tinnitus is well compensated. No psychological strain</td>
</tr>
<tr>
<td>II</td>
<td>Tinnitus appears only in silence and is disturbing during periods of stress and pressure</td>
</tr>
<tr>
<td>III</td>
<td>Tinnitus interferes continuously in the private and professional area. Emotional, cognitive and physical disturbances occur</td>
</tr>
<tr>
<td>IV</td>
<td>Tinnitus leads to the complete decompensation in the private area; disability</td>
</tr>
</tbody>
</table>
Psychologic/Psychiatric Assessment

For depression, the following two questions have been proposed:

1. During the past month have you often been bothered by feeling down, depressed, or hopeless?
2. During the past month have you often been bothered by little interest or pleasure in doing things [17]?

If the patient answers with “yes” to one of the both questions, depression is likely and referral to a psychiatrist or psychologist should be made. Similar screening questions for anxiety disorders are used in standardized, semi-structured diagnostic interviews like the MINI International Neuropsychiatric Interview (M.I.N.I.; [18]; see Table 54.2) and can be used for screening for anxiety disorders.

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Screening question</th>
</tr>
</thead>
<tbody>
<tr>
<td>Panic disorder</td>
<td>Have you, on more than one occasion, had spells or attacks when you suddenly felt anxious, frightened, uncomfortable or uneasy, even in situations where most people would not feel that way? Did the spells surge to a peak, within 10 min of starting?</td>
</tr>
<tr>
<td>Agoraphobia</td>
<td>Do you feel anxious or uneasy in places or situations where you might have a panic attack or panic-like symptoms, or where help might not be available or escape might be difficult: like being in a crowd, standing in a line (queue), when you are away from home or alone at home, or when crossing a bridge, traveling in a bus, train or car?</td>
</tr>
<tr>
<td>Social phobia</td>
<td>In the past month were you fearful or embarrassed being watched, being the focus of attention, or fearful of being humiliated? This includes things like speaking in public, eating in public or with others, writing while someone watches, or being in social situations</td>
</tr>
<tr>
<td>Specific phobia</td>
<td>In the past month have you been bothered by recurrent thoughts, impulses, or images that were unwanted, distasteful, inappropriate, intrusive, or distressing? (e.g., the idea that you were dirty, contaminated or had germs, or fear of contaminating others, or fear of harming someone even though you didn’t want to, or fearing you would act on some impulse, or fear or superstitions that you would be responsible for things going wrong, or obsessions with sexual thoughts, images or impulses, or hoarding, collecting, or religious obsessions</td>
</tr>
</tbody>
</table>

What to Do if a Patient with Tinnitus is Suspected to also Have Depression and Anxiety

A patient who is suspected of suffering from depressive symptoms, anxiety, or any other psychiatric disorder should be referred to a psychiatrist for further diagnostic and therapeutic management. In clinical practice, this may sometimes be difficult, since patients may interpret the referral to a psychiatrist as a sign they are not taken seriously or are considered to be “crazy.” This can be easily avoided by careful explanation that tinnitus can cause a high amount of distress and that the treatment concept will include approaches for reducing tinnitus (e.g., hearing aids, noise generators). However, the patient should understand that the amount of suffering induced by tinnitus should be treated by specialists (e.g., by some form of cognitive behavioral therapy or by pharmacologic treatment). Also, a close collaboration with psychiatrists and psychologists, which are interested in tinnitus and have experience with diagnosis and management of tinnitus patients, will make it easier for otologists or audiologists to refer their patients for psychiatric diagnosis and therapy.

How to Manage a Suicidal Patient?

The suicidal patient is a rare but clinically important issue in the management of tinnitus patients. Signs of suicidal thoughts must always be taken serious. Individuals with chronic, severe tinnitus have an increased risk of suicide, especially when comorbid depressive disorders are present [19–21]. The important questions in this context are first, how to find out if a patient is at risk of suicide and second, how to find out whether asking about the patient’s suicidal thoughts may prompt the patient to commit suicide. The most prominent risk factor for suicide and suicidal ideas is a depressive disorder [22, 23]. Hence, a severe depressed mood, which cannot be modulated and is accompanied by social withdrawal, is an important warning sign of high risk of suicide. If one suspects that a patient is at
risk of suicide, the seriousness of the patient’s suicidal thoughts should be evaluated by directly asking the patient about suicidal ideations or even asking if the patient has concrete plans of how to commit suicide. Discussing the issue of suicide with a patient should not increase the risk of them actually committing suicide. On the contrary, most patients feel relieved to have the opportunity to talk about their thoughts. Although there is no general rule of how to manage patients who are suicidal, one possible approach to this sensitive area is to first talk about passive suicidal ideas. This may be introduced by asking, for example, “In the past month did you think that you would be better off dead or wish you were dead?” Further screening questions to estimate the risk that the patient will actually commit suicide are given in Table 54.3. If it becomes clear from the clinical interview that the patient is suffering from suicidal ideas, the patient should be immediately referred to a psychiatrist.

A patient with serious suicidal and concrete plans of how to commit suicide is an emergency, in which many physicians would recommend hospitalization for treatment.

### References

15. Goebel, G, Miller, W [The tinnitus questionnaire A standard instrument for grading the degree of tinnitus Results of a multicenter study with the tinnitus questionnaire] HNO, 1994;42(3):166–72
Part V
Clinical Characteristics of Different Forms of Tinnitus
Chapter 55
Introduction

Berthold Langguth, Dirk De Ridder, and Tobias Kleinjung

Keypoints

1. Tinnitus has many forms and many concomitant symptoms.
2. Specific subforms of tinnitus, which are characterized by phenomenological properties of the tinnitus sound, acuity, a specific time course, specific etiologies, or specific accompanying symptoms, require specific diagnostic and therapeutic management.

Keywords Subforms • Etiology • Chronicity • Comorbidity • Concomitant symptoms • Types • Tinnitus

Introduction

There is increasing consensus among clinicians that tinnitus is not a disease entity. Rather, there are many different forms of tinnitus that vary in their pathophysiology and probably also in their response to treatment interventions [1]. This, in turn, implies that differentiation of the different forms of tinnitus is essential for successful therapeutic management. Differentiation according to clinical characteristics seems to be the best feasible strategy.

This means that diagnostic and therapeutic management should be individualized according to phenomenological characteristics of the tinnitus sound (e.g., pulsatile or non-pulsatile), comorbidities (e.g., vertigo, headache, and psychiatric symptoms), time course (e.g., acute tinnitus with hearing loss), or etiologic aspects (e.g., posttraumatic tinnitus). The chapters of this section deal with the most clinically relevant specific forms of tinnitus and their diagnostic and therapeutic management.

Acute sudden hearing loss with tinnitus (see Chap. 56) represents a specific subform, which requires immediate attention. In such a situation, therapeutic activities are primarily directed toward restoration of hearing. Based on data from animal models, it is assumed that after acute onset, there is a short therapeutic window for specific therapies [2, 3].

Hyperacusis and phonophobia occur frequently together with tinnitus [4]. An exact description is given in Chaps. 3 and 4. The focus of Chap. 57 is the management of tinnitus patients where hyperacusis and phonophobia are main complaints.

Both pulsatile and paroxysmal tinnitus have to be considered as specific entities and point at characteristic underlying pathologies. These subforms of tinnitus require specific diagnostic and therapeutic management which is described in the chapters on pulsatile tinnitus (Chap. 59) and intermittent tinnitus (Chap. 58).

Low-pitch tinnitus co-occurring with fluctuating vertigo and low-frequency hearing loss is characteristic for Ménière’s syndrome with endolymphatic hydrops as an underlying pathology (see Chap. 60).

When tinnitus is accompanied by headache, pathologies should be considered, which result in increased or reduced intracranial pressure. These include space-occupying lesions and pseudotumor cerebri as well as...
low intracranial pressure syndrome. The diagnostic and therapeutic management of these and other syndromes are outlined in the chapter on tinnitus with headache (Chap. 61).

Severely impaired tinnitus patients frequently suffer from psychiatric comorbidities with depression, anxiety, and insomnia being the most frequent [5]. Even if a patient presents primarily because of his tinnitus, sometimes the management of the psychiatric comorbidities is in the foreground. This is definitively the case when a patient reports acute suicidal ideation. The different psychiatric comorbidities and their therapeutic management are covered in Chap. 62 with the subchapters Tinnitus and Depression (Chap. 63), Tinnitus and Anxiety (Chap. 64), and the chapter Tinnitus and Insomnia (Chap. 65).

The last two chapters of this section are devoted to tinnitus with a specific etiology. Whenever tinnitus occurs in conjunction with a traumatic event, specific diagnostic management is indicated (see Chap. 66) [6]. This is not only necessary for the best possible treatment of tinnitus itself but also to avoid further complications since tinnitus after trauma can be a symptom of a severe underlying condition that may become life threatening if left untreated (e.g., carotid dissection) [7]. A separate chapter is devoted to blast injuries (Chap. 67) as a specific form of posttraumatic tinnitus. This form of tinnitus is of high clinical relevance, since tinnitus has become one of the most relevant warfare-related health problems in the last few years [8]. Furthermore, blast injuries are a particular diagnostic challenge, since the tinnitus-inducing mechanisms may include noise, ear, head, neck, and emotional trauma [9, 10].

References

1. Möller, AR. Tinnitus: presence and future. Prog Brain Res, 2007;166: 3–16
6. Folmer, RL, Griest, SE. Chronic tinnitus resulting from head or neck injuries. Laryngoscope, 2003 May;113(5): 821–7
Chapter 56
Sudden Hearing Loss and Tinnitus

Carlos Herráiz

Keywords
Tinnitus • Sensorineural hearing loss
• Hyperacusis • Sudden deafness • Transtympanic
• Steroids

Abbreviations
ABR Auditory brainstem response
AIEDA Autoimmune inner ear disease
FTA Fluorescent treponemal antibody absorbed (FTA-ABS) test for syphilis
MRI Magnetic resonance imaging
RCT Randomized clinical trials
SNHL Sensory neural hearing loss
SSNHL Sudden sensory neural hearing loss
TTS Transtympanic steroids

Introduction

Hearing loss can occur suddenly when the ear canal becomes occluded or the middle ear becomes damaged from trauma. However, the term sudden hearing loss is mainly used for suddenly occurring sensory neural hearing loss. Sudden sensory neural hearing loss (SSNHL) was first described by De Klein in 1944 [1]. SSNHL is a dramatic condition for the patient that twenty-first century medicine still has no explanation of; there is no known cure. The mechanisms, etiology, and the treatment remain hypothetical. The SSNHL definition is also controversial among authors. The most detailed criteria have been proposed by Stokroos [2], who described SSNHL as an acute deafness with abrupt onset, generally within 3 days, of more than 30-dB hearing loss at three consecutive frequencies. Different authors have used different definitions of SSNHL [3].

Incidence of SSNHL

SSNHL occurs suddenly, over less than 3 days; it normally affects only one ear. The incidence of SSNHL has been reported to be 5–20 per 100,000 inhabitants per year in the United States [4] and 8–14.6 in Holland [5]. A recent epidemiological study conducted during 2004 in Saxony, Germany, with a population of almost half a million, showed an incidence of 160 per 100,000 inhabitants [6].

Some incidences of SSNHL have been reported in childhood. The prevalence is higher in young and healthy individuals. Many individuals with SSNHL recover spontaneously. There are many causes of hearing loss similar to etiologies that could be regarded as other forms of sensorineural hearing loss (SNHL) [4], and they have to be ruled out before the diagnosis of SSNHL is made. Endolymphatic hydrops, according to the Fetterman series, is the second most common cause of acute hearing loss (5.5%) after idiopathic sudden hearing loss. Ménière’s disease was the next most frequent diagnosis (1.9%), followed by vestibular schwannoma (1.7%), perilymphatic fistula (0.7%), and autoimmune inner ear disease (0.6%). Other authors find that vestibular schwannoma accounts for 4% of acute unilateral hearing loss [7].
Causes of SSNHL

Up until now, the etiological of most SSNHL is unknown. In only 10% of the SSNHL is it possible to find a plausible cause [4]. Many causes have been suggested, such as disturbances in the cochlear blood flow, inflammatory processes secondary to viral infections, and autoimmune reactions.

Decrease of the Inner Ear Blood Flow

Blood flow in the inner ear may be reduced because of hemorrhage or arterial occlusion, which may occur from thrombosis or vascular spasm. Pathological studies describe processes of fibrosis and cochlear ossification in individuals who have had sudden deafness [8]. The vascular causes may be suspected in individuals with a history of previous thrombi-embolisms, atherosclerosis, heart surgery, or thrombocytopenia due to aplastic anemia or leukemia.

Spontaneous recovery makes impairment of the cochlea’s blood supply for longer than 1 h an unlikely cause [9].

Rupture of the Cochlear Membranes

The rupture of the cochlear membranes causes contact between the perilympha and the endolympha, altering the electrolytic balance and resulting in damage to hair cells.

A perilymphatic fistula to the middle ear is known to cause sudden hearing loss. Intense physical exercise, Valsalva maneuvers, or barotrauma can cause rupture of the oval or round window membrane. It has been estimated that perilymph fistulae may explain almost thirty percent of the SSNHL [10].

Autoimmune Inner Ear Disease

Autoimmune inner ear disease (AIED) is characterized by rapidly progressive bilateral hearing loss, usually symmetrical and fluctuant, although some individuals experience sudden hearing loss in only one ear.

Viral Theory

The viral theory is the most referred in the literature. Viral infections like mumps, rubella, herpes, or spuma retrovirus have been related to sudden hearing loss, although there is no clear evidence to confirm this theory [11–13].

Clinical Course

Sudden hearing loss starts with a rapidly progressive hearing impairment, either suddenly or within a few hours, often when the person wakes up. Sixty-three percent of people with SSNHL have ear pain initially and 41% experience aural pressure during a few days [11]. Tinnitus appears in 91% of individuals with SSNHL [4]. Tinnitus has been reported to develop days before the SSNHL occurred or it can occur simultaneously or days after to the SSNHL. Some individuals describe facial paresthesia.

Vertigo has been reported to occur in 43% of individuals with SSNHL [11]. Some patients refer a rotary motion with vegetative manifestations during a few days. Often unsteadiness and involuntary movement of the body toward the affected side occur for weeks. The presence of spontaneous nystagmus in SSNHL has been reported to occur in half of individuals with SSNHL. The recurrence rate of hearing loss in long-term follow-up has been reported to be significantly higher (51.2%) in the group who had spontaneous nystagmus than in the individuals without nystagmus (27.9%) [14].

Diagnosis

A diagnosis of individuals with rapidly occurring hearing loss requires complete auditory examination: tone and speech audiometry, tympanometry, and stapedial reflex test. Auditory brainstem responses or MRI to rule out retrocochlear diseases and laboratory tests such as the antinuclear antibodies, erythrocyte sedimentation rate, and tests for rheumatoid factor have been proposed for diagnosis of SSNHL and to detect treatment responders [15]. However, none of the
laboratory parameters have been proven to have a high sensitivity or specificity. A test to rule out syphilis (FTA) is also recommended [16].

The assessment of the vestibular system may be useful to detect possible vestibular complications for prognosis. Vestibular-evoked myogenic potentials (VEMP) have demonstrated saccular damage in patients with SSNHL without vertigo, suggesting a saccular deterioration in those patients with profound high-frequency hearing loss [17].

**Prognosis**

A spontaneous recovery has been described in 45–65% of the cases [11]. Some individuals with SSNHL have a complete recovery while most have partial improvement of hearing. Specific factors that affect the prognosis of SSNHL are the severity of the hearing loss; a greater impairment on high frequencies or the presence of vestibular symptoms significantly reduces the prognosis [11, 18]. However, Fetterman [4] did not find that the audiometric profile made any differences regarding prognosis, but “U”-shaped audiograms predict higher fluctuations and recurrences of the episodes of SSNHL. Age or speech recognition threshold did not influence the course of the disease.

**Treatment**

The high rate of spontaneous recovery and the difference in definition of SSNHL makes it difficult to compare the results presented in published studies [19].

**Blood Flow Increase**

Vasodilatation: histamine, verapamil, papaverine, novocaine, nicotinic acid, naftidrofuryl, Egb 761.

Studies that have followed a valid design do not show significant differences between treated individuals and control groups [20–22].

Reduction in the blood viscosity: Dextran, papaverine, pentoxifiline.

A recent multicenter and randomized study evaluated the benefits of rheopheresis, a method to reduce the plasma viscosity and improve microcirculation for treatment of SSNHL [23]. The rheopheresis group (two sessions within 3 days) was compared to a group that received steroid treatment (methyl-prednisolone 250 mg per day, 3 days and tapered oral dosing) and to intravenous hemodilution (500 ml 6% hydroxyethyl starch plus 600 mg pentoxifyline per day during 10 days). There was not a placebo control group in this study. None of the tested treatments were superior regarding providing overall good recovery of hearing.

**Defibrinogenase Therapy: Baxtrobin**

Administration of baxtrobin, a trombine-like enzyme that reduces the levels of fibrinogen and the blood viscosity [24] did not present better results than expected with placebo.

**Anti-inflammatory Treatment: Corticosteroids**

**Systemic Steroids**

Corticosteroids are the most effective treatment for SSNHL. A placebo-controlled study demonstrated the efficacy of dexamethasone or methyl-prednisolone. Seventy-eight percent of the patients with moderate and severe hearing loss who received such treatment had partial or total recovery of hearing compared with placebo [25].

In a descriptive study by Moon [26], SSNHL participants who showed any improvement after early steroid therapy were analyzed to evaluate the beginning time and the plateau time of hearing improvement. It was shown that 93.1% had an onset of improvement within 14 days of beginning the treatment. Complete recovery or completed improvement was achieved in 80.4% of the participants within 1 month and in 92.2% within 2 months after treatment [26].

After 1 month, the possibility of improvement decreases [2], but Stokroos did not find differences in starting a treatment within the first 24 h and during the
first 10 days [2]. Better recovery was found in participants who had the most hearing loss around 4 kHz [4].

**Transtympanic Steroid Therapy**

Many publications regarding randomized clinical trials (RCT) have demonstrated the benefit of transtympanic steroids (TTS) as a rescue treatment after systemic steroids for SSNHL. Methyl-prednisolone showed the most promising profile, when considering drug concentration in the endolymph [27]. Battaglia obtained better results in those patients who received a combined therapy, oral steroids (60 mg per day, 7 days) plus TTS (dexamethasone 12 mg/ml once per week, 3 weeks), compared to the group that received TTS plus oral placebo. This last combination was more effective than oral steroids and transtympanic placebo [28].

**Antiviral Therapy**

There was no difference in the benefit from treatment with antiviral drugs (acyclovir) or administration of steroids [5], nor have other studies with valacyclovir [29, 30] shown any benefit of the antiviral drug for SSNHL.

**Hyperbaric Oxygen Therapy**

Administration of hyperbaric oxygen treatment has been described but is controversial [2].

**Surgery**

If a perilymphatic fistula is the cause of SSNHL, surgical treatment of the fistula can improve hearing [31].

**Other Treatments**

Ozone therapy (autohaemotherapy) has been tried in a RCT for sudden hearing loss [32]. A 100 ml of the patient’s own blood with a gaseous mixture of oxygen and ozone was re-injected twice a week for 10 sessions. Seventy-seven percent of the treated patients showed a significant hearing recovery compared to 40% in the placebo group. Pure-tone averages and speech reception thresholds were also significantly better.

**Tinnitus and SSNHL**

Tinnitus is a common symptom in SSNHL. Approximately 91% of individuals with SSNHL report that they have tinnitus in the affected ear or in both ears [4]. Tinnitus occurs at the same time as hearing loss in some individuals with SSNHL and may be the first symptom before hearing loss. Tinnitus begins some days after the hearing impairment in some individuals. The tinnitus may be caused by sound deprivation caused by the hearing loss, which is known to be able to start central nervous system reorganization processes that can lead to tinnitus (see Chaps. 10, 12 and 21) and hyperacusis (see Chaps. 3 and 4).

**Tinnitus Characteristics**

We have previously shown [32] that 6.6% of the first 213 patients referred to our tinnitus clinic had SSNHL. Tinnitus onset was sudden in approximately 92% of the patients. The intensity of the tinnitus fluctuated in 46% of patients. Tinnitus psychoacoustical characteristics are shown in Table 56.5.

The tinnitus in SSNHL can imply a greater handicap than the hearing loss [2]. When the hearing resolves, either through treatment or spontaneously, the tinnitus may improve or disappear. Tinnitus may be a prognostic sign for the hearing loss [33].

**Tinnitus Management**

Tinnitus that accompanies SSNHL can be treated in a similar way as other forms of tinnitus (see chapters in Section V). All treatments for sudden hearing loss can be effective in treating the tinnitus. The use of steroids, vasodilatation drugs, and procedures or the hyperbaric oxygen therapy is often used when tinnitus accompanies SSNHL. At early stages of SSNHL,
sound stimulation is beneficial because it can prevent reorganization processes secondary to sound deprivation in the auditory nervous system. As remapping could be the physiological substrate for tinnitus development, customized sound enrichment would help to decrease the possibility of tinnitus and hyperacusis [34, 35] (see Chaps. 74, 75 and 76).

**Conclusion**

Most forms of SSNHL have no known cause. In management of patients with SSNHL, it is important to rule out other causes. Retrocochlear diseases, such as vestibular schwannoma and other central nervous system tumors, have been described to occur in 4% of SSNHL. They can be ruled out through tests such as ABR or MRI. Published results regarding the prognosis of SSNHL vary among studies. Some studies show 45–65% spontaneous recovery. The severity of the hearing loss, the audiometric profile, and the presence of vestibular symptoms affect the prognosis. Delay on starting treatment is associated with a worse prognosis. Steroid treatment is proven to be the most effective treatment, although it is not largely effective. Recent studies show promising results regarding the efficacy of steroids delivered through the eardrum. More than 50% of the patients showed a significant improvement of such treatment when administered after the failure of conservative therapy. Tinnitus accompanies most incidences of SSNHL. Tinnitus usually improves when hearing is partial or totally recovered, and individuals can also benefit from other forms of tinnitus treatment when there is no improvement of hearing.

**References**

1. De Klein A. Sudden complete or partial loss of function of the octavus-system in apparently normal persons. Acta Otolaryngol (Stockh) 1944; 32: 409–429
18. Byl FMJ. Sudden hearing loss: eight years of experience and suggested prognostic table. Laryngoscope, 1984; 94 (5 Pt 1): 647–51
Keypoints

1. Hyperacusis is a decreased sound tolerance.
2. Prevalence of the disease is described in 9–15% of the population, but increases among tinnitus patients.
3. Pathophysiological mechanisms involve some disruptions in the amplification and regulation processes of the external hair cells, disorders of the efferent system (medial and lateral olivocochlear pathways), or effects to the central sound processing at the subcortical level.
4. The role of some neurotransmitters (serotonin, GABA), which are also involved in other hyperacusis-related diseases (migraine, depression), can be relevant in this disorder.
5. Other theories confirm the effect of the endorphins that activates the excitatory function of the glutamate, the main auditory neurotransmitter, increasing its toxicity.
6. The activation of the limbic and autonomic nervous systems produces the emotional reaction of the hyperacusis (anxiety, fear, and depression).
7. Proposed treatments are based on acoustic stimulation: progressive introduction of white sound (tinnitus retraining therapy TRT) and customized sounds based on the damaged hearing frequencies.
8. Noise generators and hearing aids can be fitted in severe cases.
9. The role of some drugs involved in the metabolism of serotonin and GABA opens new approaches for the management of hyperacusis.

Keywords

- Tinnitus • Decreased sound tolerance • Hyperacusis • Recruitment • Phonophobia • Efferent system • Tinnitus retraining therapy • Hearing aid

Abbreviations

- LDL: Loudness discomfort level
- DST: Decreased sound tolerance
- OHC: Outer hair cells
- IHC: Inner hair cells
- MOCB: Medial olivocochlear bundle
- LOCB: Lateral olivocochlear bundle
- 5HT: Serotonin
- ABR: Auditory brain responses
- THS: Test of Hypersensitivity to Sound
- BBNG: Broad band noise generators

Introduction

Hyperacusis is defined as a decreased tolerance to environmental sounds or the abnormal avoidance response to sounds that they are not annoying to the general population (see Chap. 3). It is a disorder of the normal amplification process of the auditory pathways. A decrease in the loudness discomfort levels (LDL) to environmental noise is observed in individuals with hyperacusis scores below 90 dBHL for some authors [1] or below 100 dBHL according to others [2]. Auditory hypersensitivity affects all sounds, although some specific noises can be more annoying according to their frequency spectrum or intensity.

The hyperacusis has to be distinguished from other symptoms that could co-exist simultaneously or develop as isolated forms. Misophonia (from the
Greek “miso: hate”) is a “dislike of certain specific sounds,” and is different from phonophobia – a fear of certain sounds [2] (see Chap. 4). The anatomical and physiological basis is generally unknown, and these clinical entities have been regarded as belonging to the field of psychology. Phonophobia and misophonia are related to the type or the source of the sound and not specifically to its loudness. Hyperacusis is an abnormally low tolerance of sounds and may have to do with faulty gain control in the auditory pathways causing an abnormal activation of emotional reactions from the limbic and autonomous systems. Conversely, phonophobia is an abnormal reaction from the limbic and autonomous systems with normal auditory neural activity.

Recruitment of loudness is a pure cochlear physical phenomenon that depends on the outer hair cells. It is caused by the stimulation of the neighboring neural fibers to the damaged cochlear areas after exposure to intense sounds. There is a breakdown in the relation between the stimulus loudness and the intensity of the patient’s acoustic sensation. The result is a distortion of, and an annoyance to, the sound.

There are a few epidemiological studies related to hyperacusis and decreased sound tolerance (DST). Fabijanska performed a wide study, sending a specific questionnaire to the general population by postal mail. Of 10,349 returned questionnaires, the study showed that 15.2% of the population referred hypersensitivity to sound [3].

The study published by Andersson in 2002 was conducted in Sweden through the Internet. Nine percent of the 595 responders reported a DST. These data were confirmed through postal mail to 589 individuals, where 8% of the sample showed the same results [4].

Some studies have described the prevalence of DST among tinnitus patients. Between 40% [2] and 59% [5] in a tinnitus clinic sample reported symptoms of hyperacusis. The prevalence of tinnitus in DST patients rises up to 86% [6].

**Mechanisms of Hyperacusis**

The amplification of the acoustic pressure wave from the active movements of the outer hair cells (OHC) facilitates the stimulation of the inner hair cells (IHC). This mechanism can be damaged due to an increased amplification of sound from the OHC [2]. Hyperexcitability of these cells would overstimulate the action of the IHC. The OHC’s active movements would excessively amplify a sound of moderate intensity and, therefore, it will be annoying. Distortion product measurements in these patients would show increased values [2, 7].

Contralateral otocoustic emission suppression through white noise stimulation is a useful tool to test the efferent system function. We found some abnormalities in the medial olivocochlear bundle (MOCB) pathways as the cause of DST [8]. Other authors, such as Baguley, have not found any change in LDL scores after section of olivocochlear fibers (efferent fibers) when performing a vestibular neurectomy for disabling vertigo (the MOCB travels with the vestibular nerve at the point where it is sectioned) [9].

The lateral olivocochlear bundle (LOCB) originates in the lateral superior olivary complex and innervates through unmyelinated axons, the primary afferent dendrites of the cochlear nerve near their synapses with inner hair cells. LOCB terminals are more complex, with evidence for cholinergic, GABAergic, dopaminergic, and peptidergic transmission [10]. Activation of the LOCB can evoke either slow enhancement (cholinergic) or suppression (dopaminergic) of auditory nerve response. LOCB feedback maintains the binaural balance in neural excitability required for accurate localization of sounds in space [11]. Its function has been associated with the control to glutamate excitotoxicity in afferent nerve terminals in acute acoustic injury [12], and it has a protective effect over neural damage from intense sound exposition, mainly based on the dopaminergic regulation.

DST could be caused by LOCB impairment. Clinical diagnosis of LOCB function is based on auditory brainstem responses (ABR) [13], which would increase by ipsilateral stimulation and decrease in response to contralateral stimulation. Otoacoustic emissions will not be affected because the LOCB does not affect the function of the outer hair cells. It can be hypothesized that the dopaminergic LOCB synapses would be a suitable target for treatments. It has been shown that after acute acoustic injury, perfusion with dopaminergic agonists reduces cochlear damage [12].
Other possible mechanisms of peripheral disorders that could cause DST would be the recruitment phenomenon; although recruitment of loudness is regarded to be different from hyperacusis, it is included in DST. Cochlear hearing loss (which occurs in Ménière’s disease), sudden sensorineural hearing loss, or immunological inner ear disease shows a reduction in LDL and the presence of acoustic distortion. Other possible causes of hyperacusis would be damage of the acoustic middle-ear reflex mediated by the facial nerve. Bell’s palsy, other facial palsies, neuro-muscle disorders such as myasthenia gravis, or stapes surgery may present DST in many patients. This kind of DST, however, usually abates spontaneously over time. Table 57.1 shows the most relevant etiologies.

Table 57.1 Cause of hyperacusis from ear disorders range

<table>
<thead>
<tr>
<th>Category</th>
<th>Cause of Hyperacusis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cochlear diseases</td>
<td>Ménière’s disease/endo lymphatic hydrops</td>
</tr>
<tr>
<td>Perylimphatic fistula</td>
<td></td>
</tr>
<tr>
<td>Sudden deafness</td>
<td></td>
</tr>
<tr>
<td>Acoustic trauma/noise induced hearing loss</td>
<td></td>
</tr>
<tr>
<td>Otoesclerosis</td>
<td></td>
</tr>
<tr>
<td>After surgical procedures</td>
<td>Post stapedectomy</td>
</tr>
<tr>
<td></td>
<td>After transtympanic tube placement</td>
</tr>
<tr>
<td></td>
<td>After wax removal</td>
</tr>
<tr>
<td>Stapedial reflex disorders</td>
<td>Sdr. Ramsay hunt</td>
</tr>
<tr>
<td>Muscular disorders</td>
<td>Bell’s facial palsy</td>
</tr>
</tbody>
</table>

Table 57.2 Causes of hyperacusis related to central nervous system disorders

<table>
<thead>
<tr>
<th>Category</th>
<th>Cause of Hyperacusis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serotonin (5HT)</td>
<td>Migraine</td>
</tr>
<tr>
<td></td>
<td>Depression</td>
</tr>
<tr>
<td></td>
<td>Sd. Posttraumatic stress</td>
</tr>
<tr>
<td></td>
<td>Cranioencephalic trauma</td>
</tr>
<tr>
<td></td>
<td>Lyme’s disease (Borrelia burgdorferi)</td>
</tr>
<tr>
<td>Williams Sdr.</td>
<td></td>
</tr>
<tr>
<td>BZD dependence Sdr.</td>
<td></td>
</tr>
<tr>
<td>Chronic postviral fatigue Sdr.</td>
<td></td>
</tr>
<tr>
<td>Serotonin disfunction</td>
<td></td>
</tr>
<tr>
<td>Tay-Sachs Sdr. (gangliosidosis 2)</td>
<td></td>
</tr>
<tr>
<td>Multiple sclerosis</td>
<td></td>
</tr>
<tr>
<td>Benign intracranial hypertension Sdr.</td>
<td></td>
</tr>
</tbody>
</table>

Other possible mechanisms of hyperacusis are serotonin (5HT) has been involved in some diseases such as migraine, depression, or posttraumatic stress syndrome – disorders associated with DST that may modulate auditory signals [14]. (5HT) has an important role for central auditory processing (CAP) and can be decreased in older people. A study performed in elderly patients showed that treatment with a selective serotonin release inhibitor (citalopram) improved the results of auditory processing and speech discrimination tests [15].

A second mechanism described in hyperacusis is based on the role of the endogenous endorphins [16]. Anxiety and stress increase the liberation of endorphins in the IHC–auditory nerve synapses. These substances potentiate the excitatory effect of the glutamate and therefore may increase excitation in the auditory periphery. Table 57.2 gives a list of central disorders associated with hyperacusis.

The inhibitory neurotransmitter GABA acts at several levels on the acoustic pathways. Even the function of the cochlea depends on GABA transmission at IHC synapses. A decrease in the action of GABA will increase neural activity and could be a correlate for hyperacusis. GABA, receptor agonists, such as benzodiazepines, could be used for some forms of hyperacusis. The author has used pregabalin for DST management with good results in some patients. Pregabalin affects many receptors and produces a dose-dependent increase in glutamic acid decarboxylase activity, increasing neuronal GABA levels.

Diagnosis of Hyperacusis

There is no objective measurement of hyperacusis because it is a subjective symptom (Table 57.3). A complete audiological examination, however, can be useful in the diagnosis of hyperacusis. Tonal and speech audiograms, tympanometry, and the study of the acoustic middle–ear reflex should be performed in all patients. ABR can rule out vestibular schwannoma and other retrocochlear diseases (multiple sclerosis) and is also useful for the diagnosis of auditory nerve neuropathy. An increase in the amplitude of the ipsilateral ABR responses and a decrease in the contralateral ABR in normal hearing subjects would rule out a LOCB disorder [13].

The study of OHC function and the MOCB efferent system can be performed through otoacoustic emissions (OEA). Study of the MOCB efferent system can be useful for diagnosis of some causes of DST. The discomfort threshold, which is the sound intensity that is annoying and not tolerable, can be determined. Its
Table 57.3 Classification of hyperacusis according to the loudness discomfort level and dynamic range

<table>
<thead>
<tr>
<th>Degree</th>
<th>Dynamic range</th>
<th>Loudness discomfort level</th>
</tr>
</thead>
<tbody>
<tr>
<td>No hyperac.</td>
<td>≥60 dB</td>
<td>≥95 dB in all the frequencies</td>
</tr>
<tr>
<td>Mild</td>
<td>50–55 dB in any frequency</td>
<td>80–90 dB in 2 or more frequencies</td>
</tr>
<tr>
<td>Moderate</td>
<td>40–45 dB in any frequency</td>
<td>65–75 dB in 2 or more frequencies</td>
</tr>
<tr>
<td>Severe</td>
<td>≤35 dB in any frequency</td>
<td>≤60 dB in 2 or more frequencies</td>
</tr>
</tbody>
</table>

Table 57.4 List of affected or avoided activities due to DST

| Concerts | Social life | Sport spectacles |
| Going to the restaurants | Going to church | House keeping |
| Going to the cinema | Working | Taking care of the children |
| Shopping | Driving | Others |

normal values are more than 90 dBHL, which are lower than the pain thresholds. It has to be tested several times because patients may have an initial fear of sounds, which would initially give lower thresholds than the real tolerance level.

Many individuals with DST will avoid different activities, affecting quality of life. The use of visual analogue scales for evaluation of hyperacusis handicap is useful (described in Table 57.4). Another system that has been proposed, named MASH, classifies the hyperacusis in four grades according to a broad list of activities: mild (≤3), moderate (from 3.1 to 5), severe (from 5.1 to 7), and very severe (≥7) [17].

In recent years, some specific questionnaires for DST have been developed and are useful tools in clinical diagnosis. The “self-rating Questionnaire on Hyper-sensitivity to Sound” published by Nelting and Rienhoff [18], evaluated DST according to three factors: cognitive reactions to hyperacusis, behavioral changes, and emotional responses to external sound. It was based on 15 questions; the scores went from 0 to 45. Every question had four possible answers: never (0 points), sometimes (1 point), often (2 points), and always (3 points). The score obtained can be divided into four grades, as we can see in Table 57.5. This questionnaire was originally written in German and it has been translated into Spanish [19]. Another published questionnaire was written by Khalifa [20]. It is based on 14 items and evaluates three dimensions: attention, social interaction, and emotion.

Table 57.5 Grades of hyperacusis considering the GUF

<table>
<thead>
<tr>
<th>Degree</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mild</td>
<td>≤10</td>
</tr>
<tr>
<td>Moderate</td>
<td>From 11 to 17</td>
</tr>
<tr>
<td>Severe</td>
<td>From 18 to 25</td>
</tr>
<tr>
<td>Very severe</td>
<td>26–45</td>
</tr>
</tbody>
</table>

In a study on 250 consecutive patients [5], we described the clinical characteristics of hyperacusis and tinnitus in DST patients. Direct questions and specific questionnaires were used to evaluate the interference of DST and tinnitus on quality of life. Auditory and psychoacoustic measurements were done on all participants. The answer to a question “do you feel more uncomfortable with environmental sounds than a majority of people?” was affirmative for 54% of the participants. Fifty-two participants had to stop one or more activities from a list of eleven (shopping, driving, taking care of children, going to church, etc.) because of DST. Sixty-three percent of the tinnitus clinic population showed LDL ≤90 dBHL, which was our definition of hyperacusis. Sixty-one percent were women, whose average age was 51 years (±14). Anxiety or stress was reported by 65% of the group, and 15% described the presence of different phobias: height, closed spaces, or insects. Sleeping problems were also very common (51%), and in two-thirds of the cases, tinnitus was the main problem for lack of sleep. A hearing impairment over 25 dBHL in any frequency was present in 83% of the participants.

The tinnitus of the DST group was predominantly in the left ear (52%), 27% in the right ear, and bilateral or cephalic in 21%. The average time the participants had experienced their symptoms was 6.6 years. The symptoms were present all day in 81% and had fluctuant intensity in 42% of the participants. The tinnitus increased by anxiety in 63% of the participants by loud external sound (27%) and postural changes (10%). The Tinnitus Handicap Inventory (THI) was used for evaluation of the severity and degree of annoyance. An average of 47 points was obtained. The visual analogue scale on tinnitus loudness scored 6.5 ± 2 (range 1–10).

Psychoacoustic measurement of tinnitus pitch showed that 46% had high-frequency tinnitus (>2 kHz) 34% from 0.5 to 2 kHz, and 14% of the participants matched their tinnitus to low frequencies. Average loudness was 9.8 dB ± 8.5 and minimum masking level was 19.3 dB ± 18.5. Two percent reported a temporarily complete elimination of the tinnitus with residual inhibition, whereas 56% obtained a partial reduction. Forty-two percent had no changes after sound exposure.

The Spanish version of the Sound Hypersensitivity Questionnaire (THS) was evaluated in another study with 40 participants with DST who were referred to our Tinnitus and Hyperacusis Clinic.
Seventy percent of the participants were female; average age was 48 ± 11 years. Hearing loss was present in 77% of the participants. THS average was 20.1 ± 10.0 points (range 1–45). The questions “I cannot listen or pay attention when intense or annoying sounds from my surroundings are present,” “I have to leave when there are intense surrounding sounds,” and “I am worried of hearing loss because of exposure to intense sounds” were answered with “yes” by most of the participants. There was a significant correlation ($p < 0.05$) between higher scores of the THS and a higher score in the visual analogue scale and the number of affected activities. The group of DST patients with hearing loss had higher scores in THS, but there was no correlation between the degree of hearing loss (pure-tone average) and the THS scores. Ninety percent of the participants presented tinnitus. The presence of tinnitus and its handicap, according to a visual analogue scale and the THI evaluation, were also correlated with higher THS scores. There was no significant relation between THS values and sex, age, possible etiology, duration of the disease, and loudness discomfort levels [19].

**Therapeutical Approaches**

There is one basic pillar for hyperacusis treatment: acoustic stimulation. Reaching this objective requires two steps. The first one is counseling. A professional can be able to change the patient’s negative feelings about causes of hyperacusis, possibilities for its control, treatment options, and prognosis. Counseling should be focused on positive and evidence-based medicinal information, reducing the patient’s emotional reaction and behaviors.

The second step is acoustic stimulation. Controlled and progressive exposure to sound has been demonstrated to be a useful tool in hyperacusis management, as we will see later in this chapter. Patients should avoid regular use of hearing plugs, except for the activities they are not able to perform without ear protection. The continuous use of the earplugs will increase the loudness discomfort levels and will decrease sound tolerance. The combination of counseling and white noise stimulation was developed by Jastreboff on the basis of a neurophysiologic model of tinnitus and named Tinnitus Retraining Therapy (TRT) (see Chap. 73). TRT has demonstrated its efficacy for hyperacusis management [2] and is now in routine use in many clinics. According to TRT, sound therapy can be delivered using three systems.

- **Environmental sound enrichment.** Different devices are useful for sound enrichment. A progressive increase in the volume of different kinds of sounds is used to increase sound tolerance in a slow but constant way. This method is effective for mild or moderate hyperacusis.

- **Broad band noise generators (BBNG).** According to Jastreboff’s criteria, broad band noise generators should be used when LDL were 80–85 dB or less. Jastreboff reported that 30 percent of the tinnitus patients required hyperacusis management before treatment of their tinnitus [21]. The BBNG is designed to produce two sounds with different spectrums: one covers low and middle frequencies and the other one covers some high frequencies. The digital noise generators can be customized to each patient’s preferences. The patient starts the therapy at the maximum volume tolerated without feeling annoyance. In some patients, the volume and time of exposure to the generator has to be increased on a weekly or monthly basis and extended to up to 8 h a day.

- **Hearing Aids.** Patients with hearing loss and moderate or severe hyperacusis will require DST management before being fitted with a hearing aid. This symptom could lead the patient to reject the device. There is also a possibility that the hyperacusis and tinnitus could increase. In a recent study, 41 percent of DST patients in our clinic experienced increased loudness of their tinnitus after exposure to loud sounds [5] (see also Chap. 74). The fitting process has to be slow, progressive, and made in accordance to the patient’s tolerance. We recommend that patients first use their hearing aid in quiet places. The use of the device and environmental sound exposure should be increased after this initial period of adjustment. The hearing aid compression systems and the maximum output of the device should be adjusted to avoid annoyance. The use of auditory training and broad band noise generators before the hearing aid fitting helps improve the LDL, dynamic range, and the speech comprehension. This method has been used by other authors, such as Knáster [22, 23], who obtained a reduction of the LDL (recruitment coefficient) in 59% of participants who had unilateral DST and 94% in bilateral DST.

The results of TRT in the management of DST are convincing. Gold reduced the LDL for 2, 3, and 4 kHz in more than 12 dBHL [24] at the end of treatment. Hazell reported that 45 percent of the patients he treated returned to regular LDL after 6 months, and 61% of the patients
had regular LDL in 2 years. The number of activities the patient had to give up because of his DST was reduced from 3.5 to 1.1 after 15 months of TRT [25].

Noreña and Chery-Croze [26] have hypothesized that hyperacusis is caused by enhancement of neural activity in the auditory pathways caused by deprivation of input to the auditory nervous system at the hearing impaired frequencies (see also Chap. 11). The introduction of external sound limited to the impaired frequencies (inverse to the one showed at the audiogram) could progressively reduce the amplification in the auditory nervous system and thereby decrease hyperacusis. The intensity of the sound stimulus should be customized according to the hearing loss as it appears in the audiogram. The differences between TRT’s recommended noise stimulation and that suggested in Noreña’s study is that in Norena’s study, it is limited to the impaired frequencies; there is not a progression of the stimulus intensity. The intensity of the sound used should be kept the same during all training. Our method is based on stimulation with sound (or CDs) of different frequency ranges (2–8 kHz, 4–12 kHz, etc.), but there is no customized intensity for each frequency. The sound intensity is increased gradually according to patient’s improvement.

Although TRT is the most used method to treat hyperacusis worldwide, treatment with drugs can be used alone or in combination with sound treatment. Those patients we suspect of having a cochlear hyperexcitability may improve with administration of salicylates because of their ototoxicity [27]. Typical recruitment from non-compensated cochlear diseases (Ménière’s disease, sudden deafness, fluctuant sensorineural hearing loss, etc.) can be managed through steroid therapy (systemically or transtympanic delivery). The use of diuretics, beta-histine, and sulphiride are common on these clinical entities and can give some relief during acute crisis.

One of the mechanisms described for central hyperacusis has been associated with a decrease in serotonin. Drugs that are selective serotonin reuptake inhibitors (paroxetine, fluoxetine, sertraline) can therefore be helpful to some patients with DST [14, 28]. Drug or cognitive management of anxiety and depression can successfully treat the emotional component of hyperacusis. DST mechanisms based on GABA disorders can be alleviated using GABA_A agonists such as benzodiazepines. Other drugs, such as pregabalin and gabapentin, facilitate the GABA transport over the blood–brain barrier, among other effects. The authors’ personal experiences have shown pregabalin to be a useful drug for acute and severe DST in patients with normal hearing.

Conclusions

Hyperacusis is a decreased tolerance to sounds and is estimated to affect 9% of the general population. Pathophysiologic mechanisms can be cochlear diseases or disorders in the central auditory pathways, with an abnormal activation of the limbic system that increases the psychological and emotional reaction to the symptom. The combination of professional counseling and acoustic stimulation using controlled sounds (TRT) has been proven to provide relief of decreased sound tolerance in many patients with DST.

References

Chapter 58
Clinical Description of a Different Form of Tinnitus: Intermitent Tinnitus

Miguel J.A. Láinez, Anna Piera, and Alejandro Ponz

Keypoints

1. Intermittent (paroxysmal) tinnitus is a form of non-pulsatile tinnitus.
2. An intermittent nature can be the only sign that intermittent tinnitus is different from other forms of tinnitus.
3. Intermittent tinnitus may be accompanied by irregular symptoms of other neurotologic disorders.
4. Both objective and subjective tinnitus may be intermittent.
5. A wide range of pathologies may cause intermittent tinnitus, but the cause of most forms is unknown.

Keywords
Paroxysmal (intermittent) tinnitus • Myoclonus • Temporomandibular joint changes • Cerebellopontine angle changes • Migraine • Auditory hallucinations • Audiogenic seizures

Abbreviations
ABR Auditory brainstem response
AGS Audiogenic seizures
CPA Cerebellopontine angle
CSF Cerebrospinal fluid
EEG Electroencephalography
GABA Gamma amino butyric acid
IC Inferior colliculus
TMJ Temporomandibular joint

Introduction

Suddenly occurring non-pulsatile tinnitus (intermittent tinnitus) can be the only symptom or it can be accompanied by neurotologic symptoms like vertigo, headaches, visual changes, and disturbances of consciousness.

Like constant tinnitus, intermittent tinnitus can be objective or subjective, depending on the pathology; objective tinnitus is caused by physical sounds generated in the body, which can also be heard by an observer. Subjective tinnitus is caused by abnormal neural activity, and only the patient can hear the tinnitus.

Objective Intermittent Tinnitus

Some important pathology should be ruled out in patients who present with intermittent objective tinnitus. The most important disorders that may occur together with objective tinnitus are palatal and middle-ear muscle myoclonus and temporomandibular joint (TMJ) disorders.

Palatal and Middle-Ear Muscle Myoclonus

Tinnitus produced by middle-ear myoclonus is objective intermittent tinnitus, and is rare; only a few cases are reported in the literature. In middle-ear myoclonus, otoscopic examination shows visible rhythmic movements of the eardrum, and weak clicking sounds are heard in the ear by auscultation. Tympanometry confirms rhythmic changes in middle-ear compliance. Middle-ear myoclonus can be accompanied by palatal myoclonus or can be the only manifestation. Palatal
myoclonus is an uncommon rhythmic “shock-like” involuntary movement of the muscles of the soft palate, throat, and other structures derived from the branchial arcs. Objective intermittent tinnitus associated with palatal myoclonus can be related to hearing impairment; however, this relation is not always present. Examination of muscles of the soft palate and throat shows rhythmic involuntary movements and, in some cases, spontaneous clicking sounds by auscultation near the ear [1–3].

**Temporomandibular Joint Changes (Synchrony with Joint Movements)**

The TMJ is a complex, sensitive, and highly mobile joint. Millions of people suffer from temporomandibular disorders. Tinnitus associated to TMJ changes is synchronous with joint movement and is easy to provoke during the examinations. Several disorders of TMJ cause tinnitus: luxation, condyle malposition, bruxism, degenerative arthropathy, capsulitis, and many others. These disorders are also very common in inflammatory arthropathies. Regardless of the cause of a TMJ disorder, all of them can be common causes of intermittent tinnitus, and a routine examination for these disorders is needed in all tinnitus clinics [4–6].

**Subjective Intermittent Tinnitus**

Subjective intermittent tinnitus, which is much more common than objective tinnitus, can occur together with pathologies such as cerebellopontine angle (CPA) disorders.

**Cerebellopontine Angle Disorders**

CPA disorders may be suspected in patients with unilateral hearing loss and unilateral intermittent tinnitus with or without dizziness. Audiological and imaging studies of the posterior fossa are used to rule out disorders of the CPA. Lesions of the CPA are frequent and represent 6–10% of all intracranial tumors. Vestibular schwannoma (acoustic neuroma) and meningioma are the two most frequent lesions and account for approximately 85–90% of all CPA tumors. The other 10–15% encompasses a large variety of lesions including aneurysms, epidermoid cysts, arachnoidal cysts, Arnold–Chiari malformations, lipoma, and melanomas. Such lesions are now detected more frequently because of the sensitivity and accuracy of magnetic resonance imaging (MRI) [7–10].

Recently, Levine has described a subtype of intermittent tinnitus, called typewriter tinnitus, with an excellent response to treatment with carbamazepine, in which vascular compression of the auditory nerve was suspected to be the cause in five of six patients that were studied. This suggests that surgical decompression may also be effective in such patients [11].

**Audiogenic Seizures and Epilepsy**

The inferior colliculus (IC) plays an important role in many pathophysiological conditions that involve hearing (including tinnitus, age-related hearing loss, and audiogenic seizures (AGS)). AGS occur frequently in rodents and can be genetically mediated. AGS can also be readily induced in experimental animals [12]. AGS can be induced in normal animals by administration of drugs that are GABA receptor antagonists. Glutamate-mediated excitation is a critical element of neurotransmission in IC neurons, and excessive activation of glutamate receptors in the IC is implicated in AGS. Such neurotransmitter abnormalities cause excessive firing of IC neurons that act as the critical initiation mechanism for triggering seizures in response to intense acoustic stimuli, thus AGS. The IC plays a role in the integration of acoustic-motor and acoustic-limbic integration, as well as in acute and chronic AGS. García-Caraisco et al. [13] have demonstrated in animal experiments that chronic kindled AGS change behavioral expressions in a similar way as those that occur in temporal lobe epileptic seizures mixed with audiogenic seizure activity, which is known to be dependent on brainstem networks. This form of AGS involves subcortical intermittent pattern of tinnitus manifestation [14, 15].

Tinnitus, as an intermittent pathologic cortical manifestation, has been described in only a few patients while electroencephalography (EEG) was monitored, and was determined that the tinnitus originated from the contralateral mid-temporal area [16].
Auditory Hallucinations

Auditory hallucinations can be simple or complex. Therefore, tinnitus could be considered an auditory hallucination. Several studies with transcranial magnetic stimulation have reported a benefit in improving both auditory hallucinations and tinnitus by modulating cerebral cortex activity. New results of studies and new questions about the neurobiological basis of mental and neural disorders have concerned whether there is a common substrate in tinnitus and auditory hallucinations [17, 18].

Migraine with Basilar Aura

Vertigo, dysarthria, and tinnitus may occur together with basilar aura in individuals with migraine. It has been reported to occur in 50% of individuals with basilar aura. Other symptoms are diplopia, bilateral visual symptoms, bilateral paresthesia, hearing loss, decreased level of consciousness, and ataxia. For management of patients with tinnitus and headache, it is important to obtain information about how the tinnitus may be regarded as a symptom that precedes headache [19–21].

Cerebrospinal Fluid Pressure Changes

Both intracranial hypotension and hypertension (pseudotumor cerebri) have been suggested as possible causes of intermittent tinnitus. A lumbar puncture is needed to measure pressure changes in cerebrospinal fluid (CSF), and in most cases, a neuroimaging technique (MRI or CT) is necessary to rule out other brain lesions [22–24].

Phantom Sensations Without Evidence of Cortical or Auditory System Dysfunctions

In spite of the many different pathologies that may cause intermittent tinnitus, in some patients no pathology can be found and the tinnitus remains a phantom intermittent sensation – an expression of an abnormally high correlation of activity in many nerve cells in cortical and subcortical parts of the auditory system [25, 26].

Techniques used in Diagnosis of Intermittent Tinnitus

Several different techniques are useful in the diagnostic workup of patients with intermittent tinnitus; the most important are MRI, auditory brainstem responses (ABR), and in some patients, EEG.

Magnetic Resonance Imaging

MRI is performed in almost all patients with intermittent tinnitus in order to rule out CPA disorders, cortical ectopias, and indirect signs of benign intracranial hypertension of licuoral hypotension.

Electroencephalography

EEG is only indicated if there are further signs of seizure and when tinnitus is accompanied by symptoms of consciousness disturbance.

Basal EEG recordings performed with provocation maneuvers like flashing light and hyperventilation in some cases may be useful. EEG can help determine if temporal lobe discharges are present.

Role of Auditory Brainstem Responses

ABR are indicated only in intermittent tinnitus for screening auditory nerve compression and can provide prognosis for microvascular compression [27].

References

Keypoints

1. Pulsatile tinnitus, in general, is not related to pathology of the auditory system.
2. Two main types of pulsatile tinnitus exist: arterial heart beat synchronous pulsatile tinnitus and venous respiratory synchronous pulsatile tinnitus.
3. The main causes of pulsatile tinnitus are related to aberrant/ectopic, stenosis, or other pathologies of blood vessels, either arterial or venous.
   (a) MRI, CT, or classical angiography are diagnostic tools for pulsatile tinnitus.
4. Benign intracranial hypertension is another common cause of pulsatile tinnitus.
   (a) Funduscopy and lumbar puncture with measurement of cerebrospinal pressure are diagnostic tests for benign intracranial hypertension.
5. In 15–30% of patients, no cause can be found for the pulsations.
6. Pseudopulsatile tinnitus groups a number of muscle-related tinnitus types mimicking pulsatile tinnitus.

Keywords Pulsatile • Tinnitus • Venous • Hum • Arterial • Intracranial hypertension

Abbreviations

ABR Auditory brainstem response
AV Anterior-venous
AVM Arterial venous malformation
BIH Benign intracranial hypertension
CT Computerized (axial) tomography
EEG Electroencephalography
ICA Internal carotid artery
IPL Interpeak latencies
MRA Magnetic resonance angiography
MRI Magnetic resonance imaging
MVC Microvascular compression
MVD Microvascular decompression
TMJ Temporomandibular joint

Introduction

Tinnitus can be subdivided into two entirely different entities: pulsatile and non-pulsatile tinnitus [1–3]. Most forms of pulsatile are heart beat synchronous, where arterial pulsations modulate the tinnitus. The arterial pulsations are most likely transmitted to the cochlea via the cerebrospinal fluid, a mechanism similar to what has been proposed as an explanation for bone conduction [4–6]. Respiration synchronous tinnitus is rare, but an individual may perceive hearing their own breathing sounds because of an open Eustachian tube as tinnitus.

Pulsatile tinnitus synchronous with the heart beat seems to be related to arterial causes; pulsatile tinnitus synchronous with respiration is most likely due to venous causes. Venous pulsatile tinnitus might be equally or even more prevalent than arterial heart beat synchronous tinnitus [2, 7], even though it is not as well known as arterial pulsatile tinnitus. Thus, pulsatile tinnitus, in general, is not associated with pathology of the auditory pathways per se, which is in contrast to the more common non-pulsatile tinnitus.
Arterial pulsations may modulate existing tinnitus or cause tinnitus. The tinnitus caused by arterial pulsations being transmitted to the cochlea are forms of objective tinnitus.

Since heart beat synchronous tinnitus is predominantly vascular in origin, almost all causes of pulsatile tinnitus can be diagnosed by magnetic resonance imaging and magnetic resonance angiography, except for benign intracranial hypertension [1–3].

Causes of Pulsatile Tinnitus

Heart Beat Synchronous Pulsatile Tinnitus

Arteriovenous malformations (AVM) of the dura are the best known causes of arterial pulse synchronous pulsatile tinnitus [8, 9]. Such abnormal communications between the arterial and venous systems may be congenital or acquired [10]. Often AVMs result from chronic mastoiditis or other causes occluding the sigmoid-transverse sinus, such as trauma. As a natural repair mechanism, vascular bypasses tend to develop around the occlusion, resulting in a dural AVM. If the dural AVM is symptomatic or if it is asymptomatic with leptomeningeal drainage, these lesions are often treated with embolization, usually in multiple sessions. If intractable with endovascular, treatment involving surgical excision of the AVM and dura is often done [11] (Table 59.1).

Posttraumatic pulsatile tinnitus can be the result of a carotid dissection, AV fistula, or caroticocavernous fistula. In 16–27% of carotid dissections, pulsatile tinnitus is experienced at the side of the dissection but is usually associated with other focal or global symptoms [12]. In contrast, in (non-traumatic) vertebral artery dissection, only 5% of patients present with pulsatile tinnitus [13].

Carotid dissection in the neck is a relatively common condition. Most dissections are spontaneous, likely related to activities that cause a sudden stretch of the pharyngeal portion of the carotid artery. Traumatic carotid dissections occur in approximately 1% of all patients with blunt injury mechanisms [14]. Carotid dissections are characterized by a triad of neck and head pain, Horner’s syndrome, and pulsatile tinnitus. Others present with transient or persistent brain ischemia. Strokes are due to the embolization of thrombus material from the lumen of the dissected artery to the intracranial arteries, most often the middle cerebral artery [15]. Carotid dissection is asymptomatic in less than 10% of the patients, whereas in more than 90% of patients, carotid territory ischemia and/or local signs and symptoms on the side of dissection develop. Local signs and symptoms on the side of dissection include head (65–68%), facial (34–53%), or neck pain (9–26%), as well as Horner syndrome (28–41%) and cranial nerve palsy (8–16%), in particular the hypoglossal nerve. The facial nerve may also be involved; dysgeusia results mainly from involvement of the chorda tympani (0.5–7.0%) or the glossopharyngeal nerve. Transient pareses of the ocular motor (III, IV, and VI) and trigeminal nerves have been observed. In three-fourths of carotid dissections, an ischemic event occurs, which includes ischemic stroke in 80–84%, transient ischemic attack in 15–16%, amaurosis fugax in 3%, ischemic optic neuropathy in 4%, and retinal infarct in 1% [12].

Treatment consists of anticoagulants or antiplatelet agents, and healing occurs within 3–6 months with a resolution of stenosis in 90% of patients, and recanalization of occlusions in as many as 50% [14]. In cerebral hemodynamicly, compromised patients without an irreversible infarct emergency stenting can be considered [14].

Table 59.1 Causes of pulsatile tinnitus

<table>
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<th>Pulsatile tinnitus</th>
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<td>Venous</td>
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<td>High jugular bulb</td>
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<td>Sigmoid sinus diverticulum</td>
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<td>Carotid stenosis</td>
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<td>Vascular lesions of petrous bone/skull base</td>
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<td>Arteriovenous malformation/fistula (dural, caroticocavernous, etc.)</td>
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<td>Intrapetrous aneurysm</td>
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<td>Hyperdynamic state (anemia, thyreotoxicosis, pregnancy, etc.)</td>
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<td>Paget’s disease</td>
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<td>Carotid or vertebral artery dissection</td>
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<td>Benign intracranial hypertension</td>
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Posttraumatic AV fistulas often result in pulsatile tinnitus. The sinus fistulas are often the result of a venous thrombosis, similarly to the non-traumatic variant of AV fistulas. The most common posttraumatic fistula is the carotid-cavernous fistula. These are characterized by pulsatile tinnitus, pulsating exophthalmia, chemosis, and visual deficit of the afflicted side.

Carotid-cavernous fistulas are the most common arteriovenous fistula. They are divided in the more common (70%) [16] direct high- and rare indirect low-flow fistulas. Low-flow fistulas are usually associated with atherosclerosis, hypertension, and collagen vascular disease, or may develop in females during the peripartum period [16]. They can be spontaneous (low-flow) [17] or high-flow posttraumatic [18]. Spontaneous high-flow fistulas are very rare [19]. Carotid-cavernous fistulas are characterized by pulsatile tinnitus (50% in low flow [20] and more frequently in high flow), pulsating exophthalmia, chemosis, and visual deficit of the fistula side [21]. Treatment consists predominantly of endovascular treatment [22], although high obliteration rates have been described 1-3 months after gamma knife surgery [23].

Dural arteriovenous malformations are the best known causes of arterial pulse synchronous pulsatile tinnitus. Often AVMs result from chronic mastoiditis or other causes occluding the sigmoid-transverse sinus, such as posttraumatic thrombosis. As a natural repair mechanism, vascular bypasses tend to develop around the occlusion, resulting into a dural AVM [24]. If the dural AVM is symptomatic, or if it is asymptomatic with leptomeningeal drainage, these lesions should be embolized, usually in multiple sessions [25, 26]. In benign lesions with only tinnitus or embolization failures, gamma knife surgery is an alternative option if no cortical venous drainage is present [27]. If intractable with endovascular or radiosurgical treatment, surgical excision of the AVM and dura can be proposed [11, 28, 29]. Not only transverse or sigmoid sinus AV fistulas can generate pulsatile tinnitus: sagittal sinus AV fistulas [30] and carotid-cavernous fistulas can generate pulsatile tinnitus as well, which disappears after embolization or gamma knife surgery.

For posttraumatic AV fistulas, the reader is referred to Chap. 66 [21].

It is not only dural fistulas or carotid-cavernous AVMs or fistulas that can present with arterial pulsations but also AVMs of the external ear [31], hypoglossal canal [32], and even the parotid gland [33] can cause pulsatile tinnitus.

Carotid stenosis is a common cause [1–3] of arterial pulsatile tinnitus. The most common cause is atherosclerotic disease [34, 35], but fibromuscular dysplasia [8, 36] can also cause pulsatile tinnitus. This kind of pulsatile tinnitus typically disappears when compressing the ipsilateral, internal, or common carotid artery. The diagnosis can be confirmed by sonography, MRI, CT, or classical angiography. Treatment of the extracranial carotid artery stenosis can consist of dilatation and stenting or carotid endarterectomy. Ipsilateral carotid endarterectomy for tinnitus is effective in reducing or abolishing tinnitus in more than 90% of patients with demonstrated ICA stenosis related to pulsatile tinnitus. Proximal lesions lend themselves to carotid endarterectomy, whereas distal lesions have been treated by stenting [37]. For the rarer intracranial carotid artery stenosis, two approaches have been used; an initial balloon occlusion test under transcranial doppler and EEG monitoring can verify whether the ipsilateral carotid artery can be sacrificed. If so, one option is to ligate the symptomatic carotid artery. The other option is to dilate and stent the intracranial portion of carotid artery, resulting in a disappearance of the arterial pulsatile tinnitus. A major problem still faced today is that stents might occlude. Thus, this elegant technique still remains experimental until the coagulation problems can be better controlled [38]. Overall, almost 70% of patients with carotid stenosis are cured by intervention, and most of these patients experience (close to 90%) immediate relief of tinnitus [37].

It is not only a stenosed internal or external carotid artery that can lead to pulsatile tinnitus; a stenotic subclavian [39] or external carotid artery [40] can also generate a treatable form of arterial pulsatile tinnitus. Reversal of blood flow in an aberrant occipital artery can also cause pulsatile tinnitus, and this condition can also be treated by stenting [41].

Hyperdynamic flow in the internal carotid artery can generate pulsatile tinnitus, as seen in basilar artery atresia with predominant flow in both carotids [42]. The hyperdynamic flow can also be due to anemia [43], thyrotoxicosis, or pregnancy.

It has been suggested that a mechanism by which one does not normally hear the pulsations of the carotids is due to a dampening effect of a pericarotid venous plexus [42]. Extensive pneumatization around the carotid artery could, however, reduce this dampening effect and result in the perception of arterial pulsations [44].
Aneurysms of the petrous carotid artery can lead to pulsatile tinnitus [8, 9, 45–47]. This may be caused by the aneurysm obliterating this venous plexus, allowing the arterial pulsations to be transmitted directly to the cochlea and resulting in the perception of the arterial pulsations. However, this is not the only mechanism involved, as aneurysms of the anterior communicating artery have also been related to pulsatile tinnitus [48].

**Congenital Vascular Anomalies**

The persistent stapedial artery, a normal fetal artery that ordinarily disappears before birth, can cause pulsatile tinnitus. The persistent stapedial artery runs through the obturator foramen between the crura of the stapes and across the promontory in the middle ear, leaving the middle ear to run along the tympanic portion of the facial nerve canal near the geniculate fossa, finally exiting the facial nerve canal to supply the territory of the middle meningeal artery, which never develops in the case of a persistent stapedial artery. Consequently, the foramen spinosum, the entry of the middle meningeal artery in the skull does not develop either [49].

Aberrant and ectopic internal carotid arteries have also been implicated in arterial pulsatile tinnitus [7]. An aberrant carotid artery is a congenital anomaly in which the cervical internal carotid artery never develops. Instead, the inferior tympanic artery (a branch of the ascending pharyngeal artery) enlarges, anastomoses with the caroticotympanic artery in the middle ear, and resumes the usual course of the internal carotid artery in the horizontal portion of the petrous carotid canal. The aberrant carotid artery may be dehiscent and visible through the tympanic membrane as it courses through the middle ear [49].

Glomus tumors, or paraganglioma, are associated with unilateral hearing loss in 80% of cases and with pulsatile tinnitus in 60% [50]. These tumors occur predominantly in women (6:1), and should thus be differentiated from benign intracranial hypertension (BIH). Diagnosis is confirmed by MRI and/or angiography. Some glomus tumors (1–3%) are endocrino-logically active and secrete catecholamines [50]. As glomus tumors are mostly benign lesions (less than 3% metastasize) growing less than 2 cm in 5 years, treatment options are either a “wait and scan” policy or embolization and surgery [51]. If the tinnitus is incapacitating, the embolization with surgery option can be helpful. The glomus tumor cell is radiation insensitive, but irradiation can reduce the vascularity responsible for the pulsatile tinnitus.

Other vascular lesions of the petrous bone or skull base – such as hemangiopericytoma [52], plasmacytoma [52], giant cell tumors [53], and neuroendocrine carcinoma [54], amongst others – are also known to cause tinnitus that can be treated by otoneurosurgical methods.

Pulsatile tinnitus that occurs in individuals with Paget’s disease can be explained in a similar way, especially when the temporal bone is involved [10, 34]. It has indeed been suggested that increased vascularization with intraosseous arteriovenous shunts may be responsible for the pulsatile tinnitus [10].

Microvascular compressions of the cochlear nerve can cause incapacitating pulsatile or non-pulsatile tinnitus [55, 56]. A meta-analysis has shown that individuals with pulsatile tinnitus are 80 times more likely to have a vascular loop in close contact with the root of the auditory nerve than individuals with non-pulsatile tinnitus [57]. Most vascular compressions, however, cause non-pulsatile tinnitus. This is similar to other disorders of microvascular compressions (MVC) of cranial nerve roots, such as trigeminal neuralgia and hemifacial spasm, which also do not have pulse synchronous bouts of pain in the distribution of the trigeminal nerve or pulse synchronous hemifacial spasms. The diagnosis of microvascular compression in individuals with tinnitus is based on the clinical picture and confirmed by auditory brainstem-evoked potentials and magnetic resonance imaging [58] [59–61] (see Chap. 40).

However, if the vascular loop extends into the internal auditory meatus, it can cause arterial pulse synchronous tinnitus via CSF/bone conduction [42, 62]. Studies of MRIs of individuals with pulsatile tinnitus, after exclusion of other causes, have shown a statistically significant high number of vascular loops in the internal auditory canal in comparison with individuals with non-pulsatile tinnitus [62]. Placement of shredded Teflon between the vascular loop and the nerve (microvascular decompression, MVD) has abolished pulse synchronous tinnitus [42]. Pathophysiologically, the sharp turn of the vascular loop in the triangular internal auditory canal creates a turbulence, which creates sound waves that are concentrically irradiating. The internal auditory canal has a cave or funnel effect guiding the sound waves towards its end – the top of the triangle where the cochlea is located. The sounds waves in the CSF–bone interface are transferred to the cochlea via bone conduction. High-frequency waves carry less
energy than low-frequency waves and are therefore reflected more easily than the longer low-frequency waves. This could explain why pulsatile tinnitus is matched to low frequencies.

A new form of pulsatile tinnitus has been described and called the somatosensory pulsatile tinnitus syndrome [63]. It is characterized by a high-pitched, pulse synchronous tinnitus, where the pulsations can be suppressed by strong contractions or normal compressions of the neck and jaw muscles (somatic testing) [63]. This form of tinnitus is hypothesized to be related to heart synchronous somatosensory activation of the central auditory pathway or to failure of the somatosensory–auditory central nervous system interactions, which normally suppresses heart somatosounds [63].

A semicircular canal dehiscence can be the cause of pulsatile tinnitus (7%) [64, 65], especially when gaze evoked. Twenty-five percent of patients with dehiscence have this kind of tinnitus.

Dehiscence of a semicircular canal is described most often for the superior canal [66], but posterior [67–70, 71, 72] and lateral semicircular canal dehiscences have also been described[73]. Hyperacusis to bone-conducted sounds is also commonly found in 39% of patients. The most typical signs of canal dehiscence are, however, autophony and “blocked ear” (94%) [74], sound-induced vertigo (97%) (Tulio’s sign), and oscillopsia. The autophony does not present with audible breathing such as that in a patulous Eustachian tube [7]. Most patients also complain of chronic dysequilibrium. In addition, most patients will have sound or Valsalva-evoked eye movements in the plane of the dehiscent canal. Tragal pressure evokes the nystagmus in 54% (Hennebert’s sign) and one-fifth of patients show sound-evoked head movements.

Patients may describe very unusual findings – hearing their joints moving, hearing their eye movements, hearing their heart beat, hearing their heels strike during walking, or the ability to hear a tuning fork placed at a distal extremity – due to increased bone conduction. Treatment of symptomatic patients consists of surgical plugging (the better method) or resurfacing of the canal [75].

Venous Hum

A venous hum was originally attributed to an impingement of the transverse process of the second vertebra in the jugular vein [76]. However, venous pulsatile tinnitus can have many different causes. BIH, also known as pseudotumor cerebri or idiopathic intracranial hypertension [77], is another possible cause of pulsatile tinnitus. Sismanis [2] showed that 40% of individuals he studied with pulsatile tinnitus were diagnosed with BIH. Other studies find that only 2% of the individuals with BIH had pulsatile tinnitus [7]. BIH almost exclusively afflicts young overweight women [77, 78]. Clinical symptoms include arterial pulsatile tinnitus, venous hum, headache, and blurry vision. Patients with BIH can also complain of aural fullness, low-frequency hearing loss, and vertigo [1, 2, 77]. Oddly enough, the venous hum presents most often unilaterally (80%) and can be the only symptom of BIH [2]. High intracranial pressure can cause more prominent symptoms that may occur after lying down (such as in the morning when waking up) or bending over or when coughing or performing other maneuvers that raise intracranial pressure. A suspicion of BIH can be confirmed by compressing the ipsilateral jugular vein, stopping the flow in the ipsilateral sigmoid sinus, which causes the venous hum to disappear. When the tinnitus disappears, the pressure-like headaches tend to increase because the intracranial pressure increases due to the reduced drainage of CSF.

About half of all individuals with BIH have low-frequency sensorineural hearing loss; this disappears on ipsilateral jugular vein compression as well. Papilledema is often present [77, 78], and magnetic resonance angiography (MRA) and MRI are usually negative. An empty sella, present in 25% of such individuals [2], should, however, raise suspicion of BIH as it can be related to prolonged intracranial pressure. Spontaneous cerebrospinal fluid leak should also be considered a sign of possible intracranial hypertension [79]. Diagnosis is usually confirmed by lumbar puncture (opening pressure >20 cm water). Treatment consists of weight loss, diuretics, or ventriculoperitoneal or lumboperitoneal shunting. BIH is usually idiopathic, but venous sinus outflow obstruction can be the cause [78] and also occurs after posterior fossa surgery (unpublished results).

The Arnold–Chiari malformation is a clinical entity in which there is a tonsillar herniation in the foramen magnum. Four different types exist, but only Chiari type I occurs frequently in the Western world. Seven to ten percent [80] of individuals with Arnold–Chiari malformations complain of tinnitus, which can be both non-pulsatile and pulsatile [81]. Pulsatile tinnitus most commonly consists of a venous hum, likely caused by raised intracranial pressure since it worsens on bending over and onValsalva maneuvers. The pulsations normally
disappear on ipsilateral jugular vein compression, which also causes improvement of the low sensorineural hearing loss, as the tinnitus may be masking normal hearing. No ABR changes have been noted in individuals with this kind of tinnitus. After surgical treatment of individuals with Arnold–Chiari malformations (decompression), this form of tinnitus often disappears [81]. The non-pulsatile tinnitus that occurs in such individuals is usually intermittent, and the cause is unknown; it may be caused by stretching of the cochlear nerve, such as by microvascular compression or brainstem traction [82]. ABR changes have been noted in 75% of patients and consist of prolongation of the IPL III–V in 100% of the patients and prolongation of IPL I–III in 30% [82]. Prolongation of IPL III–V may be due to brainstem traction and/or contralateral microvascular compression of the auditory nerve [83]; the IPL I–III may be caused by ipsilateral microvascular compression of the auditory nerve [84, 85]. Posterior fossa decompression, which consists of opening the foramen magnum and widening the dura mater, can therefore result in improving the non-pulsatile tinnitus in three out of four patients with Chiari malformation (De Ridder, unpublished results), similarly to what is seen in trigeminal neuralgia in patients with Arnold–Chiari malformation [86]. The improvement of such patients may be due to a secondary auto decompression of the vestibulocochlear nerve, analogous to what is suggested in surgical removal of posterior fossa tumors [87].

**Sigmoid Sinus Diverticulum**

A common cause for venous pulsatile tinnitus is a sigmoid sinus diverticulum [7], in which a diverticulum enters into the mastoid bone. The perceived pulsations most likely result from turbulent flow in the diverticulum, which is transmitted to the cochlea via bone conduction. Transmastoid reconstruction of the sigmoid sinus, as described for venous aneurysms, can result in a permanent cure for the pulsations [88].

**High Jugular Bulb**

A high jugular bulb can also cause venous hum, as a result of its close and direct contact with the cochlea. When the high bulb is dehiscent, it can be seen as a bluish mass in the hypotympanum on otoscopy [89], in contrast to the reddish mass in the anterior middle ear, which suggests a dehiscent/aberrant carotid artery [89]. In the same way as described for benign intracranial hypertension, the venous hum disappears on compression of the ipsilateral jugular vein. A high jugular bulb can be diagnosed by CT imaging. Surgically, ligating or lowering the jugular bulb and interposing Teflon or bone-wax can abolish or diminish this form of tinnitus [90, 91]. Transvenous stent-assisted coil embolization has been used as well for treating this condition [92]. Abnormal veins, such as an abnormal posterior condylar emissary vein [93] and abnormal mastoid emissary veins [94], have been described as surgically treatable causes of venous pulsatile tinnitus as well.

Like arterial aneurysms, a sigmoid-transverse venous aneurysm can be causally related to venous pulsatile tinnitus [95–97]. After coagulation of the aneurysm and reconstruction of the sinus wall [96], or after endovascular treatment [97], the pulsations can disappear.

**Pseudopulsatile Tinnitus**

There exist pathologies that mimic pulsatile tinnitus. Causes for this non-vascular tinnitus are palatal myoclonus [98], tensor tympani spasms [99, 100], stapedial muscle myoclonus [101], and a patulous Eustachian tube [102]. These pathologies generate neither arterial pulse synchronous nor respiratory rate synchronous tinnitus, but tend to fluctuate in intensity (as in a stormy wind) or are perceived as clicks. The clicking palatal myoclonus is often bilateral and is caused by contractions of the peritubal muscles, especially the levator veli palatine muscles which snap the Eustachian tube open, breaking surface tension. It can be effectively treated by botulinum toxin [103] or radiofrequency lesioning [104] of the peritubal muscles. Tensor tympani spasms and stapedial muscle contractions together are called middle-ear myoclonus. It generates rhythmic contractions (40–200 Hz) of the tympanic membrane coinciding with the tinnitus. It can be perceived as a “rushing wind” noise by the patient and can be treated by sectioning of the tendons of both these muscles resulting in an immediate improvement [100, 101, 105]. In some patients, stapedial muscle contractions can be treated by selective sectioning of stapedial tendons [101]. The stapedius contractions
create a buzzing sound, whereas the tensor typani has a clicking sound. The objective tubal tinnitus in a patulous Eustachian tube is often associated with autophonia and audible breathing. The somatosounds arise as a result of the walls of the Eustachian tube snapping together [102], with respiration associated with tympanic contractions. Botulinum toxin [106] and/or transsection of the tensor veli palatini muscle tendon may be a useful method of treatment if the patient experiences objective tinnitus, which is very distressing [102].

Conclusion

Pulsatile tinnitus can be divided into arterial heart beat synchronous pulsatile tinnitus and respiratory synchronous venous hum. It is important to look for a cause of the pulsations by neuroimaging tools, as many causes can be treated successfully. Idiopathic intracranial hypertension should be excluded as a non-vascular possible cause for the pulsations.

Pseudopulsatile tinnitus mimics pulsatile tinnitus but is not synchronous with the heart beat and therefore not of vascular origin. Instead, it is mostly of muscular origin.

References


68. Krombach GA, EDi Martin, S Martiny et al (2006) Dehiscence of the superior and/or posterior semicircular canal: delineation on T2-weighted axial three-dimensional turbo spin-echo images,


Chapter 60
Ménière’s Disease and Tinnitus

Michel Paolino and Vénéra Ghulyan-Bedikian

Keypoints

1. Ménière’s disease is a clinical syndrome that comprises vertigo, sensorineural hearing loss, subjective tinnitus, and aural fullness.
2. The tinnitus is classically low pitched and evolves with the progression of the disease.
3. The diagnosis of Ménière’s disease is based on patient history, a clinical examination, a complete oto-neurological assessment, and a MRI.
4. The differential diagnosis distinguishes Ménière’s disease from vestibular schwannoma, microvascular conflict of the VIII cranial nerve, and migraines.
5. Antivertiginous, antihistamines, loop diuretics, antiemetics, and benzodiazepines are effective in managing the acute attacks.
6. The following therapies are recommended for inter-cises periods:
   (a) Dietetic recommendations including low-salt diet
   (b) Medication (antihistaminics, diuretics, osmo-regulators, vasodilators, antiemetics, corticoids, benzodiazepines)
   (c) Relaxation therapy, tinnitus retraining therapy, sound therapy, etc.
   (d) Coordinated treatment of temporomandibulaires and cervical disorders
   (e) Intratympanic therapy with gentamycin or steroids
   (f) Surgical approach by endolymphatic mastoid shunt or endolymphatic sac decompression, vestibular neurotomia, and labyrinthectomy.
7. Some medications as well as transtympanic therapy seem particularly interesting, because they provide improvements of both vertigo and tinnitus while preserving the hearing in the majority of patients who have this treatment.
8. Conservative treatments should be exhausted and surgery reserved for patients with disabling and refractory vertigo, but surgery cannot prevent the progression of the tinnitus.

Keywords Ménière’s disease • Tinnitus • Vertigo • Hearing loss • Endolymphatic hydrops • Psychosomatic incidence

Abbreviations

AAO-HNS American Academy of Otolaryngology – Head and Neck Surgery
ABR Auditory brainstem response
BPPV Benign paroxysmal positional vertigo
CBT Cognitive-behavioral therapy
DPOAEs Distortion product otoacoustic emissions
ECoG Electrocochleography
MRI Magnetic resonance imaging
SP Summating potential

Ménière’s Disease: Pathogenesis, Symptoms, and Clinical Manifestations

Ménière’s disease represents one of the many causes of tinnitus. Since its first description in 1861, several etiological theories have been proposed to explain the pathogenesis of this disease: endolymphatic hydrops, autoimmune disorders, viral infections, allergic processes,
and activation of neural plasticity. Ménière’s disease is a syndrome that comprises of three (or four) symptoms:

- Recurrent episodes of spontaneous vertigo lasting from several minutes to a few hours, which can be followed by residual unsteadiness.
- Fluctuating and slowly progressive sensorineural hearing loss, usually unilateral and initially prevailing at low frequencies.
- Subjective tinnitus and sensation of aural fullness, pressure, or discomfort. The tinnitus is referred to the affected ear and described as a low-frequency “buzz” or “roar,” but as the disease progresses, it sometimes includes a high-pitched component.

Ménière’s disease usually starts in only one ear but can evolve into a bilateral form. It is characterized by periods of exacerbation and remission. The crises are frequently severe, incapacitating, unpredictable, and usually accompanied by anxiety, headaches, and autonomic manifestations (nausea, vomiting, diaphoresis, pallor, tachycardia, diarrhea, etc.).

At the beginning, the tinnitus complaint is secondary. Classically, it becomes worse during vertigo attacks but may significantly improve or even disappear afterward. However, with the progression of the disease, the tinnitus can become permanent, persisting between attacks. Its evolution is then unfavorable due to a significant increase in anxiety.

Besides the above-described classic form, some physicians have also included patients with incomplete clinical features as forms of Ménière’s disease. In fact, the frequency and the intensity of the crises, the association of symptoms, and their impact on the patient’s quality of life can vary from one patient to another. These various clinical forms can be classified into three groups:

- Predominantly cochlear forms where hearing loss and tinnitus are “in the foreground” and the vertigo is either absent or atypical
- Predominantly vestibular forms with typical vertigo crises, which are not necessarily preceded by tinnitus or aural fullness, and the hearing loss does not always affect the low frequencies
- Separate forms with initially typical vertigo crises without cochlear signs.

These symptoms that have similarities with the vertiginous crises of migraine should be distinguished from those of Ménière’s disease. Later, the cochlear symptoms usually occur together with a pattern of vestibular signs. Signs of endolymphatic hydrops sometimes are later added by vestibular symptoms, and the disease evolves into a typical Ménière’s disease.

**Diagnostic Criteria**

The diagnosis is established with patient history, a clinical examination, a complete oto-neurological assessment, and possibly a MRI of the brain.

**Patient History**

Patient history is very important for correct diagnosis of Ménière’s disease because it can provide information about symptoms during acute attacks. Clinical testing is usually only done between acute attacks. On completing the initial history, the onset, duration, frequency and intensity of crises, the association of symptoms, and their impact on the patient’s quality of life should be determined. A typical vertigo attack with no associated hearing loss suggests that the disease is possibly Ménière’s disease. A single definitive episode of vertigo that occurs together with the other symptoms of Ménière’s disease makes the diagnosis probable but an exact diagnosis of Ménière’s disease requires two or more definitive episodes of vertigo and hearing loss associated with tinnitus and/or aural fullness [1].

**Clinical Examination**

Clinical testing is usually only done between acute attacks and can be normal at the earlier stages of the disease. As the disease progresses, it reveals audio-vestibular abnormalities.

The Romberg test\(^1\) shows axial deviation while the Babinski–Weil test\(^2\) and the Fukuda stepping

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1. Romberg test: The patient stands with feet together, eyes open, and hands by the sides. The patient closes their eyes while the examiner observes for a full minute to note occurrence of a fall or axial deviation toward the affected side.
2. Babinski–Weil test: the patient walks with eyes closed, ten steps forward and ten steps backward several times; the examiner looks for a deviation from the straight path, bending to the affected side when walking forward and to the other when walking backward.
Ménière’s Disease and Tinnitus show drift toward the affected side. However, these tests are not always reliable when the patient has myo-articular and/or orthopedic problems.

Audiometric and Oto-Neurologic Examination

Pure-tone audiometry should be obtained and should show low-frequency sensorineural hearing loss that gets better or disappears after crises. However, as the disease progresses, the hearing loss often reaches high frequencies and can even change to a flat hearing loss.

Speech audiometry should show normal speech intelligibility and can confirm that the hearing loss is indeed of cochlear origin. Tympanometry can rule out middle-ear problems. The Weber-test should be lateralized toward the healthy or better ear. The Rinne test should be positive. In patients with Ménière’s disease, it usually does not indicate any difference between the auditory thresholds for air and bone conductance.

Metz-test shows objective recruitment that is more marked in Ménière’s disease compared to the other cochlear pathologies.

The Reflex Decay Test shows that the stapedial reflex is well maintained and can confirm the cochlear origin of the disease.

If the patient accepts, a Glycerol test may be helpful if the patient’s history and tests are inconclusive. Hearing thresholds, particularly at low frequencies, often improves after administration of glycerol to patients with Ménière’s disease.

Recording of distortion product otoacoustic emissions (DPOAEs) provides information about the function of the outer hair cells which may be impaired by abnormal pressure (or rather volume [2]) of the endolymphatic fluid. The combination of vestibular-evoked myogenic potentials and DPOAEs with the glycerol test is suggested for early diagnosis of Ménière’s disease and for the differential diagnosis in patients presenting a first attack of vertigo with or without hearing loss [3].

Studies have disagreed regarding the value of recording DPOAEs and cochlear microphonics for differential diagnosis in patients with and without hydrops [4]. The auditory brainstem response (ABR) typically does not show abnormalities in agreement with the assumption that the disease is not affecting retrocochlear functions. The diagnostic value of recordings of the summating potential (SP) that is a component of the electrocochleogram (ECoG) has been advocated by some investigators [5, 6] while others, Eggermont 1979, [7] have been critical regarding the value of ECoG in diagnosis of Ménière’s disease, in particular for hearing loss less than 50 dB. The large individual variation in the SP is an obstacle in its use as a diagnostic criterion.

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1 Fukuda stepping test: The patient stands with eyes closed, arms outstretched and wearing ear muffs. The patient marches in place 50 steps at the pace of a brisk walk while keeping the eyes closed. The observer looks for any rotation. Rotation of 30° or more is considered a positive test.

2 Weber test: A test in which the stem of a vibrating tuning fork is placed on the midline of the head to ascertain which ear the sound is heard by bone conduction. The sound will be perceived in the affected ear when a unilateral conductive hearing loss is present or in the unaffected ear when there is a unilateral sensorineural hearing loss. The result of this test is combined with the result of the Rinne test to interpretation of the type of hearing loss. (From Stedman’s Electronic Medical Dictionary).

3 Rinne test: Tests the ability to hear by air conduction with the ability to hear by bone conduction. By placing the tines of a vibrating tuning fork near the pinna, the acoustic stimulus is presented by air conduction; by placing the stem of a vibrating tuning fork on the mastoid process, the acoustic stimulus is presented by bone conduction. In conductive hearing losses, the stimulus is heard louder and longer by bone conduction. In sensorineural hearing losses, the stimulus is heard louder and longer by air conduction. The result of the test is reported for each ear as air conduction and is found to be greater than bone conduction, or vice versa. This information is combined with the result of the Weber tuning fork test in interpreting the type of hearing loss.

6 Metz test: The test compares the threshold of the acoustic middle-ear reflex and loudness perception.

7 Glycerol test: Glycerol is administered orally 1.5 ml/kg of body weight dissolved in the equal amount of the physiological saline. A positive result is defined as a threshold improvement of the audiogram of 10 dB or more in at least three frequencies (500, 1,000, or 2,000 Hz). The speech audiometry must show an improvement of 10% of the discrimination. However, the oral glycerol test prohibits food intake before the testing, requires a long examination time, and is associated with side effects such as headache, nausea, and vomiting. The intravenous glycerol test (intravenous injection of 100 ml glycerol (10%) over 30 min) is known to have no such disadvantages.
Vestibular Examination

Nystagmography can quantify the vestibular abnormalities. Spontaneous horizontal or horizontal-rotatory nystagmus is common in individuals with Ménière’s disease. As soon as the crisis starts, nystagmus beating toward the affected side occurs. But the direction of the eye movements is quickly reversed, confirming a unilateral vestibular deficit. The nystagmus is then very intense and occurs even when the eyes are directed toward the affected side (grade III nystagmus). Typically it diminishes with time. After the crisis, the nystagmus which is sensitized by the suppression of ocular fixation, disappear from direct observation, but can always be reactivated by the Head shaking test or by vibratory stimulation of the mastoid. If high-frequency vibratory stimulation of the mastoid reveals a latent nystagmus, it is an indication of a unilateral vestibular deficit. In patients with Ménière’s disease who have a normal caloric test, the direction of the nystagmus triggered by the head shaking and vibratory tests is usually discordant and informs us on the evolution of the disease.

The Dix Hallpike test is typically negative in Ménière’s disease. However, detailed attention should be paid to the realization of this test because the spontaneous or latent nystagmus due to the Ménière’s disease can be sensitized or revealed in the decubitus position. In addition, the possibility of benign paroxysmal positional vertigo (BPPV) should not be overlooked. BPPV can be either idiopathic or, due to the mechanical disruption and distortion of the utricle and saccule, related to the progression of Ménière’s disease.

Magnetic Resonance Imaging

A MRI of the brain with Gadolinium contrast can determine whether the internal auditory canals are open and that the morphology of structures in the posterior fossa is normal.

Differential Diagnosis

An expert clinical judgment is required to distinguish between true Ménière’s disease and several other conditions characterized by vertigo, hearing loss, and tinnitus, such as cochlear otosclerosis; bacterial or viral labyrinthitis; temporal bone trauma; V, VII, and VIII neuroma; meningioma; cholesteatoma; and migraine. The complete oto-neurological assessment including an ABR, ECoG, and MRI of the posterior fossa makes it possible to differentiate Ménière’s disease from these etiologies.

Distinguishing Ménière’s disease from vestibular schwannoma and vascular conflict of the VIII cranial nerve are the most important factors. Kentala and Pyykkö (2000) compared test results from 128 individuals with vestibular schwannoma and 243 with Ménière’s disease and found that 38% of patients with small and medium-sized vestibular schwannoma had an association of all the symptoms of a typical Ménière’s disease. In 69% of the patients, the attacks lasted from 5 min to 4 h and occurred only once or twice a year. In addition, half of the patients had spontaneous nystagmus and 61% of the patients had caloric asymmetry. Tinnitus in these patients was either mild or intense (in 49 and 12% of cases, respectively).

Vascular conflict of the VIII nerve is characterized by intermittent paroxysms of dizziness and unilateral tinnitus, which can become more frequent over time (see also Chap. 40). In a chronic stage, this condition
Ménière’s Disease and Tinnitus

induces persistent unsteadiness [9, 10] often associated with constant tinnitus. Unfortunately, this condition, which has a similar course as Ménière’s disease, is often not recognized. In a study by Ryu and coworkers, it was even shown that up to 73% of the patients diagnosed preoperatively as having Ménière’s disease were successfully treated for vascular conflict of the vestibular nerve [11].

Patients with Ménière’s disease have normal ABR with normal interpeak latency I–III [12].

Ménière’s disease also has similarities with migraine (see a recent review by Minor, 2004 [13] and Chap. 38). However, the glycerol test is negative for migraine, the crises of dizziness always start in the early morning, and there is often a family or personal history of migraines. In addition, individuals with migraine-associated dizziness usually have normal hearing, and when a sensorineural hearing loss is present, it rarely progresses, thus, different from individuals with Ménière’s disease [14] (see also Chap. 38).

Treatment

There are several different treatments available for Ménière’s disease, and the choice of treatment requires careful consideration. Both medical and surgical treatments are in general use, but there is not a consensus regarding the specific treatment and a divergence of different protocols currently in use.

Management of Acute Attacks

During an attack, the treatment is aimed at alleviating the acute symptoms.

Vertigo is often the most disabling symptom of Ménière’s disease, and the medical treatment seeks, above all, to control these symptoms. The intravenous injections of Acetylleucine used in some countries are effective in alleviating vertigo attacks. The action of Acetylleucine is not understood, but studies in animal models suggest that it acts mainly on abnormally hyperpolarized and/or depolarized vestibular neurons by restoring their membrane potential [15]. Administration of Acetylleucine does not have any proven effect on tinnitus.

A randomized double-blind clinical study showed that betahistine dimesylate 12 mg as well as a fixed combination of cinnarizine 20 mg and dimenhydrinate 40 mg are highly effective and safe treatment options for Ménière’s disease and may be used in both the management of acute episodes and long-term treatment. These drugs, commonly used to treat vestibular disorders, reduce tinnitus in approximately 60% of patients with Ménière’s disease [16].

Some physicians prescribe loop diuretics to normalize the balance of fluid volumes in the inner ear. For example, intravenous injections of 40 mg furosemide in the morning during 3–5 days are effective treatments but require checking the blood electrolytes. Tinnitus should be watched since furosemide can give tinnitus.

During crises, intravenous injections of 40 ml of 30% Glucosé-hyper in the morning and evening for 3 days are also effective treatments. Corticosteroids (such as Methylprednisolon 20 mg in intravenous perfusion) are used by some physicians for the management of acute vertigo attacks. Treatment for preventing nausea and vomiting, which can be very intense during a crisis, should also be available. The following antiemetics are often prescribed:

- Compazine (per os or suppository) – 5 mg every 12 h as needed.
- Meclizine (per os) – dose ranges from 12.5 mg twice a day to 50 mg three times a day.
- Métopimazin (per os) – 1 or 2 (15 mg) tablets three times a day.
- Métoclopramid – intramuscular or intravenous injections of a 10 mg/2 ml vial three times a day.
- Benzodiazepines (for example: Lorazepam sublingual tablets, 0.5 mg twice a day) are used to relieve the anxiety accompanying Ménière’s attacks.

Therapies for Inter-crises Periods

The purpose of treatment is to reduce the number of attacks while trying to prevent further hearing loss and damage to the vestibular system. This form of treatment depends on the inter-crises symptoms, their intensity, and their impact on the patients’ quality of life.
Non-invasive Therapies

If the symptoms disappear after the crisis, the patient only needs dietetic recommendations such as to avoid caffeine, alcohol, tobacco, and aspartame, which worsen tinnitus and other Ménière’s disease symptoms. A low-salt diet is also important.

To reduce the frequency of vertigo attacks and alleviate the inter-crisis symptoms, Betahistine (16 mg three times a day or 24 mg twice a day) is often beneficial [17]. Diuretics (such as furosemide 20 mg a day, two times a week with a control of electrolytes), osmoregulators (such as glycerol or mannitol), or vasodilators (for example, Buflomedil 150 mg twice a day) can also be effective in selected patients. However, there is insufficient evidence that this medication has any significant effect on Ménière’s disease-related tinnitus [18, 19].

A randomized and controlled clinical study showed the effectiveness of the combination of diphenidol (25 mg/d), acetazolamide (250 mg/48 h), and prednisone (0.35 mg/kg) on the tinnitus, as well as the frequency and duration of vertigo [20].

Corticosteroids are especially helpful in bilateral forms, in particular, if an autoimmune cause is suspected. Desensitizing therapies for allergies have been shown to be effective to relieve the Ménière’s disease symptoms, including tinnitus in some patients [21, 22].

Depending on the presence of psychosomatic components, the social–professional impact of tinnitus, and other Ménière’s disease symptoms, a patient’s regular follow-up by a multidisciplinary team may be beneficial. Relaxation therapy may be beneficial in some patients because of its beneficial effects on unsteadiness, as well as on tinnitus. It should be associated with standard methods of tinnitus management (see Part V – Management of Tinnitus). Balance rehabilitation can improve a patient’s balance.

Intratympanic Therapy

When vertigo persists despite optimal medical management, an intratympanic therapy with gentamycin or steroids may be proposed to control the vertigo. The intratympanic administration of low-dose gentamycin provides long-term vertigo control, whilst preserving hearing and vestibular function in the majority of patients [23]. In addition, it is effective to treat the tinnitus in Ménière’s disease [24, 25]. New protocols have been developed to reduce the risk of permanent gentamycin ototoxicity. The one-shot injection protocols present a minimal risk to hearing, whereas repeated or continuous application protocols result in higher gentamycin doses in the cochlea and can cause damage to hearing [2, 26]. In a review of literature, Dodson and Sismanis (2004) [27] suggest that this therapy should mainly be proposed to patients with Ménière’s disease who do not have useful hearing. The authors recommend intratympanic therapies with steroids for Ménière’s patients with normal hearing, which have some success in controlling vertigo.

Treatments that can control vertigo may not always improve tinnitus in Ménière’s patients. A prospective double-blind placebo-controlled trial by Garduno-Anaya et al. (2005) [28] showed relief of tinnitus in 48% of the patients who were treated with intratympanic dexamethasone. It was also shown that the inner-ear perfusion via transtympanic delivery of dexamethasone 4 mg/ml improves hearing, tinnitus, and aural pressure in patients with a cochlear form of Ménière’s disease [29]. Nonetheless, Araujo et al. (2005) [30] reported that a prospective randomized placebo-controlled but single-blind trial showed that intratympanic dexamethasone had no significant effect on severe tinnitus compared to placebo.

Surgical Approach

Surgical treatment should be a last resort and is reserved for Ménière’s patients who are refractory to medical therapy. Conservative and destructive surgical procedures are used according to the severity of the crises, the degree of serviceable hearing, and the condition of the contralateral ear.

Endolymphatic Sac Surgery

Conservative surgery by endolymphatic mastoid shunt or endolymphatic sac decompression without sac incision is the operation most often practiced. It can lead
to a temporary decrease in vertigo occurrence and intensity, while generally preserving hearing [31]. However, the literature reveals disagreement regarding the effectiveness of this approach in reducing vertigo.

**Sectioning of the Vestibular Nerve**

Sectioning of the vestibular nerve is effective in controlling vertigo while preserving hearing in most patients. Thus, it is available for patients with serviceable hearing who have failed all other treatments and are especially incapacitated by Ménière’s disease. Dandy (1941) [32], described how he treated patients with Ménière’s disease by sectioning the eighth cranial nerve. Later, this technique has been refined and now, most typically, only the vestibular nerve is sectioned. Different techniques are in use for sectioning the vestibular nerve, such as a retromastoid (retrolabyrinthine) approach to the cerebello pontine angle, and a middle fossa approach has been used as well. Endoscope-assisted, minimally invasive retrosigmoid approach that is now recommended rather than the middle fossa or retrolabyrinthine approaches is simpler, more reliable, and has lower risk of complications [33–35]. Analysis of 18 publications mentioning tinnitus status after vestibular neurotomy in a total of 1,318 patients shows that the tinnitus had worsened after the operation from 0 to 60%, but most of the patients had no change in their tinnitus (17–72%) and others even reported improvements of 6–61% [36]. Thus, vestibular neurotomy does not consistently worsen tinnitus, but the risk is present.

For patients with unilateral Ménière’s disease and total deafness, labyrinthectomy can be undertaken as a last resort.

The procedures that control the episodic vertigo by destroying the vestibular function in the affected ear should be reserved for patients who have handicapping vertigo, which persists in spite of conservative treatments. Typically, the balance improves significantly after these procedures, thanks to compensatory and substitutive mechanisms. The ability to compensate for loss of vestibular input decreases with age, and for people over the age of 50 years, the compensation takes a long time and is rarely complete. These operations, however, cannot prevent the progression of hyperacusis or tinnitus.

**Microvascular Decompression**

Cranial nerve roots in contact with a blood vessel have been associated with specific diseases such as hemifacial spasm, trigeminal neuralgia, glossopharyngeal neuralgia, and geniculate neuralgia. Also, blood vessels in close contact with the root of the vestibular nerve have been associated with a specific disorder such as a specific vestibular disorder (disabling positional vertigo (DPV)) [37, 38], and blood vessels in contact with the auditory nerve have been associated with some special forms of tinnitus [10]. Microvascular decompression operations (MVD) for DPV have shown beneficial effect in about 85% of patients [39] (see Chap. 40). MVD operations for tinnitus are effective in giving relief of tinnitus in some patients with this condition [10] (see Chap. 84).

The fact that vascular loops have been reported to be in contact with the vestibular nerve in patients with Ménière’s disease does not mean that contact with a blood vessel is associated with symptoms. Studies have shown that vascular loops in contact with cranial nerve roots occur frequently without giving specific symptoms from the respective cranial nerve [40] (see Chap. 40).

**Other Forms of Treatment**

**Applying Air Puffs to the Inner Ear**

It has been shown that applying air pressure to the inner ear can relieve some of the symptoms of Ménière’s disease [41, 42]. This was first realized by placing individuals with Ménière’s disease in a pressure chamber. These findings have later been explored, and a practical device that a person can wear was developed (the Meniett, now marketed by Medtronic, Inc.). This device provides a series of air puffs to the sealed ear canal. Using this device requires that ventilation tubes (PE tubes) are inserted in the eardrum to make it possible for the air puffs to reach the middle-ear cavity. The Meniett device is now in use for management Ménière’s disease.

The efficacy of such treatment was studied by Odkvist et al. (2000) [43] in a prospective randomized placebo-controlled, multicenter clinical trial. The study had
56 participants with active Ménière’s disease, age 20–65 years, with a hearing loss of 20–65 dB PTA. Thirty-one participants completed 2 weeks using the Meniett device and 25 patients completed the 2 weeks with the placebo device. A grommet (PE tube) was inserted in the eardrum on the affected side 2 weeks before the study began. The active group experienced significant improvement concerning the frequency and intensity of vertigo, dizziness, aural pressure, and tinnitus, assessed using a visual analogue scale (VAS). The placebo group experienced no difference from the normal course of their disease. Pure-tone threshold improved at the frequencies 500 and 1,000 Hz after active treatment, but there were no improvement of hearing after placebo treatment. Boudewyns et al. reported a significant decrease in the median number of vertigo spells without any improvement in hearing status, tinnitus and functional level, or self-perceived dizziness handicap [44].

In another study, Densert and Sass (2001) [45] found beneficial effect on the symptoms in 37 individuals with Ménière’s disease, 31 of whom had failed to respond to medical treatment in a 2-year follow-up; 19 were free from vertigo spells; 15 had a significantly fewer vertigo spells; and 3 did not respond to pressure treatment. These three individuals later had treatment with gentamicin injections, one of these three became deaf in the affected ear. None of the patients’ conditions when treated with air puffs became worse [45]. All participants in the study reported improvement in functionality of at least two levels, according to the AAO-HNS functionality scale.

References

Chapter 61
Tinnitus with Headaches

Miguel J.A. Láinez, Anna Piera, and Alejandro Ponz

Keypoints

1. Patients with tinnitus frequently have headaches, but the relation between these two disorders is not always casual.
2. Headaches and tinnitus could be symptoms of the same disease.
3. Idiopathic intracranial hypertension is a syndrome in which headaches and tinnitus often occur together.
4. Headaches and tinnitus often occur together with other focal symptoms in symptomatic intracranial hypertension.
5. Intracranial vascular abnormalities such as arteriovenous malformations (AVMs) can occur together with any kind of headache with paroxysmal tinnitus.
6. Tinnitus may be one of the signs of a basilar migraine.
7. Headaches are a very frequent symptom after head trauma, and tinnitus is also common in the posttraumatic syndrome.
8. When a patient with tinnitus presents with headaches, a careful neurological examination that may include neuroimaging should be completed.

Keywords  Tinnitus • Headache • Idiopathic intracranial hypertension • Arteriovenous malformations • Symptomatic intracranial hypertension • Brain tumor • Basilar migraine

Abbreviations

AVM Arteriovenous malformation
CPA Cerebellopontine angle
CSF Cerebrospinal fluid
DCN Dorsal cochlear nucleus
IIH Idiopathic intracranial hypertension
MR Magnetic resonance
MRI Magnetic resonance imaging
TNC Trigeminal nucleus caudalis

Introduction

Headaches are the most frequent reason for neurological consultation [1]. The lifetime prevalence of headaches has been estimated in 66% of the general population with a current prevalence of 47% [2]; with these high prevalence rates, it is not surprising that patients with tinnitus also have headaches. However, the relation between these two disorders is not always casual.

Headaches and tinnitus are both symptoms of the same disease; patients with increased or decreased cerebrospinal fluid (CSF) pressure, basilar migraine, or carotid dissection experience both symptoms. Headaches are also common in patients with chronic tinnitus; the relationship between these symptoms is unclear.

Headaches and Tinnitus as Symptoms of the Same Disease

Idiopathic Intracranial Hypertension (Pseudotumor Cerebri)

In an alert and orientated patient without localized neurological signs, idiopathic intracranial hypertension (IIH) is characterized by the symptoms and signs of elevated CSF pressure. It occurs most frequently in obese women of childbearing age, but women can
develop IIH at any age. A headache is the more prominent symptom; it is usually severe, daily, and referred to the entire head; it can be throbbing or pressing and often worsens with Valsalva maneuvers or postural changes. Nausea is a common symptom and, less frequently, so is vomiting [3, 4].

Tinnitus referred to inside of the head occurs in approximately two-thirds of individuals with IIH. This type of tinnitus is more often perceived as a pulsatile bruit-like sound that is synchronous with the heartbeat and not a high-frequency ringing sound. The tinnitus is often unilateral and disappears after jugular compression on the side to which the tinnitus is referred. The tinnitus has been attributed to the intensified vascular pulsations that are transmitted to the wall of the venous sinuses by the CSF. The pulsatile compressions were thought to convert laminar blood flow to turbulent flow [3].

Episodes of transient blurred vision of brief duration are frequent, and papilledema is the hallmark sign in a neurological examination. However, it is important to remember that IIH may occur occasionally without papilledema. Horizontal diplopia is also common.

An increase of CSF pressure (>200 mm H₂O in the non-obese, >250 mm H₂O in the obese) measured by lumbar puncture confirms the diagnosis. After the lumbar puncture, headaches and tinnitus improve. The diagnosis of IIH requires neuroimaging to rule out other causes of intracranial hypertension [5, 6].

MRI and MR angiography are recommended; if MRI cannot be obtained, a computed tomography scan with contrast is the second best choice.

If a patient is overweight, losing weight is recommended. In order to reduce the increase of CSF pressure, diuretics are recommended; acetazolamide is the most commonly used drug, but furosemide is an alternative. A furosemide test (40–80 mg each morning for 3–10 days) has been proposed to rule out the IIH.

Topiramate is also an alternative in combination with diuretics. If the medical treatment fails, surgery should be considered: optic nerve sheath fenestration and shunting procedures could be performed [3, 6].

**Symptomatic Intracranial Hypertension and Intracranial Hypotension**

All space-occupying lesions in the brain produce, or can produce, increased CSF pressure and, because of this, headaches are a usual manifestation. The most common type of “brain tumor headache” is a tension-type headache; usually, it is moderate or severe, worsens in the morning, and, in few cases, is accompanied by nausea and vomiting. The more typical and classically mild, early morning frontal headache that resolves after wake-up is uncommon in brain tumor patients. In other space-occupying lesions, such as subdural hematomas or brain abscesses, headaches are an earlier and more frequent symptom [7].

Tinnitus has been described as a symptom of intracranial hypertension, but the prevalence is unknown. Usually the tinnitus is described as constant and ringing high-frequency sound. Tinnitus is more common in individuals with tumors of the posterior fossa, especially in space-occupying lesions of the cerebellopontine angle (tumors, arachnoid cysts). Some individuals with such lesions complain of intermittent tinnitus [8, 9]. Headache and tinnitus usually improve or disappear after resolution of the intracranial hypertension.

Tinnitus has also been described in patients with such hindbrain abnormalities as Arnold–Chiari malformation. In these cases, headache and tinnitus should appear with manoeuvres that transiently increase intracranial and intra-abdominal pressure. Coughing, sneezing, and physical exercise typically trigger attacks of tinnitus [7].

In low CSF pressure syndrome, headache is also the main clinical feature. It is typically an orthostatic headache that is present when the patient is upright and is relieved when lying down. It may be throbbing, and the location is predominantly posterior. Changes in hearing (echoed, distant, or muffled) and tinnitus have been described as associated symptoms; these symptoms can also appear in an upright position and improve or disappear by lying down [10, 11]. These symptoms may be related with stretching the VIII nerve or changes in the pressure of the perilymphatic fluid in the inner ear [12].

**Vascular Abnormalities**

All kinds of vascular malformations can produce headache and tinnitus. Both are more prominent in arteriovenous malformations (AVM) and arteriovenous fistula [13]. The headaches in individuals with AVMs often fulfill criteria for migraine with aura, but all types of headaches have been described. The tinnitus often occurs episodically. Both tinnitus and headaches are
more common in individuals with vascular abnormalities located to the posterior fossa, and the tinnitus may be a relevant symptom in lesions near the VIII nerve [14, 15].

Carotid and vertebral dissections are other causes of acute headaches and tinnitus. Headaches are usually severe and persistent, neck pain is also common and tinnitus is frequently paroxysmal [13].

Acute headaches, papilledema, and seizures are the most typical signs of sinus thrombosis, but tinnitus can also be a prominent symptom [16].

If a vascular abnormality is suspected, imaging such as MRI and MR angiography is indicated. In some cases, a Seldinger angiography could be indicated.

**Posttraumatic Syndrome**

Headaches are a cardinal symptom of the posttraumatic syndrome [17–19] caused by either head trauma and/or whiplash injury. In the acute phase, there are autonomic symptoms like dizziness, nausea and vomiting, orthostatic reactions, and problems with regulation of body temperature. These signs are accompanied by different degrees of cognitive problems or other often poorly defined neuropsychologic deficits, such as irritability and increased low tolerance and sensitivity to light and noise [20, 21]. Other forms of pain that resemble primary headache disorders may develop after head injury. The most frequently occurring pattern resembles a tension-type headache and occurs in more than 80% of individuals who have had head trauma. Some such individuals have typical migraine with or without aura triggered by the head impact. Even a cluster-like syndrome has been described in some individuals who have had head trauma [22].

Tinnitus is also a frequent symptom in individuals who have had head and neck trauma and is part of the post-concussion syndrome. The frequency of tinnitus is also high in individuals after blast injury (see Chap. 67). Usually, the tinnitus is continuous, and after neck injury, it has the characteristics of somatic tinnitus [23, 24] (see Chaps. 9 and 43).

**Migraine with Basilar Aura**

Vertigo, dysarthria, and tinnitus may occur together with basilar aura in individuals with migraine. Tinnitus has been reported to occur alone or in combination with other symptoms in 50% of individuals with basilar aura. Other symptoms are diplopia, bilateral visual symptoms, bilateral paresthesia, hearing loss, decreased level of consciousness, and ataxia. For the management of patients with tinnitus and migraine, it is important to obtain information about how their tinnitus may be regarded as a symptom that precedes headaches [25, 26].

**Headaches in Individuals with Tinnitus**

The prevalence of headaches in individuals with tinnitus, and vice versa, is unknown. Both tinnitus and headaches have common signs, which suggest that each one of these symptoms could amplify each other and one of these symptoms might cause the other. Both disorders are frequent, under recognized, and under treated; they cause a high degree of disability, and both are often accompanied by psychiatric disorders [27, 28]. The most remarkable finding is that these two symptoms share some common mechanism in their tendency to become chronic. Studies in animals have shown that DCN neurons receive input from the trigeminal system [29].

Lack of normal inhibition of the caudal trigeminal nucleus (TNC) may be an important mechanism in the cause of headache [30]. In both headaches and tinnitus, an increase of the somatosensory influence might play an important role in making these symptoms chronic [29, 31]. In a series of 149 patients with chronic tinnitus (mean duration: 1.5 years), we found a prevalence of headaches in 47%; in the patients with unilateral tinnitus who could modulate their tinnitus by

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1 Seldinger angiography: A method of percutaneous insertion of a catheter into a blood vessel or space. A needle is used to puncture the structure and a guide wire is threaded through the needle; when the needle is withdrawn, a catheter is threaded over the wire; the wire is then withdrawn, leaving the catheter in place. Stedman’s Electronic Medical Dictionary.

2 Dysarthria: A disturbance of speech due to emotional stress, brain injury, or paralysis, in co-ordination, or spasticity of the muscles used for speaking. Stedman’s Electronic Medical Dictionary.
somatic stimulation, the prevalence of headaches was 89%. These patients had tension-type headaches, predominantly unilateral and related with the tinnitus side. More results are necessary to establish the exact relation between the evolution of both these symptoms and the response to treatment.

References

Chapter 62
Tinnitus and Psychiatric Co-morbidity

Michael Landgrebe and Berthold Langguth

Keypoints

1. Tinnitus is often accompanied by psychiatric co-morbidity, especially in severe forms of tinnitus.
2. Many different co-morbid psychiatric disorders have been reported in individuals with severe tinnitus; among them are depression, anxiety, somatoform disorders, psychosis, personality disorders, and body-concept disorders.
3. The exact relationship between tinnitus and psychiatric disorders may vary from patient to patient. Psychiatric disorders may evolve as a consequence of tinnitus, may represent vulnerability factors, may be consequences of another causal event (e.g., trauma), or may just co-occur with tinnitus without any known cause.
4. All forms of co-morbid psychiatric disorders should be treated specifically.

Keywords Tinnitus • Psychiatric co-morbidity • Affective disorders • Tinnitus severity • Psychotherapy • Psychopharmacotherapy

Introduction

Tinnitus, the perception of a phantom sound, is reported by up to 20% of the general population (see Chap. 5) but most of those who have tinnitus are not severely affected by their tinnitus. Nevertheless, there is a group of individuals with tinnitus, who are severely suffering and sometimes become even suicidal because of their tinnitus [1]. For management of the patient with tinnitus, it is important what causes tinnitus-related distress and disability. It has been shown that individuals with tinnitus with low and high distress do not differ regarding the character of their tinnitus such as pitch, loudness, or tinnitus variability [2, 3]. What, however, differs between these two groups is the presence of psychiatric co-morbidity. Patients with high scores in tinnitus questionnaires such as the tinnitus handicap inventory suffer much more frequently from psychiatric disorders than those with low scores in the tinnitus questionnaires [2, 4–7]. The most frequently reported psychiatric symptoms in this patient group are symptoms of depression and anxiety [4, 7–9]. The incidence of psychiatric disorders such as posttraumatic stress disorder [10, 11], somatoform disorder [12], psychosis [13], obsessive-compulsive disorder [14], or body-image disorder [15] is high among individuals with tinnitus. Published studies agree that there is a high degree of correlation between severe tinnitus and symptoms of depression and anxiety, but the results from different studies vary. This is probably due to differences in kinds of studies (e.g., population survey vs. primary care vs. highly specialized tinnitus centers; [16]). Differences in the used diagnostic criteria [17] also contribute to the variation in results. The exact relationship between tinnitus and psychiatric disorders vary from patient to patient; it may depend on the order of onset of the two symptoms. Psychiatric disorders may develop as a consequence of the individual’s tinnitus or because of the individual’s vulnerability factors. The interplay between tinnitus and psychiatric symptoms is often complex, and that makes it difficult to determine if tinnitus has caused a reactive psychiatric co-morbidity in an individual patient or if a pre-existing...
but compensated psychiatric disorder flares up due to the tinnitus, or whether a well-managed tinnitus reappears due to the onset of a psychiatric disease. Recent studies indicate that assessment of psychiatric co-morbidity should not be restricted to chronic tinnitus, but that affective disorders may also occur in acute stages of tinnitus [18]. Also therapeutic interventions should be considered in acute tinnitus, since the amount of distress in the acute stage (<1 week duration) seems to predict the amount of suffering in the chronic situation [19].

Finally, psychiatric disorders that occur together with tinnitus should always be treated specifically, independent on how they are related to the patient’s tinnitus. Efficient treatment of the concurrent psychiatric disorder reduces the burden of disease and improves the quality of life of the tinnitus patient. Special emphasis must be directed to emergency of treatment in tinnitus patients who are suicidal (see Chap. 54).

The topic of the following chapters concerns the most frequent co-morbid psychiatric conditions, i.e., emotional trauma, depression, and anxiety including etiological and pathophysiological considerations and the therapeutic management.

References

Chapter 63
Tinnitus and Depression

Berthold Langguth and Michael Landgrebe

Keypoints

1. There is an increased prevalence of depressive symptoms in individuals with tinnitus.
2. Affective disorders, together with personality factors, play an important role in creating the distress experienced by many individuals with tinnitus.
3. The co-occurrence of tinnitus and depression may be explained by the involvement of limbic brain structures in the pathophysiology of tinnitus.
4. Tinnitus is associated with neuroendocrine alterations, which are characteristic for depressive disorders.
5. Patients with tinnitus and depression should be efficiently treated.
6. Efficient treatment of co-morbid depressive symptoms has to consider a large variety of possible underlying disorders.

Keywords
- Tinnitus-related distress
- Depression
- Psychiatric co-morbidity
- Limbic brain areas
- Quality of life
- Suicide

Abbreviations
- DCN: Dorsal cochlear nucleus
- PTSD: Post traumatic stress disorder
- ICD: International classification of diseases
- DSM: Diagnostic and statistical manual of mental disorders

Introduction

There is abundant evidence of increased prevalence of depressive symptoms in individuals with tinnitus [1–10]. This is especially the case in individuals with disabling tinnitus [2, 3]. Also, high correlations between scores in tinnitus severity measures and depression scales have been reported [6, 9]. Thus, the occurrence of co-morbid depression may explain to some extent why some individuals suffer severely from tinnitus, whereas other individuals are not bothered by their tinnitus. Further studies have shown an additional role for specific personality traits such as anxiety, obsessiveness, neuroticism, or agreeableness [9, 11, 12]. However, it is important to note that all these findings are mainly derived from studies that used self-report questionnaires for the assessment of depressive symptoms, not from structured interviews. This difference is important, since the presence of depressive symptoms does not automatically mean that a patient fulfills diagnostic criteria for a depressive disorder. Depressive symptoms may be indicative of a depressive disorder, but they can also occur in the context of a large variety of other psychiatric disorders, such as bipolar disorders, personality disorders, anxiety disorders, dementia, or addiction, just to name a few. Therefore, further studies using structured interviews will be needed to determine exactly the prevalence of co-morbid psychiatric disorders in tinnitus.
Depression

Diagnosis of Depression

For non-psychiatrists, detection of depressive symptomatology is obviously difficult. However, the use of simple screening questions may help to identify potential depression. The following two questions have been proposed as a screening method for depression: “During the past month have you often been bothered by feeling down, depressed, or hopeless?”; “During the past month have you often been bothered by little interest or pleasure in doing things?” [13]. If the patient answers with “yes” to one of the questions, depression is likely and further diagnosis by a psychiatrist or psychologist should be initiated. A more detailed screening instrument is the Mini-International Neuropsychiatric Interview [M.I.N.I., [14]]. As mentioned above, depressive symptoms may occur in a variety of psychiatric diseases. The exact differential diagnosis is of importance, since it has important consequences for the therapeutical management. For example, depressive symptoms can occur in the context of a bipolar affective disorder or a posttraumatic stress disorder, which requires completely different therapeutic management than major depression. Thus, when screening questions suggest potential co-morbid depression, a psychiatrist should be involved in further diagnostic assessment and therapeutic management.

The exact diagnostic classification may further depend on the classification system. The most widely used classification systems are the “Diagnostic and Statistical Manual of Mental Disorders” of the American Psychiatric Association (DSM-IV) [15] and the “International Classification of Diseases” of the WHO (ICD-10) [16]. Classification criteria for major depression (DSM IV) and depressive disorders (ICD 10) are displayed in Tables 63.1 and 63.2.

Interplay Between Tinnitus and Depression

Given the association between tinnitus and depressive symptoms, the question about the nature of the relationship arises. Clinical experience suggests that all kinds of relationships may occur: Depressive symptoms may develop as a reaction to tinnitus. Specific vulnerability factors such as anxious or obsessive personality traits may contribute to the development of such depressive reactions. In other cases, where tinnitus exists for a long time without causing any substantial distress, a depressive episode may lead to decompensation of the tinnitus with subsequent impairment in quality of life. There is also the possibility that tinnitus and depressive symptoms are consequences of a third condition (e.g. traumatic event). Finally, tinnitus and depressive disorders may also co-occur incidentally since both are relative frequent conditions. In clinical practice, differential diagnosis of all relevant factors contributing to tinnitus distress is of importance, since all these factors may represent potential targets for treatment. As an example, let us consider a patient who complains about chronic tinnitus with variations of perceived distress, ranging between severe disturbances to none at all. Psychiatric exploration may reveal a co-morbid seasonal affective disorder or a co-morbid bipolar disorder, which explains the variations in tinnitus distress. In this case, the co-morbid psychiatric condition should be specifically treated with the primary aim of mood stabilization, which in turn will lead to decreased variations of distress and impairment.

Similarities Between the Pathophysiology of Tinnitus and Depression

Pathophysiologic models of tinnitus have claimed the involvement of frontal and limbic brain regions. In his neurophysiological model of tinnitus, Jastreboff hypothesized in detail that the prefrontal cortex may be the brain structure that integrates sensory and emotional aspects of tinnitus and may be involved in the emotional and autonomic reaction to tinnitus [17]. The activation of the non-classical (extralemniscal) ascending auditory pathways in some forms of tinnitus [18] (see Chap. 10) may explain the coactivation of cortical association areas, limbic areas, and the autonomic nervous system [19]. Recent neuroimaging studies (in non-depressed individuals with tinnitus) have confirmed the involvement of the prefrontal cortex, the subgenual frontal cortex, and the amygdalo-hippocampal area in the pathophysiology of tinnitus (for review see [20, 21] or Chap. 17, 18, 19).
Tinnitus and Depression

These brain areas are well known as critical parts of brain networks, functionally altered in individuals with depressive disorders [22, 23]. Thus, imaging data suggest that the neuronal correlates of tinnitus and depression overlap in limbic networks, which provides a possible explanation for the co-occurrence of tinnitus and depressive symptoms.

In this context, an important role for the dorsal cochlear nucleus (DCN) has also been proposed [24]. There is increasing evidence from animal studies that the DCN is an important contributor to tinnitus. There are direct projections from the DCN to brain stem structures such as the locus coeruleus, reticular formation, and raphe nuclei which are the principle sites for synthesis of serotonin and noradrenaline and are implicated in the control of attention and emotional responses. Thus, attentional and emotional disorders, such as anxiety and depression, commonly associated with tinnitus may result from an interplay between these non-auditory brainstem structures and the DCN [24].

Also, serotonergic dysfunction, which is assumed to play an important role in the pathophysiology of depression [25], has been suggested to be involved in
tinnitus [26, 27]. However, evidence for this hypothesis is scarce.

Finally, neuroendocrine alterations such as a hypothalamic–pituitary–adrenal axis dysfunction, which are pathognomonic for stress-related disorders such as depression or posttraumatic stress disorder (PTSD) [28, 29] have been described in tinnitus patients [30, 31], indicating another pathophysiological overlap between tinnitus and affective disorders. Interestingly, a recent study has shown that tinnitus patients differ only slightly from controls in physiological reactivity during stress tests, indicating relatively normal psychophysiological reactivity [32].

**Depression**

**Treatment Options for Depression**

There are several reasons why patients with tinnitus and depression should be promptly and efficiently treated. Efficient treatment depends on the exact etiology of co-morbid depressive symptoms to tinnitus. If diagnostic assessment reveals a major depression (see Table 63.1), standard treatment options include antidepressants and psychotherapy. There is a large variety of antidepressants available, which differ in their mechanisms of action and side effects. Also, there are some antidepressants that specifically address specific symptoms of depression. For example, amitriptylin and mirtazapin have sedative effects and are preferentially used in patients with insomnia, whereas venlafaxine, duloxetine, and bupropion exert an activating effect and are preferred in patients who suffer from loss of energy. Thus, the choice of the best antidepressant is complex and depends on previous patient’s experience with specific drugs, the predominant symptoms of depression and co-morbitities. Also, cognitive behavioral therapy has been shown to be efficient in the treatment of depression. Further non-pharmacologic treatment options include light therapy, sleep deprivation, aerobic exercise, transcranial magnetic stimulation or electroconvulsive therapy.

Several antidepressants have been investigated for their use in tinnitus [33, 34] (see Chap. 78). Two randomized double-blind placebo-controlled studies investigated the effects of antidepressants in patients with tinnitus and co-morbid depression [35, 36]. The tricyclic nortriptyline significantly reduced depression scores, tinnitus disability scores, and tinnitus loudness relative to placebo [1]. Also the serotonin reuptake inhibitor sertraline was significantly more effective than placebo in reducing tinnitus severity [2]. In both studies, reduction in tinnitus disability scores correlated high with reduction of depression scores, suggesting that antidepressants have beneficial results in depressed tinnitus patients, but that this is mainly due to the antidepressant effect of the drug.

However, induction or worsening of tinnitus has also been reported in the context of treatment with antidepressants, both as a side effect of drugs such as phenelzine, amitriptyline, protriptyline, doxepin, imipramine, fluoxetine, trazadone, bupropion, and venlafaxine. Worsening of tinnitus has also been associated with withdrawal of antidepressants (venlafaxine and sertraline) [37]. Interestingly, transcranial magnetic stimulation of the dorsolateral prefrontal cortex, performed for the treatment of depression, has also been described to induce and worsen tinnitus in rare cases [38]. This suggests that tinnitus can be generated or worsened by modulation of neural activity in the frontal cortex. Thus, the complex interactions between antidepressant treatment and tinnitus may be explained by antidepressant-induced modulation of frontal cortex networks [39], which in turn may result in altered top–down control of activity in the central auditory pathways.

**Suicidal Tendency**

Depression can become life-threatening by leading to suicidal ideation. The most important risk factor for suicide is depressive disorder, both in tinnitus patients [40–42] and in non-tinnitus patients [43, 44]. Further risk factors include male gender, elder age, and social isolation [3]. In tinnitus patients with depression, suicidal tendencies have to be assessed because a high risk of suicide requires immediate action. This sensitive area can be approached by asking the patient about passive suicidal ideations (for more details, see Chap. 54). It is important to know that asking about suicidal thoughts does not increase the risk of committing suicide.
Tinnitus and Depression

References

44. Hawton, K, van, HK Suicide Lancet, 2009 Apr 18;373(9672):1372–81
Chapter 64
Tinnitus and Anxiety

Michael Landgrebe and Berthold Langguth

Keypoints

1. Individuals with tinnitus often suffer from anxiety.
2. Such co-morbid anxiety is associated with increased severity of tinnitus and distress as well as general impairment of quality of life.
3. Similar brain areas are involved in the pathogenesis of tinnitus and anxiety disorders indicating a close interrelationship between these two disorders.
4. Hyperacusis and phonophobia represent specific problems in a subset of tinnitus patients who require special attention.
5. Treatment of concurrent anxiety symptoms in tinnitus patients is essential.
6. Treatment options include pharmacological (e.g. antidepressants) and psychotherapeutic (e.g. cognitive behavioral therapy, tinnitus-retraining therapy) approaches.

Keywords Anxiety • Chronic tinnitus • Pharmacotherapy • Psychotherapy

Abbreviations

CIDI-SF Composite diagnostic interview
ICD International classification of diseases
SNRI Selective serotonin and noradrenaline reuptake inhibitor

Introduction

Anxiety belongs to the most basic physiological emotions of human beings. It represents a biosocial signal contributing to the development of normal interpersonal relations and a risk-sensitive interaction with the environment. Anxiety is an indicator of threat and points to potential dangers. The subjective perception of anxiety and its related threats, however, is very much influenced by learning processes, which gradually leads to modified perceptions and cognitive evaluations of internal and external dangers. This, in turn, determines the individual level of tolerated anxiety and forms the behavioral styles to recover safety. Hence, anxiety subserves important physiological functions, which are essential for survival. In this respect, anxiety does not represent a pathological symptom. However, it gains clinical relevance in cases of too much or too little anxiety.

Symptomatology, Epidemiology, and Etiopathogenesis of Anxiety Disorders

Anxiety as a symptom may be part of almost every psychiatric disease. If anxiety reaches an extraordinary level, is subjectively perceived as unrealistic, causes a high burden, and impairs social functioning, it may be regarded as a disease and the diagnosis of an anxiety disorder may be fulfilled. In general, anxiety appears
always on different levels; i.e. an emotional, cognitive (e.g. subjective beliefs regarding danger), a motor (e.g. behavioral attitudes such as fight, fright, or flight), and an autonomic level (e.g. somatic reactions such as tachycardia, stress hormone secretion, etc.). In the International Classification of diseases of the WHO (ICD-10), diagnostic criteria for several anxiety disorders have been defined. These are phobic disorders (ICD-10: F40; e.g. agoraphobia, social, and specific phobias) and other anxiety disorders (ICD-10: F41; e.g. panic disorder, generalized anxiety disorder, anxiety and depression mixed). Phobic disorders are characterized in that way, the anxiety mainly occurs in specific situations or in response to specific things which are normally not dangerous to the individual (e.g. fear of public places, busses, etc. or fear of specific things, e.g., spiders). In contrast, in the other anxiety disorders, symptoms occur independently of specific triggers. Both forms of anxiety disorders have in common that individuals try to avoid situations associated with anxiety. This, in turn, stabilizes the symptoms, because avoidance behavior prevents them from realizing that the anticipated danger is not real and hampers them from learning adaptive strategies.

Similar avoidance behavior is also found in tinnitus patients. Especially, in people suffering from hyperacusis, phonophobia develops frequently and noise is avoided. However, the avoidance of sound is counterproductive, since there is clear evidence that reduced environmental sound increases phantom auditory perceptions such as tinnitus [1]. Hence, this avoidance behavior represents one core target of cognitive behavioral therapeutic interventions, both in anxiety disorders and in tinnitus.

Anxiety disorders are prevalent. Lifetime prevalence rates range from about 1.2% for panic disorder up to 5% for specific phobias. Due to diverging diagnostic criteria, prevalence rates vary substantially between studies. The most frequent anxiety disorder seems to be social phobia, where lifetime prevalence rates of 13.3% have been reported [2]. The lifetime prevalence for anxiety disorders overall is about 10% [3]. Anxiety is even more frequent in individuals with tinnitus. Various studies reported anxiety in 19% up to 45% [4–6]. A recent study using an Internet-adapted version of the WHO short form of the Composite Diagnostic Interview (CIDI-SF [7]) found 12-month prevalence rates of 60% for generalized anxiety disorder, 83% for specific phobia, 67% for social phobia, 58% for agoraphobia, and 21% for panic disorder in a self-selected population of tinnitus patients [7]. Although these data have to be interpreted with caution, because they have been collected by internet survey and verification of the diagnoses by a psychiatrist was not performed, this study, together with the others, indicates that anxiety is a common phenomenon among individuals with tinnitus. Its clinical relevance is further underlined by the fact that tinnitus severity and distress are linked to co-morbid anxiety and depression [8], which is very often also a co-morbid condition in anxiety disorders. Hence, detection and diagnosis of anxiety symptoms in tinnitus patients represents an important task for every healthcare professional dealing with tinnitus patients. Especially, in severe forms with substantial impairment in quality of life, the probability of a co-morbid psychiatric disorder such as anxiety or depression is increased. In those cases, detailed exploration for anxiety symptoms should be performed. Screening questions, which are used in structured clinical interviews, may be helpful (see Chap. 54). However, in case of suspicion on an anxiety disorder, further diagnosis and treatment should be done by a psychiatrist or psychologist. Potential treatment options will be discussed later in this chapter.

The etiopathogenesis of anxiety symptoms is multifactorial. Anxiety patients are often characterized by some accentuated personality traits (e.g. introversion and neuroticism). Other important factors are dysfunctional cognitive processes, anxiety sensitivity, and attentional bias. A behavioral consequence of these dysfunctional cognitions is biased perceptions of potential dangers, misinterpretation of physiological signs (such as heart beat) as signals of danger, and the feeling of being unable to control these symptoms. Many aspects of such dysfunctional cognitions are also frequently found in tinnitus patients, even if they do not suffer from anxiety symptoms. For example, being unable to control their tinnitus and the fear that the tinnitus will get worse are very often mentioned by tinnitus patients as one major reason to seek medical help. It is important to note that these fearful beliefs depend to a large extent on the interaction and communication between patient and health care providers. By exaggerating the risk of dangerous diseases, which may underlie tinnitus (e.g. brain tumors), medical doctors may cause anxiety, and can also do so by informing the patient that there is no help for their tinnitus and that they have to live with it. Hence, the way
information about tinnitus – its pathophysiological basis and potential treatment options – is provided may be critical for the way a patient learns to deal with tinnitus. Informing the patient in an empathic way and providing hope are important factors for preventing the development of anxiety (see Chap. 70). This is even more important since there is evidence that anticipatory processes and dysfunctional activation of a cortical distress network seems to play a role in the pathophysiology of tinnitus [9]. Cognitive behavioral treatment strategies in tinnitus patients focus on these dysfunctional cognitive and anticipatory processes and have been shown to be able to improve quality of life in tinnitus patients [10].

On a neurobiological level, many different brain regions have been identified to be involved in the generation of arousal and anxiety. The arousal level is under the control of brain stem nuclei, mainly the locus coeruleus [11]. But arousal alone is not equal to anxiety. Anxiety depends also on connoted emotions, which are generated within the limbic system and involves structures such as the amygdale, hippocampus, septal nuclei, and the hypothalamus. Recent studies point to a critical role of the amygdale in generation of anxiety, especially in the formation of emotional, anxiety-related memories [12]. There is emerging evidence that the limbic system is also involved in the pathogenesis of tinnitus. A variety of studies have shown functional activations [13–15], as well as structural alterations [16, 17], in limbic areas in individuals with tinnitus, pointing to a role of these structures in the pathogenesis of tinnitus. This is further underlined by the fact that the severity of tinnitus is closely related to concurrent symptoms of anxiety and depression [18]. Just by treating these symptoms, the functional impairment of tinnitus patients may be improved [19].

In summary, anxiety is a frequent symptom, which may occur as a normal physiological phenomenon but also in the context of almost every somatic or psychiatric disorder. Anxiety is a common phenomenon in individuals with tinnitus, and prevalence rates of anxiety disorders are very high. Furthermore, anxiety is often associated with depression. Neurobiology shows that tinnitus and both depression and anxiety share similar neural circuits underlining the close interrelationship of these syndromes. Finally, tinnitus severity is associated with co-morbid anxiety and depression, indicating the necessity to detect and treat these symptoms when treating patients with tinnitus.

### Hyperacusis and Phonophobia

Hyperacusis and phonophobia, which often occur together with tinnitus, may complicate treatment of tinnitus in some patients. Hyperacusis has been defined as lowered tolerance to ordinary environmental sounds, or as a consistently exaggerated [20] or inappropriate response to sounds that are neither threatening nor uncomfortably loud to a typical person [21]. In phonophobia, specific sounds evoke negative emotions such as anxiety and fear. Phonophobia represents a form of a specific phobia. Hyperacusis may occur in context of some underlying somatic conditions such as disorders of the facial nerve (e.g. Bell’s palsy) or neuropsychiatric disorders such as migraine, depression, or posttraumatic stress disorder (see Chap. 3, see (22)). Prevalence rates of hyperacusis in the general population vary between 9% [23] and 15% [24]. Among tinnitus patients attending a tinnitus clinic, prevalence rates of up to 40% have been reported [22].

The pathophysiological mechanisms of hyperacusis are not yet identified, and several potential mechanisms have been discussed [22]. Among these, alterations of serotonergic neurotransmission have been postulated [25]. Evidence for this hypothesis derives from findings that hyperacusis tends to occur in diseases where serotonergic neurotransmission is altered (e.g. migraine, depression, or posttraumatic stress disorder), indicating again a close pathophysiological relationship of tinnitus and hyperacusis to depression and anxiety. But also via direct connections from the central auditory system to the amygdale, specific or more general sounds can induce emotions of fear and anxiety. These unconscious conditioning processes, linking sounds to emotions, have been postulated to be one major mechanism by which hyperacusis may occur [26]. Treatment strategies such as tinnitus-retraining therapy aim to disconnect these dysfunctional connections [27].

### Treatment Strategies

The importance of efficient treatment of co-morbid anxiety symptoms in tinnitus patients has already been extensively discussed in this chapter. However, before treatment can be initiated, a clear diagnosis of the anxiety disorder should be made, which requires
in most cases the consultation of a psychiatrist. In cases of a diagnosed anxiety disorder, principal treatment options are pharmacotherapy and psychotherapy. First-line pharmacological treatment options for anxiety disorders such as panic disorder, social phobia, and generalized anxiety disorder are antidepressants, predominantly serotonin reuptake inhibitors (SSRI) and combined serotonin and noradrenalin reuptake inhibitors (SNRI; e.g. venlafaxine or duloxetine). But also mirtazapine or tricyclic antidepressants may have beneficial effect. For patients with tinnitus and anxiety, improvement of tinnitus has been reported under treatment with antidepressants (see Chap. 63). The selection of an antidepressant for the individual patient may depend on other symptoms such as agitation and/or sleep problems or concurrent other medical conditions or the necessity of other medication, which may increase the risk of unfavorable drug–drug interactions. Therefore, it is advisable that pharmacological treatment is managed by a psychiatrist. Besides classical antidepressants, some other substances may sometimes have beneficial effects in treatment of tinnitus patients with anxiety. For example, the antiepileptic drug pregabalin has been shown to be effective in the treatment of generalized anxiety disorder and has recently also been approved for this indication. Thus, pregabalin represents an additional option for the treatment of anxiety disorders in tinnitus patients.

In contrast to antidepressants and pregabalin, which exert their anxiolytic effects after continuous treatment over weeks to months, benzodiazepines have acute anxiolytic properties. Several pilot studies also suggest beneficial effects of benzodiazepines on tinnitus (see Chap. 78). However, benzodiazepines are regarded to be addictive, and therefore regular use over longer periods of time should be avoided. Also, protracted tinnitus has been reported after discontinuation of benzodiazepines [28]. Thus, the administration of benzodiazepines in treatment of anxiety in tinnitus patients should be restricted to use as rescue medication. For this purpose, short-acting substances such as alprazolam should be preferred. Sometimes, already, the availability of this efficient rescue medication gives the patient some form of control over his/her symptoms and reduces the fear of developing severe anxiety or panic attacks.

Psychotherapy is the other important treatment option, especially in patients with tinnitus who are severely affected and have psychological distress symptoms [29]. Cognitive behavioral therapy aims at modifying dysfunctional cognitions and avoidance behavior in the context of tinnitus. A recent meta-analysis of 285 tinnitus patients showed that tinnitus-specific cognitive behavioral therapy contributes to a positive management of tinnitus and leads to improvement in quality of life [10]. However, this meta-analysis did not reveal improvement in the subjective loudness of tinnitus or on associated depression. The latter suggests that patients with tinnitus and co-morbid anxiety or depression may require specific forms of psychotherapy, which focus primarily on their co-morbid psychiatry disorders.

Avoidance of unpleasant sounds represents a substantial problem in individuals who have hyperacusis and phonophobia in addition to their tinnitus. Both hyperacusis and phonophobia can be efficiently treated by cognitive behavioral therapy [30]. Another treatment option represents tinnitus-retraining therapy, which is based on the neurophysiological model of tinnitus [26] (see also Chap. 73). This model postulates involvement of the limbic and autonomic nervous systems in all cases of clinically significant tinnitus and points out the importance of both conscious and subconscious connections between limbic and auditory pathway structures. Tinnitus-retraining therapy aims at extinction of these dysfunctional unconscious connections in order to allow habituation to tinnitus. Similarly, tinnitus-retraining therapy is also assumed to be efficient for the treatment of hyperacusis [27].

In summary, in patients with tinnitus and concurrent anxiety symptoms or anxiety disorder, a broad variety of pharmacological and available psychotherapeutic treatment options can have beneficial effect on both tinnitus and anxiety. Decisions for the best individual treatment require psychiatric experience and depend on individual symptoms that may be concomitant with other medical conditions and depending on the individual patient’s acceptance.

References

Tinnitus and Anxiety


Chapter 65
Tinnitus and Sleep

Tatjana Crönlein, Peter Geisler, and Göran Hajak

Keypoints

1. Disturbed sleep is a frequent problem in persons suffering from tinnitus.
2. Insomnia may persist over years even after successful treatment of tinnitus and specific therapy of sleep.
3. Sleep-disturbed tinnitus patients are more impaired the more severe their tinnitus is.
4. Since disturbed sleep is a risk factor for mental and somatic health, sleep-disturbed tinnitus patients need special therapeutic and diagnostic care.
5. The results of sleep tests in sleep-disturbed tinnitus patients show similarities with insomnia patients, and their psychological symptoms are similar.
6. The prevalence of organic sleep disorders is high in older persons, and therefore differential diagnostic measures are important in this population.
7. Insomnia in sleep-disturbed tinnitus patients can be treated with hypnotics or with insomnia-specific psychotherapy.
8. Insomnia-specific cognitive behavior therapy may improve both sleep and tinnitus.

Keywords Tinnitus • Sleep • Insomnia • Behavior therapy

Abbreviations

BDI Beck Depression Inventory
CBT Cognitive behavior therapy
CPAP Continuous positive air pressure
ESS Epworth sleepiness scale
PI Psychophysiological insomnia
PLMS Periodic limb movements in sleep
PSQI Pittsburgher sleep quality index
RLS Restless legs syndrome
SAS Sleep apnea syndrome
SDTP Sleep disturbed tinnitus patients
TP Tinnitus patients

Introduction

“I know how my night sleep is going to be just from the way my tinnitus is during the day.” This sentence uttered by a patient in our sleep disorder center describes the special combination of tinnitus and insomnia. The definition of sleep includes decreased responsiveness to external stimuli. The first studies on sleep depth had been performed on the basis of acoustic threshold. E. Kohlschütter described in his dissertation “Zur Festigkeit des Schlafes” (Henle Zeitschrift) in 1862 that the acoustic arousal threshold increases at the beginning of sleep and declines at the end, a result that has been later confirmed [1]. Noise level beyond this threshold prevents or interrupts sleep. Tinnitus, as an internal stimulus, is a special phenomenon in sleep research. However, only a few studies about insomnia and tinnitus have been published. Is disturbed sleep that is experienced by tinnitus sufferers a logical consequence of the tinnitus? Then why do only some tinnitus sufferers develop insomnia? And how can it be treated if tinnitus is a permanent sleep-preventing stimulus?
Prevalence

Disturbed sleep is a major problem associated with tinnitus. Hallam reported that disturbed sleep is one of the three most important components of tinnitus complaints, next to difficulties in hearing and emotional stress [2]. The prevalence of disturbed sleep in persons with tinnitus varies from 25 to 77% [3–8]. Tyler and Baker found that 57% of 72 tinnitus patients (TP) experienced difficulties getting to sleep [4]. In a larger sample of 436 TP, 15% reported disturbed sleep [5]. Epidemiologic data are dependent on whether samples are tinnitus sufferers [6] or individuals with tinnitus [7,8], and whether the sample is representative for the population in general [6] or whether it regards a defined subpopulation, for example military personnel [3]. Another reason that different investigators arrive at different values of prevalence may be that the quality of sleep had varied over the time of an investigation.

Insomnia occurs more frequently in individuals with recent onset tinnitus [8]. While 45% of individuals experiencing tinnitus onset less than 1 year reported disturbed sleep, only 26% of individuals who had tinnitus for more than 11 years reported similar sleep issues. Thus, there seems to be an acute and a chronic type of insomnia in individuals with tinnitus. However, there are no published data that support whether these are two distinct forms of insomnia or merely a variation of one form. Nevertheless, the chronic form seems to be more severe since tinnitus and insomnia become more pronounced the longer a patient suffers from these conditions. In a follow-up study in Oregon, 43 of 175 participants still reported having sleep problems after being treated. In this group, loudness and severity were significantly greater [8]. In a follow-up study, after 5 years of treatment in a university clinic in Sweden, 62% TP still reported having sleep problems [9].

The fact that insomnia becomes a persistent problem only in some individuals with tinnitus indicates a special relationship between tinnitus and disturbed sleep. This is considered to be a diagnostic and therapeutic challenge.

Clinical Data of Sleep-Disturbed Tinnitus Patients

Subjective sleep studies of sleep-disturbed tinnitus patients (SDTP) show that there is a higher incidence of problems falling asleep than being awakened by tinnitus. Furthermore, less than half of tinnitus sufferers who had problems falling asleep also reported being awakened by their tinnitus [6]. Results from studies of the relationship between the loudness of tinnitus and sleep are contradictory. Some studies report an influence of loudness on disturbed sleep [7,10] while others do not [11]. Surprisingly, only a minority of tinnitus sufferers with disturbed sleep reported that their tinnitus interferes with their sleep [6]. Studies indicate that SDTP predominantly suffer from delayed sleep onset and that individuals with tinnitus have a tendency to perceive disturbed sleep independent of their tinnitus.

Subjective sleep studies provide only limited information about the effect of tinnitus on sleep because they are based on memory. Individuals with insomnia often underestimate their sleep [12,13]. Only three published studies used polysomnographic techniques to assess the participants’ sleep.

One study shows that individuals with insomnia but no tinnitus have a shorter sleep duration compared to that in healthy controls [14], thus similar to SDTP. Another study [15] compared objective and subjective sleep data as well as clinical data to investigate the hypothesis that SDTP resemble individuals with primary insomnia, daytime vigilance, depression, and daytime tiredness of individuals with primary insomnia and SDTP. No differences in objective sleep measurements were found, but both groups had low sleep efficiencies and long sleep onset latencies. In addition, no differences were found in subjective ratings of daytime sleepiness (ESS) or depression (BDI), both scores being within the normal range. Similar results were obtained in tests of sustained attention performance (QM), which is sensitive to effects of sleep deprivation. These results and those of other studies [7] support the hypothesis that insomnia in TP is not a consequence of depression. Furthermore, polysomnographic1 of tinnitus patients [14–16] measurements reflect subjective disturbed sleep in SDTP. The polysomnographic results in SDTP and individuals with insomnia are similar.

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1 Polysomnography: physiologic measures of sleep which are electroencephalography (EEG), electroencephalography (EOG), and electromyography (EMG).
Since not all tinnitus sufferers develop insomnia, disturbed sleep and tinnitus probably do not have a common somatic mechanism, and it is not clear whether insomnia is a consequence or a comorbid condition to TP. Patients, however, regard the relationship between tinnitus and sleep disturbance to be clear, and many patients believe that their tinnitus is the cause of their disturbed sleep. This mechanism is probably valid in acute tinnitus and is in line with a commonly expected relationship between a disturbing noise and sleep. A higher prevalence of insomnia in persons who suffer from acute tinnitus can be explained with the so-called adjustment insomnia.

The essential characteristic of “adjustment insomnia” is the existence of an identifiable stressor. The insomnia is expected to resolve as soon as the stressor is eliminated. The fact that sleep disturbance becomes less frequent with the duration of tinnitus [8] can be explained by habituation to the tinnitus in some SDTP.

Nevertheless, there is a chronic form of insomnia which cannot be explained with “adjustment insomnia” criteria, where insomnia exists as a comorbid condition. Tinnitus is erroneously held responsible for disturbed sleep by the patient as well as the physician, and insomnia persists without a precipitating factor. Similarities between SDTP and patients with psychophysiological insomnia found in clinical studies support this assumption [15]. Psychophysiological insomnia essentially is a conditioned sleep disorder with a heightened somatic and mental level of arousal that results from a cycle of symptoms such as the urge for sleep, focus on the inability to sleep, and focus on sleep-related cues [17].

There seems to be an overlap between several psychological aspects of tinnitus and features of psychophysiological insomnia. Andersson and Westin proposed classical conditioning, selective attention, and appraisal of tinnitus as mediating factors for the distress experienced by tinnitus sufferers [18]. Similar factors exist in psychophysiological insomnia [19, 20]. Insomnia and tinnitus distress in its chronic form reinforce each other [15]. SDTP perceive tinnitus in the acute phase as a sleep-preventing stimulus. The perspective that it is permanent and inescapable triggers a vicious cycle of insomnia with an obsession on sleep, selective attention to tinnitus, and the inability to sleep in addition to mental and somatic hyperarousal [15]. While the actual sound becomes less prominent after some time, it is rather the attitude that tinnitus prevents sleep that reinforces insomnia. It is therefore understandable that such patients feel their tinnitus is severe [11]. In a study of tinnitus sufferers attending our tinnitus center, we found a clear correlation between tinnitus severity and degree of sleep disturbance (Fig. 65.1).

Restorative sleep is a basic condition of health, and impaired daytime functioning is one of the diagnostic criteria of insomnia (ICSD-2, 2005). An increasing number of studies have indicated that insomnia is a risk factor for physical and mental health disorders [21–23]. Insomnia is associated with impairment of vitality, general health, and a physical ability to function that is not insignificant, even when compared to disorders such as depression or congestive heart failure [22]. Furthermore, studies have shown that insomnia is associated with an increased risk for arterial hypertension [24], coronary heart diseases, psychiatric disorders such as depression [23, 25], accidents, and impaired productiveness on the job [26, 27]. Insomnia itself, therefore, is a major health problem, and disturbed sleep in tinnitus sufferers is a serious medical problem.

Several different treatments, such as pharmacologic agents [28] and cognitive behavior therapy, are available to treat insomnia [29].

Two kinds of pharmacologic agents are in common use: benzodiazepines and sedative antidepressants. Both have problems, especially for long-term use. Benzodiazepine, a GABA_A receptor agonist, such as zolpidem, zopiclone, and zaleplon [30] has limited duration of beneficial effect [31]. While intermittent intake of hypnotics has shown to be effective [32], adaptation (or tolerance) is a problem when used for long periods. Despite that recent studies suggest that an intake of up to 6 months of Z-substances is not harmful [33]; chronically, sleep-disturbed patients
such as SDTP normally need a medication for an even longer period of time.

Especially in the case of chronic insomnia in SDTP, cognitive behavior therapy (CBT) should be applied since it targets the symptoms that modulate distress. Components are bedtime restriction [34], stimulus control, relaxation therapy, and sleep education [35]. Several studies have shown that CBT has a positive and long-lasting effect on primary insomnia [29, 36]. In a follow-up three months after participating in a CBT program specially designed for insomnia and tinnitus, an improvement in sleep as well as tinnitus could be seen.

Alternation of dysfunctional attitudes and behavior patterns leads to patients’ regaining control of pre-sleep ease and as a consequence results in a better likelihood for falling asleep.

**Assessment of Results of Treatment**

The Pittsburgh sleep quality index (PSQI) [37] is the most preferred sleep questionnaire in sleep research so far. It is designed to measure sleep quality, and it also assesses symptoms of sleep-related breathing disorders. The insomnia severity index is a commonly used instrument for measuring severity of insomnia syndrome [38]. Sleep protocols are a useful tool for quantifying therapy effects as well as diagnostic purposes.

**Other Syndromes That Occur Together with Sleep Disturbances**

**Sleep Apnea**

One study [16] showed that sleep apnea syndrome (SAS) occurred in 10 out of 26 SDTP and PLMS in three of these patients. Detecting organic sleep disorders in SDTP is very important since even a mild severity of SAS or PLMS may have an impairing effect on insomnia in SDTP because of its sleep fragmentation effect. Once sleep is disturbed and leads to nocturnal awakening, SDTP have problems falling back asleep. People may attribute nightly awakening to tinnitus and present it to the therapist that way. Studies of the prevalence of sleep-related breathing disorders in individuals with tinnitus have not been published, and there are no data available about the effect of CPAP therapy on the severity of SDTP with SAS. Sleep apnea is not always associated with a high body mass index, excessive snoring, and/or regular alcohol consumption.
Especially in older persons, sleep apnea may occur without these symptoms. Sleep medication such as benzodiazepines and other sedative agents may worsen sleep-related breathing disorders.

Restless Legs Syndrome

Restless legs syndrome (RLS), especially if associated with PLMS, may also have an impairing effect on sleep in SDTP. Periodic limb movement disorder (PLMS) is characterized by episodes of repetitive, stereotyped leg movements during sleep. It may but need not lead to sleep fragmentation. A study found PLMS without PLMS to occur in 15% of individuals with insomnia, and its occurrence increases with age (34% of people over 60 years) [39] (reference needed). Certain sedative drugs, such as benzodiazepines, may worsen insomnia when an SAS or PLMS is present.

Conclusion

Insomnia in tinnitus sufferers is a condition that decreases the quality of life and may be regarded as a serious health problem. It is important to rule out organic sleep disorders such as sleep apnea and RLS as a cause for insomnia. Disturbed sleep in persons with tinnitus should be treated specifically. An early treatment of disturbed sleep is useful to prevent insomnia and its occurrence increases with age (34% of people over 60 years) [39] (reference needed). Certain sedative drugs, such as benzodiazepines, may worsen insomnia when an SAS or PLMS is present.

References

10. Slater R, Jones D, Davis B, Terry M. Project into psychological aspects of adjustment to subjective tinnitus and the effectiveness of tailored masking. London 1983
Chapter 66
Posttraumatic Tinnitus

Dirk De Ridder and Berthold Langguth

Keypoints
1. Posttraumatic tinnitus can be both non-pulsatile and pulsatile.
2. Posttraumatic non-pulsatile tinnitus can result from trauma to the ear or neck. Trauma to the ear includes temporal bone fracture, labyrinthine concussion, ossicular chain disruption, perilymphatic fistula, barotraumas, or noise trauma.
3. Posttraumatic pulsatile tinnitus is related to vascular lesions. Posttraumatic carotid dissection, AV fistula or caroticocavernous fistula, can cause pulsatile tinnitus.
4. Posttraumatic stress disorder can worsen the tinnitus percept and distress.
5. The cause of posttraumatic non-pulsatile and pulsatile tinnitus may be a sign of life-threatening diseases, some of which are treatable.

Keywords Pulsatile • Tinnitus • Trauma • Vascular

Abbreviations
AV Arteriovenous
AVM Arteriovenous malformation
CCF Carotid-cavernous fistula
CD Carotid dissection
CSF Cerebrospinal fluid
MVC Microvascular compression
OChD Ossicular chain disruption
PTSD Posttraumatic stress disorder
TBI Traumatic brain injury

Introduction
Tinnitus often arises after, or is associated with, a trauma to the head – especially to the ear. Mechanical, pressure-related, noise-related, or stress-related trauma can cause tinnitus. Posttraumatic tinnitus can be either non-pulsatile or pulsatile.

Noise-related trauma is the most common cause of tinnitus and hearing loss. It is discussed in detail in Chap. 37.

Head [1] and neck injuries [2] are common causes of tinnitus [3, 4]. 53% of individuals suffering traumatic brain injuries (TBI) develop tinnitus; hyperacusis (intolerance to sudden or loud noise) develops in up to 87% of all TBI cases [4].

Non-pulsatile tinnitus in head injury can be due to injuries to the ear or brain. Injuries to the ear may consist of petrous bone fractures, ossicular chain disruption, and perilymphatic fistulas, as well as barotraumas and noise trauma. Posttraumatic damage to the auditory nerve and brain injuries can cause tinnitus as well. About 10–15% of whiplash injuries develop a whiplash syndrome consisting of persistent tinnitus combined with one or more of the following symptoms: headache, vertigo, instability, nausea, and hearing loss [5] (Table 66.1).

Pulsatile Tinnitus
Posttraumatic pulsatile tinnitus can be the result of a carotid dissection, AV fistula, or caroticocavernous fistula. Traumatic carotid dissections (CD) occur in approximately 1% of all individuals who have had blunt traumatic injury [6]. Pulsatile tinnitus is experienced in 16–27% of carotid dissections at the side of the dissection.
Table 66.1 Causes of posttraumatic tinnitus

1. Non-pulsatile tinnitus
   (a) Ear
      i. Temporal bone fracture
      ii. Labyrinthine concussion
      iii. Ossicular chain disruption
      iv. Perilymphatic fistula
   v. Barotrauma
   vi. Noise trauma
   (b) Nervous System
   (c) Auditory nerve
   (d) Brain injury
   (e) Posttraumatic stress disorder
   (f) Neck
   (i) Neck trauma
   2. Pulsatile tinnitus
      (a) Carotid dissection
      (b) AV fistula
      (c) Carotidocavernous fistula

CD is asymptomatic in less than 10%, whereas more than 90% of individuals with CD develop carotid territory ischemia and/or local signs and symptoms on the side of dissection. Signs and symptoms from the side of dissection include head (65–68%), facial (34–53%), or neck pain (9–26%), Horner syndrome (28–41%), and cranial nerve palsy (8–16%) of the hypoglossal nerve in particular. The facial nerve may also be involved; dysgeusia results mainly from involvement of the chorda tympani (0.5–7.0%) or the glossopharyngeal nerve. A metal-like taste is typical after chorda tympani lesions. Transient pareses of the ocular motor (III, IV, and VI) and the trigeminal nerve have been observed. In ¾ of carotid dissections, an ischemic event occurs, which includes ischemic stroke in 80–84%, transient ischemic attack in 15–16%, amaurosis fugax in 3%, neuropathy in 4%, and retinal infarct in 1% [7].

1Horner syndrome: Ipsilateral myosis, ptosis, and facial anhydrosis; usually unilateral and due to an ipsilateral lesion of the cervical sympathetic chain or its central pathway; an ominous sign when it accompanies an ipsilateral traumatic brachial plexopathy because it usually indicates an avulsion of the C8 and T1 primary roots from the spinal cord. From Stedman’s Electronic Medical Dictionary
2Dysgeusia: Distortion or perversion in the perception of a tastant. An unpleasant perception may occur when a normally pleasant taste is present, or the perception may occur when no tastant is present (gustatory hallucination). From Stedman’s Electronic Medical Dictionary.
3Amaurosis fugax: A transient blindness that may result from a transient ischemia resulting from carotid artery insufficiency or retinal artery embolus, or to centrifugal force (visual blackout in flight). From Stedman’s Electronic Medical Dictionary.

Posttraumatic AV fistulas often result in an audible bruit, thus objective tinnitus or pulsatile tinnitus [8–10]. AV fistulas can develop after days, weeks, or even years [11]. The incidence at the middle meningeal artery in head injuries is 1.8% [9]. However, they can also occur along the superior sagittal sinus [10], the posterior auricular artery (internal jugular vein) [8], vertebral artery-vertebral plexus [11], sigmoid and transverse sinuses [11], or even the scalp [11]. The middle meningeal artery fistula to the sphenoparietal sinus is often the result of linear fractures and are most common in elderly people who have an adherent dura [9]. The sinus fistulas developing after trauma are often the result of a venous thrombosis and are similar to the non-traumatic variant of AV fistulas.

The most common posttraumatic fistula is the carotid-cavernous fistula (CCF). They are divided in the more common high- and rare low-flow fistulas. In 3.8% of traumatic skull base fractures, a traumatic carotid-cavernous fistula is seen, especially in middle fossa fractures, where up to 8.3% develop a CCF [12]. These are characterized by pulsatile tinnitus, pulsating exophthalmia, chemosis, and visual deficit of the afflicted side [11]. Endovascular treatment is the most commonly used treatment.

Non-pulsatile Posttraumatic Tinnitus

Ear

Temporal Bone Fracture

Tinnitus develops in nearly 50% of individuals with temporal bone fractures [13]. The common causes of temporal bone fractures are road accidents, falls, beatings, and gunshot wounds [14–16]. Forty-four percent of temporal bone fractures occur after a motor vehicle accident [15]. Head trauma occurs in 75% of traffic accidents; in 5% of these, petrous bone fracture is noted [17]. Of all head injuries requiring hospitalization,
9% have a skull fracture and 2% of all these have temporal bone fractures [15]. Three kinds of fractures are noted after a substantial trauma to the temporal region: transverse, longitudinal, and mixed fractures. The most common fracture is the longitudinal fracture (82%) [14], characterized by visible laceration and fracture line of the external ear canal. Lateral impact to the head can cause tympanic membrane perforation or blood in the middle-ear cavity with an ossicular chain disruption in about half of the patients [14]. Facial paralysis occurs in 3% [13]. Hearing loss is predominately conductive but may have a sensorineural component as well. CSF leaks occur in 36% of patients with longitudinal temporal bone fractures [13].

Transverse fractures (11%) [14] resulting from antero-posterior impact have a higher rate of sensorineural hearing loss (53%) [18], vertigo [18], and facial paralysis (63%) [14]. Tinnitus develops in 41% of transverse fractures [18]. CSF leaks occur in 25% of patients with transverse fractures [13]. Mixed fractures occur in 7% [14] of the traumata. Some head injuries can severely damage the auditory nerve causing deafness and sometimes tinnitus.

**Labyrinthine Concussion**

Labyrinthine concussion may occur after less serious blows to the head [19], on the side of the trauma or, sometimes, on the opposite side [20]. Tinnitus, dizziness, vertigo, and high-frequency sensorineural hearing loss (4,000–8,000 Hz) are particularly common [21]. In some individuals, onset may be delayed for several days [19], and concomitant conductive hearing loss may occur from disruption of the ossicular chain or from bleeding into the middle-ear cavity [21]. A blow to the mastoid or occiput may damage labyrinth membranes [19] causing the symptoms, as has been suggested by animal experiments [22, 23].

**Ossicular Chain Disruption**

Ossicular chain disruption (OChD) may occur without rupture of the eardrum or temporal bone fracture and result in conductive hearing loss and tinnitus. Traffic accidents are the most common cause of OChD [24]. Twenty-two percent of the OChD are associated with a temporal bone fracture [24], and OChD occurs in 15% of such fractures [25]. There is often a long delay between the injury and treatment (average of 5.7 years) [24]. The most common disruption is the incudostapedial joint followed by the incudomeatal joint [14]. The stapes is most commonly fractured followed by the malleus, with the incus almost never fractured [14]. There is no literature available on tinnitus in posttraumatic ossicular chain disruption specifically, but any hearing loss may cause tinnitus because it activates neural plasticity. Treatment consists of ossicular chain reconstruction if symptoms persist after 2–3 months of recovery or a disappearance of blood in the middle-ear cavity.

**Perilymphatic Fistula**

A perilymphatic fistula results from disruption of the membranes of the labyrinth, most often at the round or oval window [19]. In half of the patients, barotraumas such as blowing the nose, lifting heavy goods, and landing in an airplane are the cause [26]; in about 40% a trauma is the cause [27].

The most prominent symptoms are tinnitus (61–76%), sudden or fluctuating hearing loss (83–93%), vertigo and dizziness (77–91%), and aural fullness (31%) [26, 28]. Subjective positive signs (i.e., vertigo and nystagmus induced by pressure changes in the external ear canal or with coughing or straining [19]) of fistula are present in 71% of these patients [26].

Treatment consists of bed rest while elevating the head, preventing stressful physical activity, and packing both cochlear windows with soft tissue graft [19, 26–28]. If the symptoms persist, a ventriculoperitoneal shunt can be inserted [29, 30]. In general, vestibular symptoms respond to treatment better than auditory symptoms [19, 26–28].

**Barotrauma**

Barotrauma to the ear may occur during rapid change in pressure, such as descent from high altitudes or during underwater diving. It is usually associated with sudden severe ear pain [19]. The cause is that the Eustachian tube fails to equilibrate the pressure in the middle-ear cavity to that of the increasing atmospheric pressures [31]. It causes inward displacement of the tympanic membrane, increased blood flow, and swelling with fluid, sometimes, even blood [19], oozing into the middle-ear cavity, which may lead to hearing loss.
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and tinnitus [32]. In severe situations, it can cause rupture of the eardrum and ossicular chain disruption and even rupture of the round window causing a perilymphatic fistula [31].

Blast injuries are special forms of combined baro- and noise trauma resulting in hearing loss (55–72%) and tinnitus (66–88%) in most individuals exposed to large explosions; both tinnitus and hearing loss occur in 41% [33–35]. Other symptoms include ear pain (41%) and distortion of sounds (28%) [34]. Two-thirds have a perforation of the eardrum, often on both sides (70%) [35]. In seventy-five percent of cases, the perforation heals spontaneously [35, 36].

Treatment of barotraumas is conservative, but when ossicular chain disruption has occurred and a perilymphatic fistula is present, the treatment may be surgical.

Noise Trauma

There is a significant correlation between a history of exposure to noise trauma and the presence of a high-pitched “whistling” tinnitus; the presence of such tinnitus is significantly correlated with high-frequency hearing loss [37] (see also Chap.37). The most commonly observed frequency of tinnitus on pitch matching is the same as the worst frequency for hearing [38], most often at 4,000 Hz [39]. The effect of exposure to noise on hearing loss has been well studied (see Chap.37), but the relationship between noise exposure and tinnitus has been researched to a lesser extent. One study shows that the prevalence of tinnitus in noise-exposed workers is 24% [40], significantly higher than in the general population [41]. It has also been shown that noise-induced hearing loss usually has a steep slope, which is a risk factor for tinnitus prevalence and intensity [42]. Furthermore, the more pronounced the hearing loss is the more discomfort the tinnitus generates [43] and the louder it is perceived [42]. The patients presenting with noise-induced tinnitus are mainly male and on average were 10 years younger than other tinnitus patients suffering from bilateral high-pitched “whistling” tinnitus in correlation with their high-frequency hearing loss [37]. Between 50 and 70% of young people who expose themselves to loud recreational noise have temporarily experienced tinnitus [44]. Disc jockeys develop hearing loss both at high frequencies and at low frequencies and have tinnitus of the same sound spectra [45]. Up to 75% of DJs develop tinnitus [45].

Cervical

Whiplash-Associated Tinnitus

Ten to fifteen percent of individuals who have suffered a whiplash injury develop symptoms such as tinnitus, deafness, and vertigo [5, 46, 47]. It has been hypothesized that tinnitus might develop because of the somatosensory influences on the dorsal cochlear nucleus [48]. (For more information on somatosensory tinnitus, see Chap.43.)

Neuropsychological

Posttraumatic Stress Disorder

Posttraumatic stress disorder (PTSD) is an anxiety disorder caused by exposure to terrifying events. It is often accompanied by tinnitus. The prevalence of tinnitus in individuals with PTSD is much higher than the prevalence in noise-exposed workers (24%) [40]. For example, 50% of Cambodian refugees suffer from tinnitus [49], and the prevalence of PTSD in these individuals is significantly higher than among individuals who do not have tinnitus. Of those patients who seek help for their tinnitus at a veterans tinnitus clinic, 34% have PTSD [50]. It is not known if it is the increased vigilance that causes the tinnitus.

Summary

Tinnitus can be both non-pulsatile and pulsatile after a trauma. Non-pulsatile tinnitus is a common symptom after head and neck injuries, after noise trauma, barotrauma, and in PTSD. Some of the causes involved (e.g., ossicular chain disruption, perilymphatic fistula) can be treated successfully. Posttraumatic pulsatile
Posttraumatic Tinnitus


Chapter 67
Traumatic Brain Injury and Blast Exposures:
Auditory and Vestibular Pathology

Michael E. Hoffer and Carey Balaban

Keywords  Tinnitus • Traumatic brain injury • Tinnitus
• Vestibular disorders • Hearing loss

Abbreviations
mTHB  Mild traumatic brain injury
TBI  Traumatic brain injury

Introduction

Brain injury has been associated with a variety of neurologic sequelae including the auditory symptoms of hearing loss and tinnitus. Traditionally, we think of brain injury as being secondary to head impact and classify the resultant neurologic damage as mild, moderate, or severe [1]. This classification depends on a variety of factors including length of alteration of consciousness, force of the impact, associated injuries, and neuropathology (such as bleeding). This classification is important since it guides management of the injury and gives health care providers some information about the expected pathologies and best practices for management. There has been a great deal of work done over the years on blunt head injury; however, not all brain injury is secondary to blunt head impact.

The most common etiology of injuries in modern warfare is blast exposure. The use of explosives for terrorism has extended this threat to the civilian world. Such as blunt injury, blast exposure can produce traumatic brain injury. This chapter describes differences between blast injury and blunt head injury from a clinical perspective. We will then consider the audiologic sequelae of blast injury, including tinnitus.

Pathophysiological Features of Traumatic Brain Injury

A heuristic diagram for understanding the progression of signs and symptoms of traumatic brain injury is shown in Fig. 67.1.

1. The direct injury to the brain is presumed to be the “textbook” neuropathological hallmarks of concussive brain injury, which include subdural hematoma, cerebral contusion, and subarachnoid hematoma.

2. The subdural hematoma can be delayed, emerging later in subacute or chronic stages after injury [1]. Tissue injury responses include, at the cellular level, cellular repair and metabolic pathways and, at the tissue level, wound healing and vascular regulatory responses. Secondary damage includes ischemia and excitotoxic events that reflect imbalances in homeostatic control of both the intracellular and extracellular environments. Plasticity of intact neuronal pathways can also contribute to recovery. The outcomes (functional recovery and permanent functional loss) will obviously depend upon the severity (and location) of the primary trauma and the efficacy of the biological responses to the primary and secondary damage. The signs and symptoms of a patient at any given time will reflect the interplay
between these dynamic mechanisms. One example of this approach is the growing recognition that subarachnoid hemorrhage can contribute to both early and delayed mechanisms of secondary brain injury, including vasospasm, transient ischemia, oxidative stress, excitotoxicity, cortical spreading depression, microcirculatory dysfunction, and delayed thromboembolism [2–4].

An Introduction to Blast Injury

A shock wave, a blast wind, and an electromagnetic pulse are generated by detonations of explosives. Primary blast injury is defined as the effects of shock wave propagation through tissue. The blast front is a supersonic over-pressure wave, followed immediately by a negative pressure component termed “the under-pressure” [5]. The blast wave produces a positive-negative shift in intracranial pressure that mirrors the incident waveform [6–8]. Unlike primary blunt or acceleration–deceleration brain trauma, low-level blast exposure produces a global compression–decompression of the cranial contents rather than localized brain contusions from impact with the skull. Secondary blast injury is produced by shrapnel or fragments. Tertiary blast injury can produce blunt trauma by impact with objects in the environment. Quaternary blast injury is produced by other detonation products such as heat, electromagnetic pulses, and detonation toxins.

Clinical Contrasts: Neurologic Aspects of Mild Blast Trauma Vs. Mild Blunt Head Trauma

A clinical picture is now emerging from a series of studies conducted with active duty military personnel who sustained a mild traumatic brain injury (mTBI) as a consequence of pure blunt head injury or pure blast head injury. These studies have been presented in detail in other publications, but will be summarized here [9, 10]. The mTBI was defined by the Department of Defense Policy for Mild Traumatic Brain Injury (October 2007) criteria as the presence of a documented head trauma or blast exposure event followed by a change in mental status, which could include nausea, dizziness/balance problems, temporary headache, sensitivity to noise or light, tinnitus, vomiting, fatigue, insomnia/sleep disturbances, drowsiness, blurred vision, memory problems, or poor concentration. One study examined males with purely blunt head injury during service in Iraq (34 individuals) or with purely blast injury during service in Iraq (21 individuals) within 9 months of injury. The clinical characteristics varied markedly between the blunt and blast-exposed patients. Specifically, the blast mTBI group had a much higher prevalence of clinically significant hearing loss (43% vs. 7% of the blunt head injury group) and cognitive impairment (90% vs. 17% of the blunt head injury group). Rotational chair balance test results also suggested a different pattern of functional impairment in the two groups with more unilateral, peripheral vestibular symptoms in the blunt group than the blast group [9, 10]. A second study used dynamic posturography to assess postural control after mTBI. The 33 blunt head injury patients and 39 blast injury patients in this study all received mTBI in Iraq and entered the study within 9 months after injury. There was a significant difference in the sensory organization test results of a portion of the blunt patients as compared to all the blast patients. The group mean scores of the motor
control test of the patients with blast injuries were markedly worse than the group mean score of the patients who had suffered blunt injuries [9, 10]. In summary, our laboratory results demonstrate that the head injury and resultant sequelae seen after blast injuries are markedly different than those seen after blunt head injury. The implication of these findings is that we cannot utilize our decades of knowledge on blunt head injury to predict the pathologies or discern the best management practices in individuals with blast exposure.

**Auditory Pathology after Blast Exposure**

The rate of hearing loss, documented by pure-tone audiometry, increases slightly as a function of time from injury to presentation in individuals with blast exposure and resultant head trauma [9]. Tinnitus was noted initially by from 33% and in 43% of those seen later than 1 month after their most recent blast exposure. However, the occurrence of tinnitus is greater than the occurrence of hearing loss in both groups. Nearly 70% of individuals with documented mild traumatic brain injury report tinnitus in the first 72 h after the blast. This number decreases over time, but the rate of tinnitus exceeds the rate of hearing loss at all time points.

There are many factors that might account for the tinnitus seen after blast injury in our mild traumatic brain injury population. Of course, in many individuals, the tinnitus occurs along with the hearing loss, and the postulated etiology would be from primary damage to the ear. However, as stated earlier, tinnitus is more common than hearing loss, and many individuals who have been exposed to a blast wave have normal pure-tone hearing tests but show abnormalities in hearing noise and in central auditory processing. Most of these individuals complain of tinnitus despite their aforementioned normal audiograms. In this regard, it is critical to note that the subjective tinnitus can be produced by mechanisms that range from localized disturbances in the peripheral auditory system and central nervous system to systemic metabolic disturbances [11, 12]. Particularly, germane to blast TBI is the association of tinnitus with stroke and cerebral hemorrhage [11, 13–16]. Somatic tinnitus can accompany acceleration–deceleration injuries, such as whiplash, in the absence of hearing loss [17]. Blast injury is also associated with a higher than expected rate of posttraumatic Ménière’s disease. Thus, it is quite possible that the central and peripheral sequelae of blast injury produce tinnitus independent of direct ear damage. These factors may contribute to a higher than expected rate of tinnitus and suggest the need for more comprehensive diagnostic tests and a broader range of therapeutic approaches. At the same time, this may allow us to intervene specifically in the primary etiology of the tinnitus and/or more effectively manage the tinnitus after it develops.

**Conclusions**

Ultimately, the tinnitus seen after blast exposure and brain injury is likely multi-factorial and a product of end organ damage, brain injury, and/or a pathology that develops over time. Several factors remain unclear. We have very little data documenting the rate of tinnitus in those with blast exposure who do not have resultant mild traumatic brain injury. Given the rate of blast exposure in current operational settings, this is a very important piece of information. Also, while we have candidate pathologies to account for the tinnitus seen in blast-exposed individuals with mTBI, we still have a great deal of work to do in this area. More targeted and specific tinnitus tests need to be done on this population. We are obligated to better characterize the disorder, so that we can help develop diagnostic and management strategies to initially treat and, in the future, prevent tinnitus associated with blast exposure.

**References**

Part VI
Management of Tinnitus
Keypoints

1. Therapeutic tools include manipulating sensory inputs, modifying psychological influences and a variety of direct approaches to the central nervous system including drugs.
2. A combination of these therapeutic opportunities constitutes a package of care.
3. In this section appropriate experts clarify almost every possible therapy.

Keywords Tinnitus therapy • Package of care

Introduction

For clinicians like me, the following section on management of tinnitus is the first section we look at and the one to which we shall refer most often. We want to know if we can improve the way we implement the therapeutic interventions we already use. We want to know if we should seek to adopt therapies which are already available but are ones we do not use. We want to know about new therapies which are being investigated and may, in the future, help our patients.

The editors are to be congratulated on bringing together such a comprehensive team of authors. Each is recognized as an authority on at least one aspect of tinnitus management. Collectively, they provide a detailed description, analysis, and instructions on almost every aspect of tinnitus management available to help us in the care of our patients.

Earlier sections of this book are also relevant to the management of tinnitus. However, as clinicians, we do not really manage tinnitus. We manage a patient who has tinnitus, and we help that patient to cope with this disorder. Our first contact with a patient, the obtaining of a history, items in a questionnaire, our clinical examination, and the tests performed are all part of our management of that patient. During a well-handled assessment process, anxiety may start to subside. Alternatively, new concerns may be raised. The first contact, even the making of a first appointment, may influence our patient’s concerns and affect the therapeutic outcome. Every component of the assessment process is part of the therapeutic management of each patient troubled by tinnitus. However, this section is focused on the various therapeutic tools we may utilize or recommend following assessment.

Tinnitus is most likely always multi-factorial. However, sometimes, one factor is dominant, and correcting that factor alone may be almost all that our patient requires, as far as their tinnitus management is concerned. Clearing the ear canals of wax or debris, surgical correction of hearing loss, the withdrawal of a drug, or facilitating the treatment of a psychiatric disturbance may stop tinnitus from being a problem. However, even in these “dominant factor” situations, other factors helped determine that awareness of tinnitus became a major feature and influence whether the tinnitus persists as a problem, even after the dominant factor has been treated.

In most of our patients, several factors are important in their awareness of tinnitus and the distress they experience. I find it helpful to group these factors into three broad categories.
• Changes in sensory input. These usually predisposed to the onset of tinnitus and help maintain it.

• Psychological influences. These include emotional state and emotional associations, lack of understanding and resultant anxiety, and unconscious conditioning (“neurophysiological model”).

• Changes in neural activity within the brain. These have usually been triggered by the above factors, sometimes by direct injury, but then become self-perpetuating and are now regarded as the actual “generators” of tinnitus.

I then view the same three broad categories as distinguishing the avenues available for treating each patient who is troubled by tinnitus.

1. Manipulating and where possible normalizing sensory inputs. This applies most often to auditory input where hearing loss may be corrected or compensated, or therapeutic auditory stimulation applied. A lot of attention has recently been focused on somatosensory inputs, their ability to modulate and sometimes trigger tinnitus, and how these effects may be reduced. Visual, olfactory, vestibular, taste, and other sensory inputs may have influences on tinnitus, but have received little study in this context.

2. Controlling emotional factors. Successful management of tinnitus almost always requires reduction in concern about implications and often separation from anger about perceived causes. Disassociation of tinnitus from emotional factors, especially depression, anxiety, fear, and anger is essential. Explanation and understanding reduce anxiety and fear and the tendency for a patient to dwell upon their tinnitus. The most sophisticated and validated approach to achieving this “de-concerning” is cognitive behavioural therapy. At a less conscious level, de-conditioning techniques such as tinnitus-retraining therapy and desensitization with music are useful in reducing physiological changes associated with troublesome tinnitus.

3. Direct approaches to the central nervous system. Once tinnitus has become intrusive and distressing, then treatment through control of sensory input and psychological factors may be insufficient. Neuroplastic changes within the brain may need to be approached directly as well. The most readily available route is through the bloodstream, providing access for drugs and dietary factors. However, changes within the brain can also be approached directly by surgery, by direct electrical stimulation, and, especially in this context, by transcranial magnetic stimulation. Even if such direct approaches can reverse neuroplastic changes, they almost certainly need to be used in conjunction with the control of sensory input and psychological factors if relapse is to be prevented.

We are fortunate that the experts who have contributed to this section of the book have, between them, examined all avenues and clarified almost every possible therapy. Generally, they acknowledge, explicitly or implicitly, that each therapy described needs to be part of a package of care incorporating other approaches if it is to be of long-lasting benefit.

In managing sensory input, we find chapters on auditory training (Chap. 72), sound stimulation and hearing aids (Chap. 74), music treatment (Chap. 75), middle-ear implantable devices (Chap. 76), cochlear implants (Chap. 77), treatment directed to the ear (Chap. 83), and surgical treatments (Chap. 82), all of which may improve or manipulate auditory sensory input. Some pharmacological and nutritional therapies have their effect by improving inner ear function and auditory input (Chaps 78, 79 and 92). Chapters on temporomandibular joint dysfunction (Chap. 95), cutaneous stimulation (Chap. 91), focus on techniques which probably alter somatosensory influences on tinnitus, as may neuro-biofeedback (Chap. 87) and low-level laser therapy (and Chap. 93).

Psychological factors are addressed at a conscious level through chapters on counselling (Chap. 70) and cognitive behavioural treatment (Chap. 71) but may also be important components of sensory stimulation such as in music treatment (Chap. 75). De-conditioning at an unconscious level is inherent in tinnitus retaining therapy (Chap. 73) but may also be important in some forms of sound treatment. There are benefits in more holistic approaches (Chap. 92).

Most of the pharmacological treatments described act directly on the central nervous system (Chap. 78) as may some non-conventional therapies, nutritional factors, and vitamins (Chap. 92). Principals of neuro-modulation are discussed (Chap. 86) prior to descriptions of neuro-biofeedback (Chap. 87) and of direct stimulation both magnetically (Chap. 88) and electrically (Chaps. 89 and 90).
Some specific treatments for particular problems are also described, such as treatment of vestibular schwannoma (Chap. 85) and microvascular decompression (Chap. 84). Chapter 94 is devoted to the similarities between treatment of tinnitus and that of pain. The final chapter is devoted to the methodology of clinical trials for tinnitus. Treatment of disorders that are closely associated with tinnitus such as temporomandibular and masticatory disorders can often relieve tinnitus (Chaps. 95 and 96).

This section is an authoritative description and assessment of almost all the approaches to tinnitus therapy currently in use and others with potential for future benefit. Where a potential therapeutic approach has not been addressed, it is mostly because it has not been reliably reported. There is still room for more innovation.

Our improved understanding of the influences of sensory input and psychological factors and the neuroplastic changes, which result, has given us far greater sophistication in managing our patients who feel distressed by their tinnitus. For each patient, we have to identify the most helpful ways in which sensory inputs can be manipulated, how best to improve understanding and disassociate emotional factors, and whether there is a place for centrally acting agents and other direct approaches. The chapters in this section are a reference library of all the information available to help us make the most appropriate recommendations for each patient in the context of what is available for them.
Chapter 69
The Prevention of Tinnitus and Noise-Induced Hearing Loss

Larry E. Roberts, William Hal Martin, and Daniel J. Bosnyak

Keypoints

1. Although tinnitus is more common in older individuals, it can occur at any age. Because tinnitus in most individuals is associated with hearing impairment, prevalence may be increasing among youthful populations owing to exposure to environmental and recreational sound.

2. At present, there are no effective medical treatments for chronic tinnitus. Because hearing loss is a major risk factor, primary prevention is possible. Primary prevention is effective in other health domains, although it takes time for such programs to have impact.

3. Public education programs, role modeling by parents, cooperation from employers and industry, awareness campaigns, education of health professionals about avoidable risk factors, legislated standards for sound-emitting devices, and protection strategies that are acceptable to the young as well as adults, all have a role to play.

4. “Dangerous Decibels” is an example of a successful program aimed at reducing noise-induced hearing loss and tinnitus among school-aged children and young adults.

5. Epidemiological research tracking the prevalence of hearing loss and tinnitus at all ages, and research on intervention approaches, can provide essential information about effectiveness and long-term trends.

Keywords Tinnitus • Hearing loss • Prevention • Noise exposure • Epidemiology • Dangerous Decibels

Abbreviations

HL Hearing level
TEN Threshold equalizing noise test
DPOAE Distortion product otoacoustic emissions
ABRs Auditory brainstem responses
NIOSH National Institute for Occupational Safety and Health
NIHL Noise induced hearing loss
WHO World Health Organization
OMSI Oregon Museum of Science and Industry

Introduction

It is a common perception that tinnitus is an affliction of older individuals, which is to a significant extent true. Although reported prevalence varies widely among studies (see Chap. 5), it has been estimated that between 8 and 20% of individuals over the age of 60 report a persisting tinnitus, and among these individuals approximately 25% describe their tinnitus “moderate” and another 6.6% as “severe” [1] implying an adverse effect on quality of life in the latter group, which translates into millions of Americans and many more around the globe. However, it is well documented by national surveys [2] and confirmed by clinical experience that persisting tinnitus can occur at any age. Because in most individuals tinnitus is associated with hearing impairment, prevalence may be increasing among youthful populations owing to exposure to environmental and recreational sound in our electronic age.
This state of affairs is by itself sufficient cause for concern among those who formulate public health policy. However, the problem is compounded by the fact that while treatments exist that often can reduce the impact of chronic tinnitus on individual lives, elimination of the disturbing tinnitus sensation itself remains largely beyond the reach of medicine (see Section V of this book). It is especially worrisome that although tinnitus experienced by younger individuals after noise exposure often subsides, tinnitus may return later in life as changes in brain function related to aging unmask a hidden vulnerability. The prospect of a growing cohort in future years calls not only for intensified research into the causes of tinnitus and its treatment but also for programs aimed at its prevention.

Programs and policies aimed at primary prevention have worked in other domains (see Fig. 69.1). In the three decades following publication of the US Surgeon General’s Report on Smoking and Health in 1964, the incidence of smoking (a major preventable cause of respiratory and cardiovascular disease) in the United States declined from 42% of adults in 1964 to 26% in 1998, with this decline being particularly steep among men more of whom smoked (53%) than did women (33%) in 1964, compared to 28% and 23%, respectively, in 1998 [3]. Public education, anti-smoking campaigns, government restrictions on advertising and conditions of use, and litigation have undoubtedly contributed to this outcome, which (although it is not a simple matter to quantify health benefits) is an important medical success story. Use of seat belts in automobiles and helmets for cyclists have also doubtlessly reduced the risk of injury and subsequent social and health costs associated with driving and cycling. These well-known examples illustrate some of the key ingredients of successful prevention. Public awareness is essential, and cooperation from industry (sometimes resisted) is needed. When the need is urgent, government policies, law making, and legal action can mobilize interventions to reduce risk. The personal costs associated with prevention including convenience and expense must be acceptable. Persistence and patience over the long haul are required, and monitoring is needed to gauge effectiveness.

It must be acknowledged that prevention of tinnitus does not have the same urgency as that associated with tobacco use or passenger protection, which are examples that address risks affecting a substantial proportion of the population and if ignored can have catastrophic personal consequences. However, for millions of individuals severely affected, tinnitus is a debilitating and costly condition for which no effective medical treatments are currently available. Tinnitus also shares in common with these examples evidence of a role for a causal and tractable factor that makes prevention of new cases of tinnitus a practical goal. Epidemiological and neuroscience studies indicate that among the many benefits of preserving normal hearing is likely to be the prevention of tinnitus.

**Fig. 69.1** Adult per capita cigarette consumption and major smoking and health events, United States 1900–1990. From the report of the US Surgeon General (2001) *Women and Smoking*
**Tinnitus and Hearing Loss**

One of the highest risk factors for tinnitus is noise exposure. Individuals who regularly worked in loud sound situations or were frequently exposed to impulse noise were nearly three times more likely to have tinnitus than those who did not have regular, loud sound exposures [4]. Henry et al. [5] noted that prolonged sound exposure and noise trauma represented the most commonly known factor associated with the onset of tinnitus. The Oregon Tinnitus Data Registry reported that sound exposures represented the most commonly reported onset factor in a tinnitus clinic population of 2,503 individuals [6]. Tinnitus has also been found to be an early indicator of permanent sensory neural hearing loss in work settings with prolonged loud sound exposure [7]. When measured within individuals, there is a close correspondence between the frequencies that are present in the tinnitus sensation and the sound frequencies at which hearing loss is present in the audiogram [8–11]. The nature of this relation is that ratings of sound frequencies for their similarity to tinnitus increase incrementally at the audiometric edge and continue to increase with the depth of threshold shift up to about 12 kHz [10, 11] (see Chap. 13). Konig et al. [12] reported that tinnitus is associated with steeper slopes of hearing loss, and also noted a strong relationship between the frequency with the steepest slope and the dominant tinnitus pitch for tonal cases. Restoration of hearing is often associated with a decrease in tinnitus, provided that the tinnitus has not been present for too long. It is commonly reported in the clinic and confirmed by systematic study [13] that many individuals with tinnitus experience a reduction of their symptoms when fitted with a hearing aid (see Chap. 74).

However, many people have hearing loss without having tinnitus, and many people who have “normal” hearing according to their audiograms have tinnitus. For example, Barnea et al. [14] found that 8% of their patients suffering from tinnitus had normal hearing thresholds (<25 dB HL) up to 8 kHz, and Roberts et al. [15] reported that 8 of 32 individuals with tinnitus (25%) had normal hearing similarly defined. However, in the latter study, all 32 individuals with tinnitus had hearing thresholds exceeding 25 dB HL when measured above 8 kHz, underscoring the need for more thorough audiometric assessments. In a subsequent study, Roberts et al. [11] measured hearing thresholds up to 16 kHz in two groups of individuals with tinnitus: one consisting of individuals aged 50 years or older (n=40) and the other aged less than 50 years (n=7). As expected, the older group exhibited threshold elevations commencing above about 2 kHz, but the younger group had normal hearing thresholds up to 10 kHz (see Fig. 69.2). However, when the people in these tinnitus groups were compared to age-matched controls without tinnitus, both tinnitus groups had both age groups, even though the audiograms for the younger tinnitus group were in the normal range up to 10 kHz. From Roberts et al. [11]

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**Fig. 69.2** Hearing thresholds in individuals with and without tinnitus, matched for age above (right panel) or below 50 (left panel) years. Hearing thresholds are elevated between 2 and 8 kHz in tinnitus subjects compared to age-matched controls in
hearing thresholds that were elevated by approximately 11 dB compared to controls over the frequency regions corresponding to their tinnitus. These findings suggest that tinnitus and hearing impairment are related and that the degree of impairment needed to increase the risk for tinnitus is not large [16].

An alternative interpretation of the results of Fig. 69.2 is that the threshold elevations seen in the audiograms of individuals with tinnitus do not reflect reduced hearing, but confusion of the test sound with their tinnitus, which overlaps the same frequency range. Measures other than the conventional audiogram provide another approach. Tests for off-frequency listening can indicate the presence of cochlear dead regions that may lead to the development of tinnitus. Weisz et al. [17] administered the Threshold Equalizing Noise (TEN) test for off-frequency listening in individuals with tinnitus who were selected for study because their audiometric thresholds were within the normal range. Evidence was found for circumscribed cochlear damage in the frequency ranges that were rated as being similar to the tinnitus percept. Cochlear dead regions also appear to influence the shape of tinnitus spectra when band-limited noises differing in center frequency are used to measure these spectra, implying that individuals with tinnitus are listening off frequency to sounds in the stimulus where hearing thresholds are better preserved [11]. Measurement of distortion product otoacoustic emissions (DPOAEs) is another approach to detecting changes in hearing. Shiomi et al. [18] found significant decreases in DPOAE amplitudes over limited frequency ranges in 93% of ears in individuals with tinnitus and normal audiograms, in 96% of ears in patients with tinnitus and hearing impairment, and in only 15.4% of control ears. Similarly, Gouveris et al. [19] found decreased amplitudes in the 1,650–2,400 Hz range and increased amplitudes in the 4–6.3 kHz range in tinnitus patients. These studies point to some degree of impairment of outer hair cell function in tinnitus. Studies of auditory brain stem responses (ABRs) have provided more ambiguous results, with some authors reporting shortened wave V latency [20], others prolongations of waves I, III, and V [21, 22], and others no effects on the latency of waves I–V [14, 23].

If it is accepted that hearing loss is a substantial risk factor for tinnitus, how is tinnitus generated when hearing impairment occurs? Neuroscience studies have begun to answer this question (for a review see Chap. 13). Briefly, hearing loss induced by experimental noise trauma in animals leads to a reorganization of tonotopic maps in the primary auditory cortex, as thalamocortical input to the affected region is impaired [24–26]. This reorganization likely occurs because when thalamocortical input is reduced, neurons in the hearing loss region begin to express the frequency tuning of their unaffected neighbors via horizontal connections in the tonotopic map. It has also been found that the spontaneous firing rate of the affected neurons is increased and that there is an increase in neural synchrony (temporally coupled neural activity, sometimes called temporal coherence) in the region of hearing impairment [24]. Evidence from physiological, psychoacoustic, and human brain imaging studies suggests that increased neural synchrony in the hearing loss region may underlie the tinnitus sound [27].

Notwithstanding these lines of research pointing to a role for hearing loss in tinnitus, it is undeniable that there are individuals who have hearing loss but not tinnitus (see the older control group of Fig. 69.2). This is a puzzle to be explained. One factor that might distinguish between individuals with and without tinnitus despite the presence of hearing impairment is a difference in the prevalence of cochlear dead regions in the two groups. To date, this possibility has not been investigated. Age-related changes in intracortical inhibition [28, 29] may also play a role, with lags favoring normal tonotopic structure and conferring a benefit in preventing tinnitus. Some older individuals who have high-frequency hearing loss without tinnitus may eventually come to experience tinnitus, reducing the disparity between the two phenomena. Nevertheless, what protects many elderly individuals with hearing loss from tinnitus is presently unknown.

### Hearing Loss in the Young

Noise exposure, which can lead to hearing loss, is an increasing problem among children. Blair et al. [30] reported that at some time during their young lives, 97% of 273 third graders surveyed had been exposed to sound levels that are regarded to be hazardous to their hearing. Another recent study indicated that 16% of 14- to 18-year-olds listen to their personal stereo systems at levels exceeding the recommendations of
the National Institute for Occupational Safety and Health (NIOSH) on a daily basis [31]. Thirty percent of the students said they sometimes participated in other noisy activities (such as shooting firearms or attending auto races); however, only 5.5% of the students ever used hearing protection while engaged in these activities. Sources of excessive sound exposure for children include loud music [32, 33], real or toy firearms [34], power tools [35, 36], fireworks [37], loud toys [8, 38], and snowmobiles or other loud engines such as jet skis or motorcycles [39]. The World Health Organization reported that North American children “may receive more noise at school than workers from an 8-h work day at a factory”[40]. Surveys of junior high and high school students have identified large deficiencies in their knowledge about normal hearing as well as hearing loss, and that students know little about the damaging effects of noise exposure [41, 42]. Results from the third National Health and Nutrition Examination Survey indicated that 12.5% of 6- to 19-year-olds in the United States (5.2 million) have documented evidence of elevated hearing thresholds directly attributed to noise exposure [43]. Early exposure to noise causes cumulative damage that accelerates age-related changes and long-term consequences [44].

The good news is that nearly all noise-induced hearing loss (NIHL) and related tinnitus can be prevented. Educational interventions can increase knowledge about NIHL issues. One study that evaluated the effectiveness of hearing conservation education in high school students found an average increase of 16% correct responses after participation in an educational program [45]. A second study presented an educational program on hearing conservation to elementary school children and found that their knowledge regarding NIHL improved by an average of 23% [46]. Recent work using resources from the Dangerous Decibels program (see below) has shown that several interventions, including classroom programs, museum exhibits, and online interactivities can improve knowledge, attitudes, and intended behaviors related to sound exposure and use of hearing protection strategies [47–49]. Knowledge of potentially dangerous sounds, their consequences, and simple ways to protect oneself are all significant factors in prevention of NIHL and tinnitus. Public education can promote hearing health and behavior to reduce noise-induced hearing loss, a fully preventable condition.

**Dangerous Decibels**

The health behavior literature has shown that attention to specific components of an intervention affects the success of that intervention. Strategies that tailor messages to the target group [50–53], use interactive not passive instruction [54], and incorporate teaching skills and self-efficacy [52, 53, 55, 56] have been most effective. Dangerous Decibels® is an exemplary program that has been built on health promotion theory applied to hearing loss and tinnitus prevention.

The Dangerous Decibels partnership began in 1999 and has been locally, regionally, nationally, and internationally active in hearing health promotion [48, 57]. The total number of individuals reached by Dangerous Decibels activities, including the museum exhibition at Oregon Museum of Science and Industry (OMSI), classroom education, web-based activities, OMSI Science Festivals at county fairs, and educator training workshops, approaches one million annually. It is the most extensively developed, disseminated, and evaluated hearing loss and tinnitus prevention program in the world with materials in 46 US States and 17 different countries. Between 2001 and 2006, 4,634 elementary and middle school students and adults participated in the formative and summative evaluation process for the Dangerous Decibels interventions. The results showed that the interventions were effective at changing knowledge, attitudes, and behaviors regarding exposure to loud sound and use of appropriate hearing protective strategies [47].

The Dangerous Decibels resources include the following components, some of which are illustrated in Fig. 69.3:

- A permanent Dangerous Decibels exhibition at the OMSI including 12 components covering over 2,000 ft² and providing information to approximately 670,000 visitors each year 70,000 of whom are K-12 students on school group field trips.
- A virtual Dangerous Decibels museum exhibition at the Dangerous Decibels website (www.dangerousdecibels.org).
- An interactive, inquiry-based classroom program targeting kindergarten through 12th grade students covering the physics of sound, normal hearing function, the pathophysiology and functional consequences of noise exposure, and tinnitus and hearing loss protective strategies.
- Educator training workshops that fully equip and certify individuals to present the classroom program in a manner proven to be effective, plus a Teachers Resource Guide with activities, images, and graphics intended to supplement the classroom program.
The “Jolene” system for measuring the sound pressure levels generated by personal music systems through headphones. The Jolene Cookbook [58] describes how students can make their own version of a Jolene.

These and other Dangerous Decibels activities are designed to communicate information about three questions important for the protection of hearing: (1) What are sources of dangerous sounds? (2) What are the consequences of being exposed to dangerous sounds? (3) How can I protect myself from dangerous sounds? Tinnitus is one of the potential consequences, and information about the role of hearing loss in tinnitus is essential to prevention.

Conclusion

Noise-induced hearing loss and tinnitus prevention activities have historically been emphasized in, or perhaps even limited to, occupational and military settings with the assumption that those settings provided the highest risks. However, recent epidemiologic evidence [59] indicates that cumulative hearing loss in the population has not declined over the past 30 years despite expected decreases in NIHL due to mandatory hearing conservation programs in occupational settings, suggesting that sound-related hearing loss may be resulting from exposures in non-occupational settings. Teaching individuals from an early age to cherish and protect the gift of hearing and equipping them to do so provides the highest likelihood of reducing the incidence of tinnitus.

Primary prevention takes time (Fig. 69.1), and education about noise exposure, while fundamental to success, is not the only factor that may bring benefits. Role modeling by parents, cooperation from employers and from industry, public awareness campaigns, education of health professionals about avoidable risk factors, legislated standards for sound-emitting devices, and protection strategies that are acceptable to the young as well as adults, are needed for a successful...
outcome. Epidemiological research into the prevalence of hearing loss and tinnitus at all ages, and research on the effectiveness of intervention approaches, can provide essential information about the magnitude of the problem and long-term trends. In addition to reducing the incidence of tinnitus, other benefits of hearing protection are reductions in health care costs and in disability claims as well as improved social and workplace communication. Primary prevention is especially important for tinnitus, because while some treatments exist that may reduce the impact of tinnitus on individual lives, elimination of the tinnitus sensation itself remains largely beyond the reach of medicine.

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References

Chapter 70
Counseling and Psycho-Education for Tinnitus Management

Grant D. Searchfield, Jane Magnusson, Georgina Shakes, Eberhard Biesinger, and Orianna Kong

Keypoints

1. Tinnitus is a dysfunction of the auditory system that has proven to be highly resistant to a wide variety of treatments (Laryngoscope 109:1202–1211, 1999) making it a difficult condition to treat and to live with (The psychological management of chronic tinnitus: a cognitive-behavioral approach. Allyn & Bacon: Boston, 2001).

2. As there are no easy cures for tinnitus, the tinnitus patient has to adjust to not only the perception of internal noise but also to the often negative beliefs and consequences that accompany it (Psychological aspects of tinnitus, in Contributions to medical psychology. Pergamon: New York, 1984).

3. Some of the difficulties that tinnitus patients encounter include high levels of emotional distress, sleep difficulties, loss of concentration, attention problems, and disruption to their personal, occupational, and social lives (J Speech Hear Disord 48:150–154, 1983).

4. The need to address these “psychological” aspects of tinnitus has been known for many years (Lancet 36:828–829, 1841) but has only recently been given adequate consideration.

5. Fundamentally, the goal of tinnitus treatment is to reduce the negative impact this condition has on the patient’s life. To facilitate this, counseling helps individuals understand their tinnitus, which can reduce the occurrence and level of distress.

6. Providing patients with education about what their tinnitus is, and what it is not, helps to demystify the condition, which can greatly change how they perceive and respond to their tinnitus.

7. This chapter focuses on one counseling approach and provides resource materials that will enable the practitioner to provide support for their patients’ efforts to reduce tinnitus distress.

Keywords Tinnitus • Counseling • Treatment • Attention • Education • Habituation

Abbreviations

ASA Auditory Scene Analysis
CBT Cognitive Behavioral Therapy
CD Compact (audio) Disc
COSI Client Orientated Scale of Improvement
MP3 MPEG-1 (Moving Picture Experts Group) Audio Layer 3
OAEs Otoacoustic Emissions
S.M.A.R.T Specific Measurable, Attainable, Realistic, and Timely

Introduction

Men are disturbed not by things, but by the view which they take of them. (Greek philosopher Epectueus)

To help people cope with their tinnitus and its consequences, counseling is recognized as a vital component of virtually all tinnitus management options [6]. Yet despite the important role that counseling plays, it can be difficult to ensure that this aspect of tinnitus management is undertaken. Nonpsychologists often feel uncomfortable in their role as a patient’s counselor, frequently feeling uncertain as to how far their counseling efforts should
It is therefore the intent of this chapter to clarify the need to provide counseling for tinnitus patients, the role of counseling, and who should deliver this very important component of a tinnitus treatment program.

**What is Counseling?**

For the purpose of the chapter, we define counseling as the process of facilitating change by informing, advising, and empowering individuals who need support. To help patients understand tinnitus and facilitate their coping with the condition, clinical approaches to the management of tinnitus include the use of education, psychological interventions, and counseling approaches. As the term “counseling” has many connotations and is used by many professionals, it is important to be clear what “counseling” refers to, what role it has in the management of tinnitus, and who should be providing the counseling. Just as the term “counseling” can cover a variety of topics, those who undertake counseling can include a wide range of professionals including psychologists, audiologists, counselors, social workers, general physicians, nurses, and medical specialists. In this chapter, we refer to counseling in the broad psycho-educational context that can be provided by any number of health professions.

With regard to its role in tinnitus, it has been said that counseling is the single most important component in the management of tinnitus [6]. Virtually, all treatment strategies incorporate some form of counseling. These treatments include the use of hearing aids [8] tinnitus retraining therapy [9], tinnitus masking [10], and cognitive behavioral therapy [11]. The importance of counseling was emphasized by Tyler [12], who encouraged all sound-based therapies to go hand-in-hand with counseling. The rationale and the form of counseling may differ across treatments [13], but regardless of which strategy is employed, it is necessary to help the patient understand and learn to cope with their tinnitus [6].

**Tinnitus is a Complex Condition:**

**Why is Counseling Needed?**

Tinnitus is the involuntary perception of sound originating in the head (or ears) [14]. Tinnitus is experienced as an occasional slight irritation by the majority of the population [8, 14]. Between 6 and 17% of the population have tinnitus to a significant degree, with 0.5–2% reporting tinnitus that produces sufficient annoyance to interfere with day-to-day activities and quality of life [15–17]. To date, there is no cure for tinnitus. However, there are ways of minimizing the effects of tinnitus on the patient’s life [18]. While today it is accepted that tinnitus can impact the patient’s life in many ways, awareness of the broad-ranging consequences and potential contributors to distress caused by tinnitus was facilitated by studies designed to assess how tinnitus patients experienced this condition. One of the early attempts to investigate the problems experienced by tinnitus patients was undertaken by Tyler and Baker [4], who asked tinnitus sufferers in a self-help group to list the difficulties they experienced as a consequence of their tinnitus. The primary problems reported included negative effects on lifestyle (93%), general health (55.6%), hearing (52.7%), and emotional problems (69.4%). Participants particularly noted difficulties with the persistence of tinnitus (48.6%), and sleep (56.9%) [4]. Further demonstrating the distress that tinnitus can cause the patient, 6.9% of the respondents in Tyler and Baker’s [4] study had considered suicide. The findings of this study (i.e., the potentially negative impact that tinnitus can have on the patient’s life) have been confirmed in other studies, which have reported that severe tinnitus is often associated with depression [19] and, rarely, suicide [20, 21]. Clearly, tinnitus has widespread effects on the lives of those with this condition, which would normally require a multidisciplinary approach to manage it. Consideration must therefore be given to both the physiological aspects of this condition and the psychological factors that can impact the experience of the disorder and, hence, the level of distress it creates for the patient.

When considering how people react to tinnitus, people with tinnitus generally fall within two distinct groups: those who have marked distress or handicap associated with their tinnitus and those who do not [22]. Why this difference occurs between patients is not always clear. For example, vulnerable people exposed to significant stressful events, such as war and accidents, may suffer tinnitus related to posttraumatic stress disorder [23]. Also, personality traits may play a significant role in tinnitus [24]. For those working with tinnitus, it is important to appreciate the influence of factors that can impact on the experience of tinnitus, as these factors can increase the level of distress caused
by the tinnitus as well as the patient’s ability to benefit from treatment (see Fig. 70.1).

Of those who experience distress and disability related to their tinnitus, there is considerable variability regarding the nature and extent of the psychological distress they experience [4, 25]. It is therefore essential that the difficulties experienced by those tinnitus, patients negatively affected by their tinnitus, be carefully assessed in order to determine the factors that may cause and/or maintain their difficulties [2]. When assessing the impact of tinnitus on the patient’s life, it is important to realize that how, and why, a person experiences distress is variable and may not relate to the more “obvious” elements of their condition. For example, it may appear obvious that the loudness of the tinnitus is the factor most likely to influence the degree of distress experienced by a person with tinnitus [2], yet this is not always the case. Several studies have considered features of tinnitus such as loudness and unpleasantness and have found that the loudness of tinnitus (either self-rated or determined by loudness matching) was unrelated to complaint dimensions [26–28]. This highlights the importance of understanding that the perception of tinnitus is only one dimension of tinnitus and it is the psychological dimension that leads to the emergence of tinnitus-related distress [2].

Although tinnitus is a sensory experience, how individuals respond to their tinnitus tends to be more multidimensional, involving their perceptual, attentional, and emotional processes [29]. In describing the impact of psychological factors on tinnitus, Hallam and colleagues [3] proposed a psychological model based on the process of habituation. They suggested that the distress caused by tinnitus is due to an individual’s inability to habituate to the signal, which should occur as it does to any other constant stimulus that does not present as something harmful to the individual [30]. The significance associated to the signal or any arousal-elevating condition can be influenced by the person’s emotional state and/or personality, slowing the natural progression of habituation [3]. For example, if the person is someone who experiences negative thinking, this can overlay all processing of incoming sensations. Such persons may perceive the tinnitus as distressing, harmful, and something that they will be unable to cope with. The importance of understanding how people interpret their situation is eloquently summarized by the Greek philosopher Epictetus’ quote at the beginning of this chapter. It is, after all, the person’s perceived disability that is going to have the greatest impact on their life.

The treatment of tinnitus patients can be further complicated by a considerable delay in patients

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**Fig. 70.1** The effect of counseling on tinnitus annoyance. (a) Distress caused by tinnitus is greatest when a person has high stress levels; mental health issues (anxiety, depression) poor coping strategies and little support. (b) Counseling can break a cycle of distress and provide the patient with the resources to accommodate the tinnitus
seeking medical attention from the onset of their tinnitus. It is not always clear why the person has not sought help for their tinnitus earlier and why their tinnitus has now become distressing [18]. The delay in seeking assistance may be due to people’s developing strategies to distract themselves from their tinnitus to help them cope with the condition [18]. Alternatively it may be that their resources to endure and manage their tinnitus become weakened over time, and as the condition persists, they require assistance in adapting or strengthening their resources. Furthermore, patients often report difficulties in accessing appropriate information and referral to specialist services for tinnitus.

The negative consequences of tinnitus may include emotional states such as depression, anger, and anxiety, resulting in sleep disturbance, concentration difficulties, and interference with personal and social activities [4, 29, 30, 31]. Accordingly, psychological treatments aim to reduce the negative impact of tinnitus; often through the use of cognitive behavioral therapy (CBT see Chap. 21). CBT attempts to address the negative or unhelpful thought patterns and consequential behavioral problems accompanying tinnitus. The therapeutic approach of CBT has been shown to be effective in reducing the negative impact of tinnitus (i.e., distress and tinnitus annoyance) [32, 33] through cognitive restructuring and behavioral modification [34]. While CBT has been shown to be an effective treatment approach for tinnitus, some of the techniques are considered beyond the scope of practice for nonpsychologists; a more general approach is required for those who are working with tinnitus patients, but are not trained in CBT.

**The Role of Counseling in Tinnitus Management**

The goal of tinnitus management is the reduction of either the tinnitus itself or the patient’s perception of the annoyance related to the tinnitus [7]. As reactions to tinnitus and the ability to cope with this condition vary from person to person, tinnitus is a complex condition to treat. Counseling should be the cornerstone of all tinnitus consultations. To facilitate the treatment of tinnitus, the practitioner must work toward establishing as in-depth an assessment of the individual’s complaints as possible, including a thorough tinnitus interview as well as assessment measures. The assessment allows better understanding of the person’s experience of their tinnitus, the impact it has on their life, and their ability to cope [12]. Patient’s perception of their tinnitus, their ability to cope with their tinnitus, their overall level of disability, and their ability to benefit from treatment interventions should be evaluated. As tinnitus can be associated with psychological distress from anxiety and/or depression, it is appropriate that an initial assessment determines their presence. The Beck Depression Inventory [35] and State-Trait Anxiety Inventory [36] have been used to assess baseline states of anxiety and depression. A useful alternative measure of the presence of psychological distress is the Hospital Anxiety and Depression Scale [37], which provides an easy to administer and well-validated measure of psychological distress in physical health conditions and is suggested as a crucial first step in identifying when to refer patients for psychological assessment [38].

The strong relationship between psychological symptoms (e.g., distress, depression, and anxiety) and tinnitus means that psychological approaches have been included in the treatment of tinnitus. There are many approaches to treating complex health conditions such as tinnitus [4]. Which psychological approaches are incorporated into the treatment of tinnitus will reflect a variety of factors, including the resources available to provide the patient and the training of the clinician. Common elements of a management approach include providing education and means to cope with the tinnitus and its effects. How these are provided to the patient varies considerably between programs and practices (again due to resources, practitioner experience, and practicality of program delivery). For those that do not have access to a multidimensional team approach, there is still a great deal that can be offered to the tinnitus patient in terms of tools to help them to understand and cope with their tinnitus.

**Who Should Provide Tinnitus Counseling?**

There is an opinion that interventions involving psychological therapy for tinnitus should include qualified psychologists [39]. It is also argued that an
Counseling and Psycho-Education for Tinnitus Management

Audiologist may be satisfactorily skilled to provide the CBT for patients with problematic tinnitus [34]. There are differences across country borders as to whom and how tinnitus management is provided. It is our opinion that provision of good counseling is important no matter the professional and that all clinicians should be aware of their own limitations and establish appropriate collegial networks. As the role of counseling and its definition will differ across professional groups, the varied background of participants in the counseling process will inevitably require that their expertise be used in different ways. Knowledge and training will determine to some extent the amount of, and style of, counseling. How the counseling is provided will also be determined by the environment (i.e., physical location, resources, type of patients, support networks, and referral options). For example, an otolaryngologist in a small rural center may be required to undertake greater counseling across a broader scope of practice than a clinician based in a large urban hospital working as part of a multidisciplinary team. While it is ideal for tinnitus patients to have access to a professional trained in the psychological management of this condition, it is not always possible or practical, as many practices do not have the resources or funding to provide such treatment. It is, however, possible to provide tinnitus patients with effective approaches to manage their condition, as audiologists or other tinnitus specialists can provide professional counseling [18] by familiarizing themselves with general counseling skills and principles (good basic texts exist for this purpose [7]).

Due to the chronic and distressing nature of the condition, tinnitus patients require engagement at a greater level than many other otologic or audiologic problems; as a consequence, clinicians should be prepared for an ongoing relationship with the patient. It is important that the professional be knowledgeable in their area of specialty, be sympathetic and caring for the patient, and demonstrate an understanding of the patient’s problem [13]. The professional also needs to provide a clear therapy plan and express their belief in the chosen treatment [13]. Clear communication processes need to be established about when and how patients can contact the clinician (e.g., email, telephone, consultation). It is important not to foster dependence on the clinician but still maintain an easy means for the parties to communicate.

Counseling Approaches for the Management of Tinnitus

As a chronic condition, a primary focus in counseling and psychological approaches to the management of tinnitus is to reduce the distress caused by the tinnitus and the impact the condition has on the person’s life. That is to say that tinnitus is a persistent condition with no easy cure and the focus of interventions therefore are to alter any negative thoughts the person has about the condition and its impact on their life, as this will decrease the role that tinnitus plays in their life.

Counseling interventions can range from simply providing information [29, 40] or educational sessions [41, 42] to psychologically influenced techniques such as relaxation training (e.g., [43]), attention control training (e.g., [44]), and sleep hygiene (e.g., [45]). Some counseling-based therapies include sound therapy as important elements. These approaches include masking and partial masking [2], tinnitus retraining therapy [46], tinnitus activities treatment [47], and audiological tinnitus management [48]. Counseling is the critical component in these therapies [49].

The Psycho-Educational Approach

Psycho-education is a patient-focused approach based on the premise that the more knowledgeable the patients are about their condition, the better the therapeutic outcome [50]. Readers are referred to Lukens and McFarlane [50] for a review of the effectiveness of psycho-education in health care. Providing information is considered by many to be a critical part of tinnitus management [29, 40, 51]. It has been suggested that an educational approach be the first step in tinnitus treatment before additional intervention is ventured into [39]. This helps with correcting the maladaptive thoughts and behaviors that can develop from false beliefs about tinnitus, which would be counterproductive to any accompanying management strategy [13]. Educating the patient about tinnitus and peoples’ responses to this condition enables both the patient and the clinician to explore the problem and clarify the purpose and expected outcomes of subsequent interventions [41]. During the education sessions, it is
important that there is opportunity for sufficient feedback and participation by the patient, as this will allow them to express any uncertainties and demonstrate any problematic patterns of thinking that could be a barrier to the success of any treatments offered (e.g., negative thinking about possible sinister causes of the tinnitus).

**Counseling Content and Context**

Tinnitus treatments use either group or individual sessions, but sometimes both have been applied. The integration of both contact styles has been effective in tinnitus management [52]. From a clinician’s perspective, group therapy is a more cost- and time-effective method; it allows for the presentation of information to more patients in less time [52]. Individuals in a group may be role models to each other, which helps with the realization that there are others in similar situations [52, 53]. Another benefit of the group educational approach is its abilities to attract those who are not drawn to counseling, per se, due to the stigma and uncertainty attached to nonmedical or psychological approaches [54]. However, a disadvantage of the group session is the lack of an one-on-one relationship between the patient and the clinician [52]. Also, unless sessions are well managed, outgoing individuals may dominate discussions to the detriment of more reserved participants. Additionally, within the group format, the observation of another group member’s success might evoke envy or confirm the uniqueness and difficulty of one’s problem [55], making the person feel more distressed. In contrast to group therapy, individual sessions allow for specific issues pertaining to each individual patient to be addressed, which might be necessary for some. The decision to provide group or individual counseling depends on factors such as the availability of groups (this is not always feasible for some practices) and the patient’s preference [56].

With regard to the content of the counseling, in an individual setting, it should be adjusted to suit each individual because a patient’s lack of understanding will be a barrier, thereby defeating therapeutic interventions [8]. In a group setting, the content should be broadly based to encompass the essential elements applicable to most patients. It has been suggested that successful counseling programs include: the capability to change the way patients think about tinnitus; the ability to alter their behavioral or emotional reactions toward tinnitus; and an understanding of each patient’s needs [8]. Shorter term counseling interventions have become increasingly favorable in a variety of clinical settings and are usually designed to be part of an overall management plan [42]. Topics usually covered include: the hearing system and hearing loss, the epidemiology and causes of tinnitus, perception (including habituation and attention), and treatment options [13]. In the following sections, we will briefly outline the main contents of one approach to counseling and the rationale for using them. Effective counseling on this basis requires that the clinician has good working knowledge of the physiology of the auditory system, as well as the mechanism and management of tinnitus and be able to convey this information in layman’s terms to de-medicalize the condition.

**Counseling Topics**

Although for presentation purposes, the topics are presented here in a linear fashion (2 follows 1, etc.), the person providing counseling should be prepared to take a very nonlinear approach – the clinician should guide and react to patient responses, rather than follow a script. The elements that the authors believe are important to convey to the patient are:

1. Needs and goal setting
2. Anatomy/neurophysiology of the ear
3. Results of audiological assessment
4. Perception of sound and tinnitus
5. Habituation
6. Attention
7. Treatment approaches
8. Self-management/coping strategies
9. Referral
10. Relapse prevention
11. Hyperacusis
12. Homework

**Needs and Goal Setting**

A technique commonly used in counseling of chronic conditions such as tinnitus is goal setting, as it is an
important skill to help patients work toward, achieve, and maintain treatment success [12]. Research has shown the importance of goals in improving self-efficacy and performance; it has been reported that the enthusiasm to match performance to goals derives from an anticipated increase in self-satisfaction [57]. Goal setting is said to positively impact an individual’s performance through a self-regulatory process. These processes include enabling the individual to focus their attention, promote effort, and initiate task-related strategies [58]. While there are many ways to set goals, one of the most successful methods is setting S.M.A.R.T goals, which require the person to make their goals specific, measurable, achievable, realistic, and time bound [59]. The clinician’s role is to help patients identify the areas they want to change and then guide them in ways to achieve these goals.

The purpose of using a goal setting technique is to help the patient focus on ways to move themselves forward, thereby reducing their focus on the negative and distressing aspects of their tinnitus. The use of the motivational effects of goal setting in the acquisition of new skills has been demonstrated in various fields (e.g., [58, 60]). It is necessary to ensure that the goals are adequately difficult to motivate, but not so difficult to discourage an individual from achieving them [61–63]. Several studies [57, 61] also emphasize the need to have explicit performance levels, including concrete and quantifiable outcomes. Regardless of the nature of an assignment, a person is usually advised to set short-term goals, as this increases their motivation and expectations toward the task at hand [64].

An important aspect of helping patients’ progress through treatment is determining their needs (i.e., what they want/expect from treatment). Many audiologists will be familiar with the Client Orientated Scale of Improvement (COSI, [65]). This tool is used to determine specific hearing needs and the extent to which they are achieved following the fitting of hearing aids. A slight modification of this scale can also be applied to help determine needs and set goals for tinnitus management [8]. Using the Client Orientated Scale of Improvement in Tinnitus (COSIT), the clinician and patient identify specific situations in which tinnitus is bothersome (e.g., “Tinnitus affects my ability to concentrate at work”) and discuss ways of reducing tinnitus in these situations (e.g., “Amplify sound to reduce tinnitus audibility”). At stages throughout the tinnitus rehabilitation process, the problems identified using the COSIT are re-examined and in each situation, the degree of tinnitus improvement is determined. If improvement is not shown, appropriate steps (change in strategy, different techniques, or referral) are undertaken to address the problem until realistic goals are achieved.

Anatomy/Neurophysiology of the Ear

Counseling based on neurophysiology will commonly attempt to explain, in some detail, the normal and abnormal physiology of the auditory system and related neural networks. In so doing, the aims are to provide knowledge of the processes occurring in the generation of tinnitus and eliminate unfounded fears or presumptions as to the underlying causes [46]. It is also vital that misconceptions are corrected and patients are given sufficient reassurance that tinnitus is not a life-threatening injury or a psychiatric disease.

The elements of anatomy/physiology of the ear thought to be important for discussion (using diagrams and scripts similar to Appendix 1 as a starting point) are outlined below. It is important to pitch the amount of detail to the perceived level of the patient’s understanding. Starting simple is best, but allow the patient to guide you as to the depth of their understanding through questions and answers. It is better that a patient leaves their consultation with a firm grasp of basic concepts, than a collection of confusing neuroanatomical nomenclature. Examples:

1. Outer and middle ear are responsible for conduction and amplification of sounds to the inner ear. The Eustachian tube as a source of repetitive sounds during swallowing. We demonstrate and say to patients:

   Although swallowing is louder than your tinnitus, it is not perceived. The brain is able to filter sound, when it has no “importance.”

2. Inner ear: hair cells, possible pathologies (e.g., noise trauma, ototoxicity) (Fig. 70.2).

3. Nerve ruling out possibility of acoustic neuroma (assuming investigation has been undertaken).

4. Brainstem: reaction to sound when detecting danger and provoking a strong and subconscious reaction. Tinnitus as a new signal to the brain creating arousal, fear, and threat-related reaction.
We explain to patients:

Our reactions to tinnitus are a consequence of hearing a new and unknown annoying sound; they are not signs of mental illness.

5. Midbrain and cortex: addition of emotions and complex association of the sound to templates of normal sounds and depending on the subconscious evaluation we may focus even more on the tinnitus sound.

We say to patients:

Tinnitus sound can be “stored” and become longer lasting the more you are focused on it.

and

It is thought that tinnitus becomes magnified because of how the brain analyses tinnitus and how we think about it. Tinnitus happens because the brain misunderstands information from the inner ear. The inner ear sends nerves (think of these as wires in an electrical circuit) to information centers in the brain. When damage occurs to a specific region of the ear, there is less activity from the ear; the brain reacts to this over time creating new activity.

and

People react to tinnitus in different ways. Tinnitus usually begins following ear injury, even small amounts of damage can start tinnitus (we relate the patient’s tinnitus to audiometry and discuss different measures such as otoacoustic emissions (OAEs)). But the parts of the brain involved in hearing and emotion are also involved downstream from the ear. Most of the “wiring” of the auditory system is involved in the development and appearance of tinnitus itself (relate to Fig. 70.3 and describe the nonauditory centers, explaining that the limbic and autonomic nervous systems are primarily responsible to a large degree for tinnitus annoyance).

**Results of Audiological Assessment**

The first step in the evaluation of tinnitus, and then its management, is a comprehensive case history including questions of onset, description of the tinnitus “sound,” location, possible cause (noise, medications, stress), and severity (Chaps. 46 and 47). If the tinnitus is objective, pulsatile, unilateral, or associated with a tempromandibular joint complaint, referral to an otolaryngologist or other specialist is recommended to the patient (Chaps. 48 and 50). We explain that the underlying cause in these cases may possibly be medically treatable, we are careful not to build expectations of a cure, nor are we pessimistic as to the potential for an effective intervention. We also explain that while there is currently no objective measure of tinnitus, psychoacoustical assessments of tinnitus qualities (pitch and loudness) (Chap. 49) and psychometric evaluations (Chap. 54) of tinnitus severity are often used by clinicians to characterize tinnitus.
Perception of Sound and Tinnitus

Tinnitus does not obey the normal rules which apply to sound perception [66]. For example, tinnitus intensity matches are out of step with its perceived loudness – tinnitus may subjectively match to a quiet external sound but be perceived by the sufferer as being extremely loud, e.g., “as loud as a train.” Although tinnitus may have a low-intensity match, it can be difficult to mask – even when using high-intensity frequency-matched sounds (Chap. 49). Also, tinnitus does not have an external source or object to relate to. One reason for the annoyance and “strangeness” of tinnitus could be its conflict with normal Auditory Scene Analysis (ASA) [66, 67] (Fig. 70.4). Using Fig. 70.5a, we address the perception principle that describes the mind’s tendency to seek figure and ground distinctions (e.g., Rubin’s figure ground vase) and how the brain extracts important features. We also use visual analogs to explain phantom perceptions (such as lateral inhibition, Fig. 70.5b).

We say to patients:

In our daily activities we are able to listen to one sound of interest, such as a friend’s voice buried in a background of competing noise. To do this we must categorize sound features occurring simultaneously (e.g., pitch and loudness) to the correct source. Tinnitus disobeys rules we would normally apply when listening to real sounds.

Feldmann [68] eloquently described that the natural reaction of people to tinnitus onset is to search for it and place it in context of a sound in the environment. With true sounds, we can localize them to something we can see, touch, and sometimes even smell. Multisensory recognition of objects is normal, tinnitus lacks this sense of reality, making it difficult to ignore.

We say to patients:

One of the reasons tinnitus is so annoying is that we hear it, but can’t see it or find where it is coming from. Imagine for a moment that tinnitus comes from this pen instead of your ears. If it was from the pen it would appear real, and be easier to ignore. It is natural for us to want to find and identify the source of sounds, when we can’t it becomes frustrating (e.g., finding the source of a dripping sound in the house, is it a water leak?)
The above examples help to relate the sufferer’s experiences within a simple philosophical framework that can be adjusted to suit the patient and the therapeutic approaches described in the following sections.

### Habituation

A decline in behavioral responses to a sound signal due to repeated exposure is known as auditory habituation [69]. It appears that habituation is not caused just by the repetition of the sound but by the meaning or association the stimulus holds in the particular situation [69]. A lack of habituation was possibly first postulated by Hallam et al. [3] to play an important role in tinnitus persistence and annoyance. Habituation has become a common feature of most counseling and sound therapy practice [46, 47].

We say:

If a person moves from the country to the city often they become annoyed by the noise of city traffic. Sometimes the noise keeps them awake and is a great irritation. Usually this annoyance reduces and the person becomes less and less aware of the city noise with time. They – automatically – learn to ignore the noise, as it is not an important sound. The sound becomes classified as unimportant by the brain. It is as if the sound is no longer there. The same thing can happen with your tinnitus, we need to find ways to help your hearing system treat the tinnitus as an unimportant background “sound.”

### Attention

Attention may play a large role in tinnitus annoyance and should be addressed in counseling [12]. Tinnitus can often become the main focus in a person’s life, consuming their attention resources and ability to
concentrate in other tasks. Tinnitus can become the dominant element in a person’s awareness. The acquisition of attention control skills, such as distraction, allows a person to shift their attention to and from tinnitus during stressful situations [13, 43]. These techniques may provide the individual with some sense of control over their tinnitus and the related distressing experiences [29, 44]. Apparently, it is the assumed uncontrollability of the tinnitus sensation which plays a key role in tinnitus being aversively interpreted [27, 68].

Attention control techniques aim to help listeners learn strategies to switch focus of attention from one thing to another, so that attention can be brought under voluntary control to direct thought to and from one’s tinnitus. Henry and Wilson [2] suggest that by exerting control over attention, tinnitus-related distress will be reduced. Their technique can be used to alternate attention from tinnitus to other sounds and is consistent with the process of ASA discussed with patients:

We say:

We need to teach your hearing system to pay less attention to the unnatural sound of tinnitus and instead listen to more other “real” sounds. Use your ears like a search light – listen for sounds around you, what do you hear? Where is it coming from? Can you tell me more about the sound? When you were listening for the sound – were you aware of the tinnitus – possibly not as the other sound was competing for attention against the tinnitus, we can’t hear everything around us all at once, we must pick and choose. Let’s practice and

Focus your awareness on the noises in your head – tune into the noises. What can you hear? Now quickly redirect your attention. Focus on external noises in the room and outside…notice you can only focus on one thing at a time.

(see Henry and Wilson page 106 [2] for complete dialog)

**Treatment Approaches**

The different treatment types (Chaps. 70–94 in this volume) can be discussed in layperson’s terms to facilitate the patient’s understanding of their tinnitus and the role that different treatment approaches can play in
the management process. We provide the following simple information to patients about treatments.

**Hearing Aids**

It is likely that tinnitus is the response of the hearing system to altered output of the inner ear following hearing loss. One treatment approach is to identify any underlying reasons for the hearing system being overly active and to interfere with how the brain analyses the tinnitus. When hearing loss accompanies tinnitus, this would involve the fitting of hearing aids to the injured ears in an attempt to normalize activity. There are a number of ways the fitting of appropriate hearing aids can help in reducing tinnitus [8]:

- Psychological benefit from reducing hearing handicap
- By reducing the attention being paid to hearing and consequently tinnitus
- Modification of neural networks responsible for tinnitus
- Amplified sound can partially mask the tinnitus

**Masking**

Masking is the process of covering, usually partially, the tinnitus with an external sound. The sound used does not appear to be crucial, but should be less bothersome than the tinnitus. Masking often allows the tinnitus sufferer to gain control over their tinnitus by determining when they do not wish to hear it. Long-term use of partial masking, along with counseling, may lead to tinnitus habituation. Some idea of the potential benefit of masking can be assessed in the clinic by listening to an assortment of sounds over headphones.

**Habituation Therapy**

If the patient has no reason to attend to tinnitus, they should get used (habituate) to it. Even loud sounds can be habituated to if they are nonthreatening, for example, people living near railroad tracks seem unaware of the sound of trains passing. The difference between a person who experiences tinnitus and one who “suffers” from it may be the person’s ability to habituate to the tinnitus. Habituation therapies sometimes combine sound therapy with counseling. Hearing aids, broadband noise generators, and devices combining both amplification and generation of sound (combination aids) are used to reduce tinnitus audibility to facilitate the habituation of tinnitus. The sound therapy is thought to help by allowing the patient to become used to tinnitus as the sound fades into the background.

**Cognitive Behavioral Therapy (CBT)**

Psychologists often use CBT. They teach strategies and techniques to enable patients to cope with tinnitus. This therapy enables sufferers to change the way they think about their tinnitus. By minimizing the impact of unhelpful or negative thoughts about tinnitus, through challenging and changing responses, tinnitus annoyance can be reduced.

**Self-Management/Coping Strategies**

It has been shown that a person vulnerable to stress is more likely to experience tinnitus distress, whereas a more stress-tolerant or resilient person might be able to handle a greater degree of tinnitus before seeking help [70]. Although tinnitus and its associated symptoms can be a frequent source of stress and distress, stress in return can often exacerbate the existing effects of tinnitus. Therefore, managing stress and learning to relax helps reduce the effects of tinnitus and prevent further aggravation. As a stress-reduction technique, relaxation training enables an individual to become calmer and less reactive, hopefully reducing tinnitus perception [40]. As a first step, relaxation exercises such as progressive muscular relaxation and abdominal breathing [2] are potentially helpful. Patients can be informed about relaxation and be provided with resources for undertaking it (Appendix 1).

**Improving Sleep**

A very common complaint amongst tinnitus sufferers is difficulty in sleeping [4, 12, 15, 27]. Sleep problems may include regularly waking during the night and difficulty in falling asleep [72]. It is possible that tinnitus
seems louder and more noticeable at bedtime due to the decrease in ambient noise at night [72]. Improving the quality and ability of individuals to sleep may reduce the adverse effects of tinnitus.

Sleep hygiene is a treatment tool for insomnia which involves behavioral practices that promote good sleep [45]. Patients should be asked about their bedtime routines and sleep patterns. There are different treatment versions of sleep hygiene [73, 74]. However, they all generally involve learning about sleep scheduling, attitudes and feelings that affect sleep, appropriate prebedtime activities, maintaining a good sleep environment, and the importance of daytime behavior [74]. Caffeine and nicotine both have stimulating effects, and intake should thus be regulated [53]. Regular exercise promotes sleep, but should not be carried out close to bedtime [53]. The use of sleep hygiene alone has produced variable success; however, when applied with other forms of intervention (for instance, relaxation exercises or cognitive behavioral treatment) greater improvement in tinnitus symptoms has been observed [75].

**Music**

One easily implemented self-help sound therapy measure is the use of low-level music played in the background in quiet situations to draw attention away from the tinnitus.

For best results, the following has been recommended to patients [76]:

- Listening to music or background sound that induces positive feelings.
- Music without vocals.
- Music without pronounced bass beat.
- Music should be pleasant but not too interesting.
- For short-term relief, when tinnitus is severe, attention capturing music can be beneficial.
- For long-term tinnitus, habituation music which induces relaxation while reducing tinnitus audibility.
- Music should be played at a low level, ideally where the music blends with the tinnitus.

Extra stimulation could be provided at night by a bedside sound generator or compact discs (CDs) designed to interfere with tinnitus detection [77]. This could involve the use of a pillow speaker in combination with a pre-existing CD or MP3 player. If these are not available, purchase of a purpose-built bedside sound generator or tinnitus reduction CD could be considered (the clinic could have these available or a source for clients to obtain them).

**Referring to Clinical Psychology**

On the basis of cost-effectiveness, it is proposed that combining education and self-help advice should produce significant tinnitus reducing benefits when used as routine treatment. Those patients not profiting from the use of the minimal-contact approach could be offered CBT [78] or another psychological intervention. A common reason why tinnitus becomes stressful and disabling relates to the persons’ perception of the auditory stimuli in terms of what could be causing the sensation and their ability to cope with it. Psychological therapy is therefore an appropriate treatment approach for tinnitus, as psychological techniques, including CBT, aim to change how a person thinks about something that will then impact how they react to that situation, stimulus, or event. Within a tinnitus management program, the intent is therefore to change how the person perceives and responds to their tinnitus, so that they are not as negatively impacted by the condition.

**Relapse Prevention**

Once made aware of potential triggers and means to manage them, the patient should be able to identify signs that a previously compensated tinnitus may re-emerge. New stressors, anxieties, and life events could retrigger tinnitus onset. Reassurance that the re-emerged tinnitus is likely a consequence of these events and that management of these issues should again reduce the salience of tinnitus is important. Changes in hearing could also trigger a resumption of tinnitus. As noise is the primary cause of tinnitus-related ear injury, it warrants attention. Hearing conservation should be addressed with caution. Care should be taken to distinguish damaging from helpful sound. An over emphasis on hearing protection may lead to an auditory deprivation effect – potentially reactive plasticity and tinnitus [79] and hyperacusis.
At the same time, patients should be made aware of dangerous sounds and how to avoid further injury [80]. One method to avoid resumption of annoying tinnitus is for the individual to be equipped to manage any re-emergence. Written materials to refer to can be useful, “The Consumer Handbook of Tinnitus” [81] and “Tinnitus. A Self-Management Guide for the Ringing in Your Ears” [82] being examples. Another way is to have access to homework tools that can be used independent of the clinician.

**Hyperacusis**

Although this chapter focuses on counseling for tinnitus, the underlying counseling principles can be applied to other symptoms of auditory injury such as hyperacusis. Care should be taken to explain the concept of hypersensitized auditory pathways, hearing protection vs. hearing isolation, and the importance of sound exposure to achieve some degree of normal tolerance to sound (Chap. 3).

**Homework**

Homework permits the practitioner to use the time between sessions effectively by engaging the patient in tasks aimed toward the therapy goals [83]. Although homework assignments are not commonly applied as part of a tinnitus intervention plan, research on homework in other disorders has demonstrated improved treatment outcomes [84]. One of the primary benefits of using homework is that the techniques learned during the intervention are practiced outside the session [83]. CBT [11, 29], cognitive therapies [85, 86], and rational-emotive therapy [87] have all incorporated homework as part of their therapy plans. Anxiety disorders [88] and certain phobias [89] are examples of clinical conditions that have been aided by homework activities. As part of a SMART approach, homework need not be complex or onerous, but it should encourage participation and ownership of the problem by the patient.

The benefits of the inclusion of homework assignments into therapies can be seen through such effects as significantly improving treatment outcomes [84] and modifying behavior without supervision of a clinician [90]. It is important to note, however, that the benefits a person gains from the homework assigned depends on the clarity of its description and rationale, as well as the degree of patient involvement and level of compliance [84]. Kong [56] investigated the effectiveness of two CBT-based homework exercises alongside group-based information sessions to manage tinnitus. Simple, stand-alone take-home tasks were specifically designed, so that they could be provided to participants without needing to have a psychologist involved in their delivery. Two experimental groups, ACTIVE and PASSIVE, received identical educational sessions, once a week, for five consecutive weeks. All participants were given information sheets with general instructions to carry out the homework tasks that were meant to help in areas of difficulty caused by tinnitus. Additionally, the ACTIVE group participants received detailed and specific assignments to complete during the week. The majority of participants tended to benefit from the participant education sessions. A slightly greater reduction in tinnitus effect was recorded for the ACTIVE group participants at the end of this study when compared to the participants in the PASSIVE group. It was concluded that group-based information sessions including specific “active” homework assignments have the potential to be used alongside audiological management to reduce tinnitus impact [56]. The decision for the weekly topics was based on areas of difficulty frequently experienced by people with tinnitus, which were identified in previous research [4, 11, 30]. Kong [56] compiled the self-help strategies presented in Appendix 1 from various psychological management publications (e.g., [29, 91, 92]).

**Resources**

We the approach, we recommend that it is useful for the clinician to have charts of the ear and the central nervous system, and cartoons/schematic diagrams of perceptual principles available. In this chapter, we have provided some examples that we, as clinicians, have found useful. Pubmed is a great source for up-to-date information on tinnitus, while hearing aid manufacturers often have excellent anatomy charts. In addition, clinicians have offered helpful counseling tools in

Summary

Education alone can be a sufficient intervention for some patients [43, 78, 93]. Due to the complexity and multiple factors which impact upon the emotional well-being of an individual, a multidisciplinary team approach is best when treating a patient with complex tinnitus [52, 94]. However, circumstances will determine which professionals will be providing counseling within different settings. We have suggested a method that works in our clinic practices. We have found the education approach a very useful counseling method to empower patients to de-attend and habituate to tinnitus. Clinicians will have their own counseling “tricks” and methods, but the essence of approaches will likely be similar, to make the patient feel less fearful of their tinnitus and provide tools to minimize and, hopefully, eradicate any negative effects from the perception of tinnitus.

Acknowledgments  Our thanks to Dr David Munoz who drew the image of the organ of Corti (Fig. 70.2b) and Kim Wise who used her clinical experience of counseling tinnitus to critique the text.

Appendix 1: Homework

The following homework exercises were compiled and trialed by Kong [56] on the basis of several previous studies [including: 2, 27, 92]. They are presented here in a format for clinicians to provide to patients.

Topics for Take-Home Tasks

1. Goal-setting
   S.M.A.R.T goal-setting strategy

2. Sleep hygiene
   Going to bed strategies
   Falling asleep strategies
   Sleeping environment
   Daytime habits

3. Relaxation techniques
   Progressive muscle relaxation
   Deep breathing exercises

4. Attention control
   Attention control techniques
   Distraction

5. Communication strategies (when tinnitus accompanies hearing loss)
   Communication tips

Task 1 Goal Setting

Background for Clinician

Tinnitus may result in a withdrawal from work and social activities that might normally provide a sense of achievement and enjoyment. The loss of these positive feelings, along with isolation, may lead to the person strongly attending to their tinnitus. Goal setting is about identifying and then overcoming barriers to participation and activity.

For Person with Tinnitus

Goal setting is the process of determining what your goals are, and making plans to achieve them. The goal-setting strategy explained here is known by the acronym S.M.A.R.T. This strategy has five components:

Specific: Goals need to be Specific in order to make reaching them easier. Specific goals have a much greater chance of being accomplished than do broad and general ones.

For example, a specific goal, “to be able to read without becoming annoyed by tinnitus,” is easier to reach, make a plan for, measure progress, and to know when it is achieved than a general goal of, “I want the tinnitus to be gone.”
Answering these questions can help to ensure a goal is a specific one:

Who is involved? What do I want to accomplish? Where am I going to do this? When will this occur? Why do I want to accomplish this goal?

Measurable: Goals need to be measurable. This way you will be able to see the progress you are making, will know when your goal is reached, and will know when it is time to celebrate! Celebrating your success is an important part of goal setting.

To determine if your goal is measurable, ask questions such as:

How much? How often? How will I know that I have reached my goal?

How will I know that I am making progress towards my goal?

For an example of a measurable goal, let us say your goal is to read in the evening without becoming annoyed by tinnitus. When you have achieved this, you will know that you have reached your goal. You can set mini-goals along the way of reading for 10 min at a time. Each time you reach one of these mini-goals you know that you are making good progress towards your overall goal. This allows you to monitor progress – and to have mini-celebrations along the way!

Attainable: The goals you set for yourself need to be achievable. The goals also need to be important for you so as to encourage you to make the commitment and put the effort in to reaching them. While goals should challenge you slightly, it is important to set goals which you are likely to achieve. This will set you up for success. Succeeding will encourage you, help to keep you motivated, and give you confidence to set and achieve further goals.

For example, setting a goal of reading an entire book without being annoyed by the tinnitus may not be feasible, whereas reading several chapters may be.

Realistic: Goals need to be realistic. Realistic does not mean easy, but it does mean do-able. The goals you set need to be reachable, relevant, and meaningful to you. You will need to devise a plan that makes reaching your goal a realistic proposition.

Timely: Put a timeframe on your goal. Setting an endpoint for your goal gives you a clear target to work towards and helps to encourage you to put in a consistent effort. Look for signposts along the way indicating progress towards your goal. Include these mini-goals in your time frame. Without a time frame in which to accomplish your goals, the commitment to achieving them becomes too vague.

TIPS:
Telling others about your goals may provide you with support and encouragement.
Take the time to look back, notice the progress you have made, and celebrate your successes!
Use these SMART strategies to get you where you want to be. Identify what you want to do. Set your goals and GO FOR IT!

Task 2 Sleep Hygiene

Background for Clinician

One of the most common tinnitus complaints is poor sleep. Good sleep practices along with relaxation exercises may improve the amount or quality of sleep.

For Person with Tinnitus

Using a number of strategies and forming new sleeping habits can improve quality of sleep. These strategies are commonly referred to as “sleep hygiene.” Good sleep refers not only to quantity of sleep, but also quality of sleep. We want to make sure you get enough, and that what you get is refreshing. In practice, this means getting to sleep and not waking until fully rested!

People tend not to spend a lot of time thinking about their sleeping habits. You might have your dinner, do whatever it is that you normally do, and then just go to bed for the night. However, there are often things that we can do to make a good night’s sleep more likely.

The quality and quantity of our sleep can be much improved by changing some of our habits!

Good sleep hygiene includes the following:

Going to Bed Strategies

Maintain a routine. Try to go to bed and wake up at the same time every day, even on the weekends. Keeping a regular schedule will help your body expect sleep at the same time each day.

Use bedtime rituals. Doing regular things before sleep tells your body that it’s time to slow down and
begin to prepare for sleep (e.g., a warm bath each night before bed).

Relax for a while before going to bed. Some quiet time can make falling asleep easier. Try relaxation techniques.

Write down all of your concerns and worries. Write down your worries and possible solutions before you go to bed so you don’t need to dwell on them in the middle of the night. This allows you to put away your concerns until the next day.

Go to sleep when you are sleepy. When you feel tired at night, go to bed.

Don’t nap through the day. If you find you have to, limit naps to 30 min, as daytime sleep can upset your body clock for sleeping at night.

Falling Asleep (or Getting Back to Sleep) Strategies

Practice your attention control techniques. This will help to keep your mind occupied, will increase your relaxation, and help you to fall back to sleep.

Get out of bed if unable to sleep. Don’t lie in bed awake. Go into another room and do something relaxing until you feel sleepy. Worrying about falling asleep actually keeps many people awake.

Don’t do anything stimulating. Don’t read or watch a stimulating TV program (as the brain receives a mixed message of having to pay attention to something and yet wanting to go to sleep). Don’t expose yourself to bright light. The light gives cues to your brain that it is time to wake up.

Drink some warm milk. Milk may help create feelings of sleepiness.

Consider changing your bedtime. If you are frequently experiencing sleeplessness, think about going to bed later so that the time you spend in bed is spent sleeping.

Sleeping Environment

Make sure your bed is large enough and comfortable.

Make your bedroom primarily a place for sleeping. Use your bed for sleeping or intimacy only. Help your body recognize that your bedroom is primarily a place for rest.

Keep your bedroom peaceful and comfortable. Make sure your room is well ventilated and the temperature is fairly constant. You could use a fan or a bedside sound conditioner to help reduce attention to tinnitus.

Hide your clock. A highly visible clock may cause you to focus on the time and make you feel stressed and anxious. Place your clock so you can’t see the time when you are in bed.

Daytime Habits

Limit caffeine and alcohol. Avoid drinking caffeinated or alcoholic beverages for several hours before bedtime.

Expose yourself to bright light/sunlight soon after awakening. This will help to regulate your body’s natural biological clock. Likewise, try to keep your bedroom dark while you are sleeping so that the light will not interfere with your rest.

Exercise early in the day. Twenty to thirty minutes of exercise every day can help you sleep, but be sure to exercise in the morning or afternoon, not evening. Exercise stimulates the body and aerobic activity before bedtime may make falling asleep more difficult.

Check your iron level. Iron deficient women tend to have more problems sleeping so if your blood is iron poor, a supplement might help your health and your ability to sleep. Check with your doctor as to whether this is a concern for you.

Establishing good sleeping habits will have many positive benefits for you. Remember to give yourself and your body time to adjust to your new sleeping routine. Some of the strategies will be of more use to you than others. The “going to bed” and “falling asleep” strategies may be the ones that create the biggest change in your quality of sleep, so focus on establishing these first.

Here’s wishing you many nights of great sleep!

Task 3 Relaxation

Background for Clinician

Tinnitus can be increased with stress and tension. Relaxation is one strategy toward overcoming the
negative consequences of stress and alleviating some tinnitus effects.

**For Person with Tinnitus**

By relaxing and becoming more calm, the stress driving your tinnitus, or resulting from tinnitus, may be reduced. This may have a positive effect on your mood and reduction in tinnitus annoyance.

*Abbreviated Progressive Relaxation*

After learning the skill of relaxation, this can be quickly tapped into at times of stress.

This exercise involves four muscle groups. You can modify this exercise if needed, simply be sure to include the areas listed below. Follow the principals of holding a muscle tense for 10–20 s and releasing, then relaxing for 15–20 s before moving on to tensing the next muscle.

1. Hands, forearms and biceps
2. Head, face, throat and shoulders, including concentration on forehead, cheeks nose, eyes, jaws, lips, tongue and neck
3. Chest, stomach and lower back
4. Thighs, buttocks, calves and feet

Find a quiet, comfortable place to sit where there are minimal distractions. Use a squeeze ball while doing these muscle-relaxing exercises, as it a useful aid to help a person to identify when a muscle is being tensed and when it is relaxed.

To begin:

1. Curl both fists
2. Tighten the upper arm and forearms as tight as possible
3. Hold them for 10–20 s and then relax them (this is the same for every part that follows)
4. Next wrinkle up the forehead. Simultaneously, press your head back as far as possible and roll it in a clockwise fashion. Then reverse the head roll
5. Now wrinkle up the muscles of your face like a walnut, and then relax them
6. Arch your back (but be careful if you have a bad back) and take a deep breath. Press out your stomach and relax.
7. Put both feet flat on the floor and now pull your toes back toward your face as far as possible. Tighten your shins; now your calves, thighs, and buttocks; now relax them.

Don’t stand up in a hurry after finishing – take a few deep breaths and stand up slowly to give your body a chance to re-orientate itself.

**Deep Breathing Exercises**

Another form of relaxation is deep breathing. This is a simple exercise that does not take a lot of time. Slow deep abdominal breathing is a useful method of reducing anxiety and causing relaxation. Abdominal breathing expands the belly as it expands the lungs. Chest breathing is shallower and does not provide the relaxation that comes with abdominal breathing.

To begin:

- Close your eyes.
- Focus on your breathing.
- Place your hand flat on your stomach.
- Take slow deep breaths, breathing in through your nose to a count of 1-2-3-4.
- As you breathe in, feel your stomach rise under your hand. If you cannot feel your stomach rise under your hand keep practicing to learn the technique – you will know when it is right because you will feel your stomach rise under your hand. Sometimes it can feel awkward to have your stomach go out when you breathe in.

To help learn this technique, imagine that when you take a breath in you are inflating a balloon in your stomach – deep breathing inflates the balloon and your stomach goes out and exhaling deflates the balloon and your stomach goes in.

- Pause.
- Exhale slowly through your nose (or mouth if you prefer) and count down 4-3-2-1.
- Repeat this breathing technique.
- After a few minutes of breathing like this, as you breathe in, think of the work “relax” and as you breathe out say the words “let go”.

**Task 4 Attention Control**

**Background for Clinician**

People may experience tinnitus as intrusive, constantly on their mind and in their thoughts. It can become the unwanted over riding focus of their attention and make
it difficult to think about anything else. This constant awareness can be overwhelming and the cause of much distress. Simple attention control exercises can be useful to shift attention from tinnitus to more useful perceptions. See also Henry and Wilson [2].

**For Person with Tinnitus**

How much of your *time and attention* does your tinnitus take from you? The answer is quite likely “Too much!” One of the most common complaints amongst those with bothersome tinnitus is that it is always on their mind. It takes up too much of their attention. *But – it doesn’t need to be this way!* Although we are not always aware of it, we have some control over what we pay attention to – and we make these decisions many times a day. For example, we might be working on a crossword puzzle while others are watching TV. In that situation there are a number of things competing for our attention, but we are able to choose to pay more attention to one (e.g., doing our crossword puzzle) and less to the other (e.g., watching TV). *With practice, it is possible to take that same control over the attention that you give to your tinnitus.*

Learning this skill of attention control means you will be able to give less consideration to your tinnitus. It won’t be constantly on your mind and in your thoughts. You will be able to better manage your tinnitus and to reduce the associated distress. You may even be able to do more of the things that you enjoy!

With tinnitus (or anything else really!) it is impossible to simply choose not to think about it anymore. But we can control where we focus our attention. We can redirect the focus of our attention from the tinnitus to something else; with practice this can become nearly second nature! There are a number of strategies that can help you learn how to direct the focus of your attention. These include attention control, imagery, and distraction. Without consciously thinking about it, you probably use some of these techniques already. You can use these very same techniques to manage your tinnitus. The aim of all of these techniques is to learn how to control the focus of your attention – to be able to direct your attention from one thing to another at will. The idea is that you will learn how to direct your attention, to and from, the tinnitus under your own control.

**Attention Control Techniques**

A characteristic of human behavior is that we can really only concentrate on one thing at a time. As we focus on a particular thing, other things become less of the focus of our attention and recede into the background of our mind. We can work this to your advantage, with tinnitus becoming less the center of attention and receding into the background of your awareness. The following are two examples of attention control techniques simplified from a self-help book “Tinnitus. A self-help management guide for the ringing in your ears” by Drs’ Henry and Wilson (2002) which you might find useful. Modify them to suit you, and practice making up your own.

**Example 1:** Focus on your breathing. Breathe in and out. Think about breathing in through your nose and out through your mouth. Breathe slowly, deeply. Become aware of each breath. As you focus on your breathing, notice that you have been less aware of other parts of your body. Gently shift the focus of attention from your breathing to your feet. Without moving your feet become aware of any sensations they are feeling. Become aware of each toe. Picture them in your mind. How do they feel? Are they warm, cold? Can you feel your toes resting next to each other? As you focus your mind on your feet, notice that you have become less aware of your breathing. Gently shift the focus of your attention back to your breathing. As before, think about breathing in through your nose, out through your mouth. Become aware of each breath.

Practice switching your attention from your breathing to your hands. Focus on the details of each hand. Then practice directing your attention back to your breathing. Do this with different parts of your body, going back to your breathing in between. Notice how you can control where you focus your attention. *Notice that as you focus your attention on one thing, other things fade into the background.*

**Example 2:** Find a comfortable place to sit. Ask yourself “Where is my attention now?” Is it focused on a thought, a feeling, or a noise outside? Now change your focus to the physical sensations of your body. Does your skin feel cool or warm? Become aware of any other sensations in your body. Spend some time exploring these. Now refocus your attention to the noises around you. Try to identify what they are. Perhaps you can hear traffic outside, birds chirping, or people talking. Now refocus again, focusing your
attention to your hands, picturing each one in your mind. Notice that you can become aware of where your attention is and that you can change the focus of your attention. You are able to deliberately change your attention from one thing to another.

**Distraction**

Distraction can be helpful in taking your mind off what is causing you distress or worry. You probably have some distraction techniques that you already use. These might include going for a walk, watching TV, or reading a book. Here are some others that you could try. Some will suit you more than others – try them all and see!

- Make a list of five things you enjoy doing most
- Listen to some nice music
- Take a walk
- Play a computer game
- In your mind, run through the alphabet backwards from Z to A
- Count backwards from 100 subtracting 6 at a time
- Search for a movie you would like to see
- Plan a shopping list
- Do something nice – for yourself or for somebody else!
- Make a list of other possible distraction techniques you could use

**Task 5 Communication**

**Background for Clinician**

Tinnitus and accompanying hearing loss can lead to communication difficulties. Communication is such an important activity we seldom think of the detrimental effects of being unable to effectively communicate. Reduced communication can lead to isolation and miscommunication can lead to negative consequences for relationships with family and friends.

For person with tinnitus and hearing loss

- Let other people know you have difficulties hearing. Tell them what they can do to help make things easier. Be specific. Let them know that you need their help because you value what they have to say.
- Place your back to the main source of background noise and face the speaker.
- Ask people to get your attention before they start talking to you.
- Face the person you are talking to so their gestures and facial expressions will help you understand what they’re saying.
- Try to choose a place that is well lit, so it is easy to see the target speaker.
- Try to keep calm and don’t panic. If you become anxious or flustered, it might be harder for you to follow what’s being said.
- Have patience, good humor, and be understanding with yourself.
- If your hearing is not the same in both ears, try turning your better side towards the person speaking to you.
- If you don’t catch what someone says, don’t be afraid to ask him or her to repeat it or say it in a different way.
- If necessary, ask people to slow down and speak more clearly.
- Don’t be too hard on yourself. No one hears correctly all the time!

**References**

52. Newman CW and SA Sandrige, (2005) Incorporating group and individual sessions into a tinnitus management


Chapter 71
Cognitive Behavioral Treatment (CBT)

Karoline V. Greimel and Birgit Kröner-Herwig

Keypoints

1. Cognitive behavioral interventions are the most widely used psychological strategies for coping with tinnitus.
2. The goal of the therapy is to alter maladaptive cognitive, emotional, and behavioral responses to tinnitus and not to abolish the sound itself.
3. There are two main components to this approach:
   (a) Cognitive restructuring and
   (b) Behavioral modification.
4. Treatment programs comprise of techniques like relaxation training, cognitive restructuring, attention control techniques, imagery training, and exposure to difficult situations.
5. The combined approach assists patients in identifying and modifying maladaptive behavior and promotes habituation to tinnitus.
6. The collaboration of patient and therapist is a prerequisite for a positive outcome of therapy.

Keywords
Tinnitus • Cognitive behavioral therapy • Relaxation training • Cognitive restructuring • Attention control techniques • Imagery techniques • Behavioral techniques

Abbreviations
CBT Cognitive behavioral therapy
PMR Progressive muscle relaxation
RET Rational-Emotive Therapy

Introduction

In the history of tinnitus research and treatment, many attempts have been directed toward abolishing or minimizing tinnitus. Despite all these efforts, until now no treatment has been found to successfully eliminate tinnitus permanently. As a consequence, increasing efforts have been undertaken by behavioral scientists and psychologists to eliminate or at least ameliorate psychological symptoms associated with tinnitus. The aim of psychological interventions is not to “cure” or to eliminate the inner noise but to reduce tinnitus-related distress and increase quality of life. If patients are no longer bothered by their inner noises and the question of how tinnitus can be removed, they might become secondary. As long as tinnitus itself cannot be eliminated, the main intention of all therapeutic interventions is to alleviate suffering from tinnitus.

Cognitive Theories of Behavior Regulation

Most interventions in reducing tinnitus-related distress are predicated on cognitive theories of behavior regulation. One of the most influential theories was developed by Beck [1, 2]. Cognitive behavior therapy is based on the “rationale that an individual’s affect and behavior are largely determined by the way in which he structures the world.”

A general cognitive framework as shown in Fig. 71.1 asserts that the emotional and behavioral consequences of an event or situation experienced by a person are modified by the way a person thinks about it. In other words, emotions and behavioral reactions are the result of
appraisals of an event and are not the result of the event itself. This model dates back to Ellis (1973) [3], who termed it the A-B-C model. A stands for activating events, B for beliefs, and C for consequences (see Fig. 71.1).

Patients have to be educated and instructed according to this model. It is made clear that mainly the thoughts, beliefs, and expectations about tinnitus are creating the problem. Tinnitus itself does not have the power to ruin one’s life. This assumption can be illustrated by the fact that the majority of individuals permanently afflicted by tinnitus— even if they describe it as loud— do not feel distressed by it. Nevertheless, the therapist should make it explicitly clear that he or she knows that the tinnitus is real, not imagined, and that the patient’s response to the abnormal tinnitus perception can be well understood.

In general, patients blame their tinnitus for their emotional impairment. They are convinced that the tinnitus is “making” them depressed, anxious, and worried and that their ways of dealing with tinnitus are of no account. Furthermore, if the patient thinks that there is nothing that can be done to alleviate the symptoms, he or she will likely become hopeless and depressed. Blaming a situation or person for the onset of tinnitus will create anger and hostility (see Fig. 71.2).

Cognitive responses to tinnitus can be very different. Regardless of the cause of tinnitus, “suffering” is a function of how the patient reacts to tinnitus— how he or she copes with it. Patients have to be made aware that their way of coping can be modified. The goal of the therapy is to alter maladaptive cognitive, emotional, and behavioral responses to tinnitus and not the sound itself.

A comprehensive model for the chronification of tinnitus, including various dysfunctional cognition and behavior, is described by Kroener-Herwig [4] based on the assumptions regarding tinnitus tolerance made by Hallam and Jakes [5] (see Fig. 71.3).

This model describes the vicious cycle of tinnitus distress and demonstrates how different cognitive, emotional, and behavioral factors interact and create positive feedback loops generating and maintaining tinnitus-related annoyance and discomfort. Attention plays a pivotal role. Focusing attention on tinnitus, accompanied by specific dysfunctional cognitive processes of appraisal like catastrophic thoughts and rumination, leads to negative emotional consequences. Furthermore, behavior resulting from illness often based on avoidance learning (e.g., exculpation from daily routine, justification for absence of work) can be reinforced by family or friends.
Cognitive Behavioral Therapy

The origin of cognitive behavioral therapy goes back to the 1950s and 1960s when Wolpe and Lazarus [6] developed new techniques for changing behavior – in particular in patients with anxiety disorders – based on experimental psychology. In the early phases, therapy (then called behavior therapy) was dominated by techniques like operant conditioning [7], systematic desensitization [8], or aversion therapy [9], which were directed at overt behavior change. In the 1970s, increasingly cognition-centered theories of behavior regulation were established and, consequently, cognitive interventions were implemented into therapy. Beck [2] has been most influential in introducing cognitive interventions into therapeutic strategies. In accordance with this trend, Ellis [10] introduced Rational-Emotive Therapy (RET) based on his A-B-C model. Meichenbaum [11] introduced the notion of the specific importance of self-talk and self-instructions for behavior regulation. Bandura [12] states in his social learning theory that self-efficacy beliefs play a most important role in guiding behavior.

Cognitive behavioral therapy was developed mainly as a treatment for affective disorders such as depression and anxiety. Subsequently, this therapy has been successfully utilized for patients with aversive medical conditions (e.g., chronic pain). A cognitive behavioral approach was first applied in the treatment of patients with tinnitus in the 1980s [13–16]. Nowadays, it is one of the most widely used and accepted psychological strategies for coping with intractable disorders [17–19].

There are two main components to this approach:

- Cognitive restructuring
- Behavioral modification

The combined approach assists patients in identifying and modifying maladaptive behavior and promotes habituation to tinnitus. The collaboration of patient and therapist is a prerequisite for a positive outcome of therapy.

Treatment programs comprise of techniques like relaxation training, cognitive restructuring, attention control, imagery training, and exposure to difficult situations.

Relaxation Training

Relaxation methods were one of the earliest psychological treatments applied to patients with tinnitus [20]. There are several forms of relaxation training. The most common is progressive muscle relaxation (PMR). In this technique, a person is shown how to decrease muscular tension and to achieve states of relaxation in a very brief period of time after detecting tension. The therapist instructs patients how to sequentially tense and relax various muscle groups, moving from practice in comfortable settings to practice in real-life
situations such as sitting at a desk, watching television, etc. Relaxation techniques may be helpful in assisting people in learning a way of coping with tension and anxiety related to tinnitus. Furthermore, it is commonly reported by patients that stress exacerbates tinnitus or causes a person to experience the tinnitus more intensely, and that a reduction in stress levels may reduce loudness and annoyance. Also, listening to one’s tinnitus in a relaxed state can foster habituation and retain serenity in the presence of tinnitus.

Despite the popularity of relaxation training in clinical practice, research shows that relaxation seems to be of limited value for most tinnitus patients when used as the sole treatment [21]. To be successful, it has to be an integral part of a larger treatment program.

**Cognitive Restructuring**

In general, cognitive therapy involves the identification of dysfunctional beliefs and negative thoughts, which occur in response to life events or sources of distress. Patients are taught methods of challenging those thoughts and substituting their catastrophic, unrealistic thoughts with more constructive ones (cognitive restructuring). Cognitive restructuring helps patients think differently and adopt a different attitude about their problem. It is used as a method to guide patients recognize and subsequently abandon rigid, unhelpful thinking patterns and replace them with constructive cognitions and thoughts. This is different from simple “positive thinking” or from “directive counseling,” a treatment component of tinnitus retraining therapy [22], because in cognitive restructuring the therapist and patient collaborate in identifying, testing, and challenging dysfunctional thoughts, beliefs, attitudes, or attributions [4, 23]. It is theorized that for patients with tinnitus, the source of distress is not the sound itself, but the way in which the person evaluates and interprets the sound. A person may have negative thoughts such as “The noise is driving me crazy” or “This is the worst thing that could ever happen.” Alternatively, he or she could think: “The noise doesn’t hurt me – it is bad, but it generally gets better by-and-by” or “Do something enjoyable, rather than being occupied with the noise in your head.”

The therapist helps the clients to challenge and test the validity of their automatic thoughts and to learn ways to substitute them with more constructive ones.

**Attention Control Techniques**

Attention control interventions make patients aware that they indeed have control over their attentional focus, and that directing attention to other aspects of the external or internal environment can make the tinnitus “disappear.” Instructing patients to switch attention to and from tinnitus illustrates that tinnitus can be “controlled.” Patients are encouraged to augment their use of other sensory modalities (e.g., smelling coffee, tasting honey). Furthermore, strategies to specifically manipulate the acoustical environment are recommended.

**Imagery Techniques**

Imagery techniques are used to change the negative associations related to tinnitus either by “masking” the noises or by integrating them into positive scenes. In imagery exercises, a patient may be asked to imagine that the tinnitus is masked by the sound of a waterfall or the waves of the sea. No real sounds are used to mask tinnitus in this exercise. Masking is achieved by imagination. Tinnitus also may be incorporated into pleasant scenes. Patients might be instructed to imagine walking through a landscape by listening to the singing of birds or lying on a blooming field and hearing the noises of bumblebees, cicadas, and other insects. Alternatively, a patient may imagine a cold and snowy winter day, sitting comfortably in front of the fire place, hearing the sizzling of a teakettle, and looking forward to enjoying a cup of hot tea.

In clinical practice, these approaches are rarely used as sole therapeutic methods, but are incorporated into relaxation training or cognitive restructuring interventions.

**Behavioral Techniques**

Tinnitus patients may tend to avoid situations where they feel impaired or distressed by their tinnitus, i.e., conversations with more than one person, a concert, a stroll in the city. This may have developed into generalized avoidance behavior. Cognitive behavioral therapy (CBT) encourages patients to expose themselves to those situations in order to realize that they can cope without major negative
consequences. These behavioral “experiments” must be well prepared and these new skills should be frequently practiced.

In some patients, “suffering” from tinnitus allows her/him to avoid situations, which were threatening, and anxiety inducing for non-tinnitus-related reasons, e.g., office work or participating in social events. Tinnitus for them is an acceptable solution or “a legitimate excuse” for avoiding these situations. Thus, tinnitus complaints are under operant control and are therefore maintained. In these cases, patients have to become aware of the underlying problem and are assisted in finding adaptive problem solutions.

Multimodal CBT has been evaluated in several studies. The meta-analysis of Anderson and Lyttkens [21] showed that psychological treatments are very effective regarding the reduction of tinnitus-related distress. The average effect size of 0.86 reveals a high efficacy. Recently, Martinez Devesa et al. [24] prepared a meta-analysis of randomized controlled group trials on CBT for the Cochrane Collaboration and came to the conclusion that the CBT is effective for improving the quality of life based on the analysis of 24 trials.

References

**Chapter 72**  
**Auditory Training in Tinnitus**

Larry E. Roberts and Daniel J. Bosnyak

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**Keypoints**

1. We reviewed sensory training studies from the point of view that tinnitus is caused by synchronous neural activity that develops in tonotopic regions of primary auditory cortex deafferented by hearing loss. Studies were classified according to whether training was conducted within the tinnitus frequency region or outside of it, and whether training was active (requiring behavioral responses) or passive (sounds were presented as background signals). Effects of training on the psychoacoustic properties of tinnitus were distinguished from those on the distress behavior that accompanies tinnitus.

2. Studies in all four categories have reported significant reductions compared to untreated controls in tinnitus distress, measured by standardized questionnaires and visual analogue scales at the first compared to untreated controls in-course assessment, with little further change thereafter. Because the particular details of sensory training do not appear to matter, these gains could reflect important nonspecific effects of the treatment procedures.

3. Psychoacoustic measures may more directly assess tinnitus sensations. Reductions in minimum masking level (MML) on the order of 5–10 dB have been reported by several studies, implying that tinnitus has become weaker. Improvements in loudness discomfort levels (LDL) have also been reported, as have changes in the frequency content of tinnitus. Improvements in MML and LDL are more gradual than those on distress behavior assessed by questionnaires, suggesting that neural plasticity may be at work.

4. Several studies reporting improvements in psychoacoustic measures and questionnaire data used passive sound presentation procedures. Hence, active sensory training requiring discriminated behavioral responses is not needed for these changes.

5. Systematic manipulation of the frequency content of trained sounds has been attempted in only a few studies. This step is needed to determine whether sound training induces specific changes in tinnitus. Alternatively, sound therapy may amplify the nonspecific effect of elements common to all tinnitus therapies.

6. Future studies should continue the practice of specifying how many participants of the total recruited contributed to a data analysis, and why and when exclusions occurred. Substantial sample sizes will be needed to establish treatment effects. Neural correlates offer the advantage of comparative immunity to patient expectations and self-report bias. When sounds are used to evoke neural responses, changes in loudness recruitment consequent on rescaling of loudness growth functions by sound exposure are a potential contributing factor.

**Keywords** Tinnitus • Sensory training • Tinnitus distress • Minimum masking level • Loudness discomfort levels

**Abbreviations**

- A1 Primary auditory cortex
- A2 Secondary auditory cortex
- ADT Auditory discrimination training
- AOIL Auditory object identification and localization
In 1995, Jastreboff [1] proposed a comprehensive model of tinnitus that addressed three clinically prominent features of this condition. These were (a) the tinnitus sensation itself, generated by pathology in the inner ear; (b) the ability of the tinnitus sensation to command attention; and (c) the patient’s disturbing emotional reaction to the tinnitus percept. Jastreboff suggested that although elimination of the tinnitus sensation by treatment of cochlear pathology was in most cases not practical, the latter two features of tinnitus were likely modifiable and if treated would benefit the tinnitus patient. Tinnitus retraining therapy (TRT) was devised to foster extinction of attentional and emotional responses by presenting low-level tinnitus-like external sounds that could be filtered out along with the tinnitus by perceptual mechanisms (see Chap. 73). Studies of TRT and clinical experience have confirmed that emotional responses diminish with time for most tinnitus sufferers, as does the extent to which tinnitus sufferers attend to their tinnitus percept [2]. These are important and beneficial effects for tinnitus sufferers. Attempts to reduce or eliminate the tinnitus sensation itself, however, have met with less success.

One approach that has gained attention with respect to the latter goal in recent years is sensory training aimed at modifying the neural basis of tinnitus sounds. The inspiration for this approach was based in part on the discovery that hearing loss induced by noise exposure in animal models leads to a substantial reorganization of tonotopic maps in primary auditory cortex, such that frequencies near the edge of normal hearing come to be overrepresented at the expense of frequencies in the hearing loss region [3–5]. Because hearing loss is a putative cause of tinnitus, it was suggested that this overrepresentation, or changes in the response properties of auditory neurons associated with it, may correspond to the tinnitus percept [6, 7]. A second foundation was laid by experiments conducted in the last 15 years that demonstrated that cortical representations for sound in the primary auditory cortex are not fixed after early development, as was once believed, but can be modified by auditory training well into adulthood [8, 9]. This phenomenon is called “neural plasticity” (see Chap. 12). These two lines of research have converged to ask whether sensory training procedures derived from animal research can be adapted to humans, with the goal of modifying neural representations that appear to underlie tinnitus.

**A Framework for Sensory Training Studies**

For this goal to be achieved, the neural modifications induced by sensory training must intersect with the neural mechanisms generating tinnitus. In Chap. 13, we reviewed evidence pointing to a role for neural synchrony (temporally coupled neural activity) in tinnitus. According to this viewpoint, tinnitus may be generated by synchronous neural activity that develops in reorganized tonotopic regions of primary auditory cortex that receive diminished input from the ear owing to hearing impairment caused by noise exposure, otological disease, or the aging process [7]. Changes in subcortical structures appear to contribute [10] and may account, as well, for some distinct properties of tinnitus including its modulation by somatosensory activity in many individuals [11, 12]. Although the thalamocortical input to the affected neurons in the primary auditory cortex (A1) is altered by hearing loss, their synchronous output remains intact and may be a driving force underlying tinnitus. This output may recruit other brain regions into a network identified by functional imaging studies [13], including frontal and limbic areas that subserve, respectively, the attentional and emotional aspects of tinnitus described by Jastreboff [1].

In this chapter, we use the neural synchrony model as a template for reviewing auditory training studies of tinnitus. This perspective suggests that it is necessary
to reduce synchronous neural activity occurring in regions of A1 that have been affected by hearing loss, in order to reduce the loudness of tinnitus sounds. Training for sounds in the tinnitus frequency region, with the aim of segregating synchronous network activity in this region, would appear to be the most direct approach. Masking sounds presented to this frequency region induce optimal post-masking suppression of tinnitus [residual inhibition, (RI)], confirming that such sounds interact with the tinnitus generating mechanism [14]. Training in the tinnitus frequency region requires that significant residual hearing be present in this region, which is the case for many, but not all, tinnitus patients. Alternatively, training can be delivered outside of the tinnitus frequency region where hearing is generally better preserved. For example, training at or below the edge frequency region may alter neural representations in these regions, which send collateral inputs into the tinnitus region that may disrupt neural synchrony. Lateral inhibition arising from augmented representations below the tinnitus frequency range could also distribute into the tinnitus region and suppress tinnitus percepts. Inhibitory interactions have been demonstrated by human electrophysiological studies [15, 16] and are known to span several octaves in primate A1 [17], suggesting the feasibility of this approach. While the neural synchrony model focuses on A1 as a preferred site of action, several brain structures are active in tinnitus, including regions of the secondary auditory cortex (A2) that may distribute re-entrant feedback into the auditory core region and disrupt neural activity underlying tinnitus [18]. Remodeling of cortical representations in A2 by sensory training appears to proceed normally in the tinnitus brain (see Chap. 13) and may confer a benefit.

Several methodological limitations should be acknowledged in advance of this review. Auditory training procedures are aimed at modifying the neural processes that generate tinnitus sensations. In order to assess whether this goal has been achieved, it is desirable to employ psychoacoustic tools that more or less directly measure the sensory attributes of tinnitus, such as loudness matching (LM), in which the loudness of an external sound in the range of normal hearing is adjusted to equal the loudness of tinnitus, and minimum masking level (MML), the minimum loudness of a masking sound required to just cover tinnitus. Loudness discomfort level (LDL) is another useful psychoacoustic method, which measures loudness growth functions that are frequently elevated in individuals with tinnitus [19], as are their audiograms. Standardized procedures for measuring tinnitus spectra are also available [14, 20] and beneficial for characterizing tinnitus. However, only a minority of studies report such measures. More often, standardized questionnaires such as the tinnitus handicap questionnaire (THQ) [21], tinnitus severity index (TSI) [22], tinnitus handicap inventory (THI) [23], and tinnitus reaction questionnaire (TRQ) [24] are employed in which tinnitus patients rate on subjective scales the loudness and intrusiveness of their tinnitus and its effect on quality of life including mood and anxiety, interference with sleep, concentration, work productivity, and interpersonal relationships. While these questionnaires—often supplemented with tinnitus ratings on visual analog scales (VAS)—likely reflect to some degree the sensory properties of tinnitus, they tend to focus on the distressful consequences of having tinnitus emphasized by Jastreboff [1]. A further limitation is that few studies have controlled for the contribution of procedural elements that are likely common to all therapeutic approaches and may affect outcome regardless of any direct effect of auditory processing on the neural substrate of tinnitus. Examples of such elements include (a) beneficial effects of discussion with informed and sympathetic staff, (b) knowledge about tinnitus, (c) investment by patient and staff in a therapeutic process, and (d) the effect of these components on a hopeful attitude and expectations for success. In this chapter, we will refer to effects of these elements as “nonspecific effects”, not to diminish their considerable importance for benefiting patients, but in order to distinguish them from effects attributable to the specific sounds incorporated into an auditory training procedure.

Notwithstanding these limitations, several approaches to auditory training have been tried or are currently under assessment. The results give a picture of the methods used, whether the goal of auditory training can be realized, and if not for all tinnitus patients, which variables may be important for treatment success. Because active training requiring explicit behavioral responses might confer a benefit in tinnitus, we categorize the studies into active procedures that require such responses and passive procedures that do not. We also categorize the studies according to whether sounds are presented to the tinnitus frequency
(hearing loss) region or outside of this region. Animal studies are included where they are relevant. One novel approach is described that does not fit into these categories.

**Active Training Within the Tinnitus Frequency Region**

Several studies have assessed the effects of auditory training procedures at or near the tinnitus “pitch” (likely resembling the modal pitch in a tinnitus spectrum). Based on their results studying phantom limb pain where it had been shown that the amount of cortical reorganization was positively correlated with amount of pain [25] and that discrimination training in sensory areas adjacent to the deafferented region reduced phantom limb pain [26], Flor et al. [27] trained seven tinnitus patients on a frequency discrimination task for tones matched to their tinnitus frequency (proximal frequency group), with an additional seven patients trained at a frequency distant from the tinnitus frequency (distal group). The participants in this study were asked to determine if two tones presented successively were either identical (50% of trials) or different in frequency and were given feedback for correctness. The difficulty of the task was increased with performance improvement across sessions. Training was to be carried out every day for 2 h over a 4-week period. Interestingly, two of the seven distant-frequency participants dropped from the study complaining of increases in tinnitus severity, suggesting an adverse effect of training below the tinnitus frequency region. At the end of training, the proximal and the remaining distal patients did not differ on any outcome measures, so they were combined for analysis. Given the unreliability of tinnitus pitch match procedures [28], some patients in the distal group may still have trained at frequencies within their tinnitus spectrum. No significant training effect on tinnitus severity was found, but not all patients complied with the training requirements. When the participants were separated post hoc into those who trained more \((n=7)\) or less \((n=5)\) over the 4-week period, the extensive training group showed significant reduction in self-reported tinnitus severity while the limited training group showed a significant increase in tinnitus severity. Cortical reorganization or changes in the psychoacoustic properties of the tinnitus were not assessed. Given that the treatment effect was not limited to the group training on frequencies within the tinnitus frequency region, it appears that potentially nonspecific factors such as focusing attention away from tinnitus might have been responsible for the lessening in severity.

Herriaz et al. [29, 30] described the results of a number of similar procedures, which they collectively referred to as ADT (auditory discrimination training). In all patients, the stimuli to be discriminated fell within the region of hearing loss. However, the procedures differed from those used by Flor et al. in that the discrimination in most cases was relatively easy (for example, discrimination between a broadband noise and an 8 kHz pure tone) and task difficulty did not increase with training (non-adaptive procedure). Training sessions were relatively short in the largest test group \((n=29)\), with the participants required to perform 10-min sessions twice daily for a 1-month period. These procedural changes allowed the patients to perform the task at home using an MP3 device. Significant improvements in self-reported tinnitus severity on a VAS scale of loudness and total score on the THI questionnaire were found compared to waitlist controls. However, because no assessments of the psychoacoustic properties of tinnitus (LM, MML, or tinnitus spectrum) were performed, it is difficult to attribute the tinnitus improvement to a reversal of the presumed cortical reorganization. In another study, participants in one group (SAME, \(n=11\)) trained at frequency discrimination at a pitch judged to be the same as the tinnitus pitch while the second group (NONSAME, \(n=11\)) trained at a frequency different from the tinnitus pitch but still within the region of hearing loss. The NONSAME group showed a larger reduction in THI score with the difference between the groups being significant. Like Flor et al. [27], these results suggest that training at the “tinnitus pitch” was not a requirement for reduction in tinnitus severity. However, because the trained pitch in the NONSAME group was in the region of hearing loss, some degree of overlap with the tinnitus spectrum was likely.

Norena et al. [20] trained a single individual on a frequency discrimination task for four frequencies within the participant’s measured tinnitus spectrum, and also measured the frequency discrimination threshold during training using an adaptive forced-choice staircase procedure. Training occurred in seven sessions over 3 weeks and was performed monaurally
although the participant had bilateral tinnitus. The tinnitus spectrum changed significantly post-training in the trained ear but not the untrained one, showing a marked reduction in likeness ratings at the highest frequencies. This individual reported informally that the tinnitus sensation shifted from the initially more salient trained ear toward the untrained ear. However, the changes in the tinnitus spectrum occurred at the highest measured frequencies rather than at the frequencies used in the training procedure. This raises the possibility that the changes observed in the tinnitus spectrum could be attributed to an improved ability of the participant to make better discriminations at higher frequencies, allowing more refined judgments of the tinnitus spectrum. The unilateral effect of the tinnitus spectrum change supports the idea that the discrimination training process induced changes in the frequency organization in the auditory cortex. Follow-up studies employing more participants are called for.

In a preliminary study of our own (see Chap. 13 and [31]), we departed from the frequency discrimination training paradigm to one requiring detection of targets of increased sound intensity that were embedded in a 40-Hz amplitude-modulated tone of 1-s duration (carrier frequency 5 kHz, in the tinnitus frequency region). This type of stimulus evokes the stimulus-driven 40-Hz “auditory steady-state response” (ASSR) that localizes tonotopically to the region of primary auditory cortex and gives a picture of events occurring in this region during auditory training. Previous research with frequency (not intensity) discrimination had shown that acoustic training advanced the phase of the ASSR (a shortened time delay between the 40-Hz stimulus and response waveforms), but the amplitude of the response (signaling a map expansion) did not change [32]. We therefore switched to the intensity discrimination procedure using a single carrier frequency, which reduced competitive interactions that may obstruct map expansions when several carrier frequencies are experienced [33]. If training at 5 kHz strengthened the thalamocortical tuning of the trained neurons, tinnitus might diminish at this frequency as the affected neurons were removed from synchronous network behavior underlying tinnitus. Measurement of the tinnitus spectrum before and after training showed little change at 5 kHz or any other tinnitus frequency after training. However, in individuals with tinnitus, auditory training did not change ASSR phase either (n=8 participants, p=0.44), although it did so in their age-matched controls (n=11 participants, p=0.006) suggesting impaired remodeling of primary auditory cortex in the tinnitus group. A different brain response that is known to be neuroplastic [32] and to localize to secondary auditory areas is the P2-evoked auditory potential (latency ~ 180 ms). P2 amplitude increased with training in both groups ([31]; see Chap.13), suggesting normal remodeling of secondary areas in tinnitus. However, this remodeling had no effect on tinnitus. The results of this study could change as additional participants and groups are tested.

Active Training Outside the Tinnitus Frequency Region

Based on the proposal that the tinnitus percept elicits abnormal levels of attention, Searchfield et al. [34] trained 10 individuals with tinnitus on an auditory object identification and localization (AOIL) task designed to refocus the participants’ attention on external stimuli. Training (approximately 30 min per day over 15 days) consisted of up to 20 listening tasks that required subjects to identify and locate in space (left, right, centre) a number of common sounds (e.g., spoken words, owl hooting, coughing, dog barking) against a variety of background noises. The frequency of the sounds and background noises were not explicitly designed to fall below the frequency region of hearing loss or tinnitus spectrum, although the dominant frequencies were likely in this region. Subjects showed a 6-dB reduction in tinnitus loudness assessed by LM, and a significant reduction in pitched matched MML (in eight of ten participants, up to 30 dB in one person). The experiment is noteworthy for its inclusion of psychoacoustic measures. This type of training explicitly targeting the attentional system (but not using sounds focused within the tinnitus region) produced changes similar to those seen in other training procedures that presented stimuli within the tinnitus spectrum.

Another approach similar to active training on sound discrimination is the restoration of behaviorally relevant input via prostheses. There are a number of studies that report cochlear implants having a suppressive effect on tinnitus (see Baguley and Atlas [35] for a review) (see Chap. 77), and hearing aids have also proven to be beneficial (see Chap. 74). Folmer et al. [36]
found that out of 50 patients purchasing and wearing a hearing aid, 46 reported at least “a little” improvement in their tinnitus, with 11 reporting “very much” after 6–48 months. The self-rated loudness of their tinnitus was significantly reduced from 7.5 to 6.3 out of 10 on a VAS. The matched pitch of their tinnitus was 4.3 kHz, which likely means that the aids (which typically have low-frequency amplification profiles) restored little input near their tinnitus frequency. However, Moffat et al. [37] fitted nine subjects with hearing aids with a high bandwidth amplification regime (20 dB threshold reductions at 6 and 8 kHz) and found no changes in the tinnitus spectrum or tinnitus loudness after 30 days. Interestingly, a second group fitted with a low-medium frequency amplification hearing aid showed a significant diminution of low-frequency components of the tinnitus spectrum, with no effect seen at middle or high frequencies. The authors suggested that the perceptual characteristics of tinnitus depend on a contrast between adjacent central auditory regions of more and less afferent activity, which was increased by the low frequency amplification profile. The limited malleability of the tinnitus percept in the high amplification group may be due to the extent of hearing loss in this region and the robustness of neuroplastic changes that give rise to tinnitus. Neither amplification group, however, reported a reduction in tinnitus when assessed by LM.

**Passive Experience Within the Tinnitus Frequency Region**

Restoration of input via prostheses restores auditory input in a behaviorally relevant manner, which supports classification of these procedures as active training. However, animal data (and training studies in normal hearing humans) suggest sound input need not be behaviorally relevant in order to effect changes. Norena and Eggermont [38] found that tonotopic map reorganization in cats exposed to traumatic noise can be prevented by subsequent immersion in an enriched acoustic environment (EAE) containing background sounds designed to compensate for the frequency-dependent decrease in sensory inputs from the hearing loss region. This procedure also led to a recovery from hearing loss between 16 and 32 kHz in the EAE cats, compared to cats exposed to an identical noise trauma but placed in a quiet environment (QE). The increased spontaneous firing rates and increased neural synchrony, which underlies the neural synchrony model, were also absent in EAE cats [39]. Subsequent research showed that passive exposure to the EAE for 6 weeks can produce tonotopic reorganization in normal adult cats in the absence of any noise trauma, suppressing sound representations in the EAE frequency region, and without inducing any threshold changes [40]. These findings accord with other data indicating that passive exposure to environmental sounds can lead to neuroplastic changes in the absence of explicit training requirements [41–44].

Is restoring acoustic input in the tinnitus frequency region, even if this input is not behaviorally relevant, sufficient to normalize frequency representations and reduce the neural synchrony possibly underling tinnitus, in subjects for whom significant residual hearing is present in this frequency region? The most direct evidence comes from three studies initiated by Neuromonics (see Chap. 75), a private company (http://www.neuromonics.com) that markets a device that delivers spectrally manipulated music tailored to augment frequencies in the hearing loss region of the patient’s audiological profile. Because the tinnitus spectrum typically tracks the hearing loss region [14], this sound (presented at levels covering fully or partially the tinnitus) would be expected to inject feed forward and surround inhibition into the relevant region, disrupting the tinnitus sound. Patients screened for residual hearing in the loss region were instructed to listen passively to the sound for at least 2 hours per day using a high fidelity sound player with ear phones over a treatment period of 12 months. In the initial months, patients were told to set the sound level so that their tinnitus was fully masked, and then in subsequent months to gradually reduce this level, so that tinnitus was intermittently heard. This sound therapy approach was combined with counseling following the method of systematic desensitization in which aversive stimuli (in this case, tinnitus) are experienced gradually and in a context conducive to relaxation. In three studies [45–47], Neuromonics treatment led to a substantial reduction in tinnitus distress measured by the TRQ at the first assessment taken 2 months into the study, with little further improvement and little remission in the 10 months of treatment following thereafter. VAS ratings assessing tinnitus severity, ability to relax, and loudness tolerance also improved, following a course similar to the TRQ data. Notably, psychoacoustic
measurements of MML and LDL were also taken in each study. In each study, MML decreased progressively over the 12-month treatment interval, while LDL levels increased.

In order to assess whether sound therapy contributed to these beneficial results, Davis et al. [46] contrasted questionnaire and psychoacoustic data among groups that received Neuromonics treatment (Neuromonics sound therapy with counseling, \(n=21\) subjects), broadband noise masking with counseling \((n=15)\), or counseling alone \((n=13)\). After 12 months, subjects in the Neuromonics treatment group reported a 66% reduction in TRQ scores, compared to reductions of 22 and 15% reported by subjects in the masking and counseling alone groups, respectively (the differences between the Neuromonics group and other two groups were statistically significant). In agreement with these results, tinnitus severity assessed by VAS was reduced in the Neuromonics group, compared to the two control conditions. The Neuromonics group also reported a reduction of 11.3 dB in MML \((p<0.001)\) at 12 months, compared to non-significant reductions of 0.4 and 1.5 dB in the masking and counseling alone groups, suggesting a benefit of Neuromonics treatment on tinnitus loudness. However, an aspect of this study that should be noted is the high proportion of subjects who were either eliminated prior to treatment for failure to meet admission criteria \((n=19/88)\) or were excluded from the final analysis for other reasons \((n=24/88,\) overall exclusion rate 48.9%). Among the exclusions were subjects with entering TRQ scores lower than 14/100 who typically show little gain from treatment [2, 47]. It should also be noted that while improvements in the psychoacoustic measures in the Neuromonics group also reported a reduction of 11.3 dB in MML \((p<0.001)\) at 12 months, compared to non-significant reductions of 0.4 and 1.5 dB in the masking and counseling alone groups, suggesting a benefit of Neuromonics treatment on tinnitus loudness. However, an aspect of this study that should be noted is the high proportion of subjects who were either eliminated prior to treatment for failure to meet admission criteria \((n=19/88)\) or were excluded from the final analysis for other reasons \((n=24/88,\) overall exclusion rate 48.9%). Among the exclusions were subjects with entering TRQ scores lower than 14/100 who typically show little gain from treatment [2, 47]. It should also be noted that while improvements in the psychoacoustic measures in the Neuromonics group suggest that sound exposure mattered, the effect of spectrally enhancing sounds outside rather than inside the tinnitus frequency region has not been investigated. Evidence on this question is needed to determine whether the specific frequency of the sounds that subjects listen to is crucial for therapeutic gains, or whether the experience of sound (regardless of frequency) amplifies nonspecific contributions by increasing patient involvement and treatment plausibility.

Other evidence supports the contention that passive listening to sounds that cover tinnitus frequencies can reduce tinnitus. In a study cited previously, Folmer et al. [36] fitted 50 subjects with in the ear sound generators producing broadband (100–8,000 Hz) noise and found that self-rated tinnitus loudness significantly reduced from 7.6 to 6.2 on a ten-point VAS scale. However, this improvement was about the same as a group fitted with hearing aids that likely did not restore much high-frequency input. TRT provides exposure to a broadband masking stimulus that resembles tinnitus but is presented at lower loudness levels (called the “mixing point”) approximating the tinnitus loudness [1] (see Chap. 73). TRT has been found to lead to decreases in tinnitus distress (measured by the TSI, THI, and THQ) that are initially less than improvements produced by masker therapy [2]. However, after 12–18 months of treatment, improvements induced by TRT exceeded those of masker therapy [2], suggesting that the listening protocol may contribute a role.

Whether covering the tinnitus frequencies are crucial remains unclear, however. In a study modeled on animal data reported by Norena and Eggermont [38, 39], Norena and Chery-Croze [19] exposed individuals reporting abnormal loudness recruitment (hyperacusis) to a background sound containing high frequencies spectrally enhanced over the region of hearing impairment, in a manner similar to EAE-exposed cats. The participants in the study listened to the sound in the background for 3 h per day over 15 weeks. Passive listening rescaled loudness growth functions in the direction of normal hearing over this interval, with some regression over a period of 1 month after passive listening ceased. Effects on tinnitus were not assessed, although a majority of subjects with hyperacusis typically report tinnitus as well [48]. The specific frequency content of the sound was not manipulated in this study (all subjects received a high-frequency amplification profile). In a study of individuals with normal hearing, Formby et al. [49] found that loudness growth functions can be bi-directionally rescaled by enhancing or reducing background acoustic environments. These results, which appear to be mediated in part by subcortical mechanisms [50], show that passive exposure can selectively remodel auditory processing in humans. Whether concomitant effects are seen on tinnitus remains to be investigated.

Passive Experience Outside the Tinnitus Frequency Region

Except for the possibility (discussed above) that effects of hearing aid amplification on tinnitus may be
attributable in part to passive exposure to sounds below the tinnitus frequency region, passive sound therapies restricted to this region have not been widely studied. However, a recent study by Okamoto et al. [51] can be discussed here.

These investigators reasoned that because hearing loss is often present in the tinnitus frequency region, auditory training may be more effective if delivered to frequency regions where hearing is better preserved. Their approach was based on an earlier series of studies by their group in normal hearing subjects [16], which showed that notched sound can suppress neural activations in the notched region by distributing lateral inhibition to these regions. Okamoto et al. [51] therefore gave chronic tonal tinnitus patients in a treatment group daily experience with their favorite music that had a one-octave notch around their dominant tinnitus frequency removed. A placebo group listened to similar musical stimuli, except that the notch shifted over the course of training but was never at the tinnitus frequency. Subjects in the treatment and placebo groups listened for about 12 hs/week over 12 months. A further control group (monitoring) received no treatment but participated in the study measurements. Tinnitus loudness measured by a VAS was significantly reduced from baseline in the treatment group, but changes in VAS ratings did not reach significance in the placebo or monitoring groups. A comparison of the VAS changes between the treatment and placebo group ratings was also significant (this comparison was not made for the monitoring group). Notably, the amplitude of the 40-Hz ASSR and the N1m response to tonal stimuli delivered at the tinnitus frequency were also reduced in the treatment group after their sound therapy, relative to these responses evoked by a control frequency (500 Hz). These brain measures did not change in either of the control groups (a comparison of the treatment and placebo groups was also significant in this measure). Hence, evidence for a brain correlate of tinnitus suppression was observed with the notching procedure. This study is notable for inclusion of control conditions designed to evaluate whether the specific frequency content of auditory training is crucial for tinnitus improvement and for carrying out brain imaging measures. A limitation, however, is that of 39 subjects that met the criteria for entry into the study, only 23 contributed data in the treatment (n=8), placebo (n=8), and monitoring (n=7) groups. Subjects were included in the final statistical analyses only if their subjective tinnitus pitch did not change over the study and if the median of repeated pitch matches fell within the notched region for subjects in the experimental group, which are reasonable criteria for a study of this design. Further research is called for to corroborate the findings and assess the limits and magnitude of possible treatment effects.

Other Approaches

Jepsen and her colleagues have proposed an alternative approach to the treatment of tinnitus based on the concept of category training [52]. This approach is modeled on studies by Guenther et al. [53] in normal hearing subjects, which found that training to classify non-speech stimuli within a particular frequency range as members of the same category (frequency categorization training) led to a decrease in discrimination ability for frequencies within the category. In subsequent research [54], frequency categorization training led to a relative decrease in neural activation measured by fMRI for the trained frequencies, whereas conventional training for discrimination among the same frequencies augmented neural activation for the trained stimuli.

Jepsen et al. [52] hypothesized that it would be advantageous to train subjects experiencing tinnitus to assign tinnitus frequencies to a common category, which might lead to a reduction in activation in this area of cortex and presumably a concomitant decrease in the tinnitus sensation. They trained 20 subjects for 30 min per day for 3 weeks, either to categorize tinnitus frequencies into a group or to discriminate among the frequencies, in each case using a take-home training device. The two groups did not differ markedly in their pre–post THI score changes, but did show differences in auditory-evoked potentials. The categorization group showed a reduction in P2-N1 amplitude post-training while the discrimination group showed an increase, which is in line with the observations of Guenther et al. [54]. However, this change was most evident for a control (untrained) frequency rather than the trained frequency, again indicating a more nonspecific effect of training rather than a reduction in cortical activation for the tinnitus region. Category training merits further investigation for its effects on discrimination ability, neural responses, and tinnitus.
Overview and Conclusion

Animal research in the last two decades has established that neural plasticity is a fundamental property of neurons in the auditory and other sensory systems. Evidence has also accumulated that hearing loss (a triggering factor in many if not most people with tinnitus) leads to changes in central auditory pathways, including tonotopic map reorganization and increased neuron firing rates that may be forged by neuroplastic mechanisms into abnormal network behavior generating tinnitus sounds. These findings have spawned renewed research into the question of whether tinnitus can be reduced or eliminated by auditory training specifically designed to normalize aberrant auditory neuronal representations that are believed to be responsible for tinnitus. For this goal to be achieved, it must be possible to modify auditory representations by acoustic training in individuals with tinnitus, and the neural modifications induced by training must intersect with the underlying tinnitus mechanisms.

In this chapter, we reviewed auditory training studies from the point of view that tinnitus is caused by synchronous neural activity that develops in tonotopic regions of primary auditory cortex that have been deafened by hearing impairments. Studies were classified according to whether training was conducted within the tinnitus frequency region or outside of it, and whether the trained sounds served as cues for behavioral responses and were therefore processed actively in attention, or whether the sounds were presented passively as background signals. We also attempted to separate the effects of auditory training on two distinct aspects of tinnitus emphasized by Jastreboff [1], namely, effects on the tinnitus percept itself and effects on distress behavior that accompanies tinnitus. The following summary statements appear to be justified.

1. The number of auditory training studies is not large, and the studies do not evenly cover the four categories we used for classifying them.
2. Studies in all categories have reported significant reductions in tinnitus distress measured by standardized questionnaires (THQ, TRQ, TSI) and VAS ratings. These reductions typically achieved their maxima at the first in-course assessment, with relatively little if any gain thereafter. A noteworthy result is that two treatment procedures that manipulated the frequency content of sounds in the tinnitus frequency region in opposite directions [46, 51] reported similar tinnitus reductions in VAS ratings. If the particular details of auditory training do not matter for these improvements, these gains would appear to be attributable to nonspecific effects of the treatment procedure.
3. Because these changes on questionnaires and VAS ratings are beneficial for patients, it is important to identify the factors responsible for them. Benefits may be greater when some form of sound therapy is employed, although further evidence on this point and particular sound therapy used is needed. Another factor relevant to a successful treatment outcome is opportunity for improvement. Several studies have reported that reductions in distress behavior are minimal when tinnitus distress is low at study commencement.
4. Changes in psychoacoustic measures have been reported that may more directly measure tinnitus sensations. Reductions in MML on the order of 5–10 dB have been reported by several studies [34, 45–47], implying that the tinnitus sensation has become weaker. MML may be a better measure of tinnitus loudness than adjusting external sounds to match tinnitus, which is known to be frequency dependent [14]. Improvements in loudness tolerance (LDL) have also been reported [45–47], as have changes in the frequency content of tinnitus [20, 37]. Improvements in MML and LDL are more gradual than those on distress behavior, suggesting that some form of neural plasticity may be at work.
5. Several of the studies reporting improvements in psychoacoustic measures used passive sound presentation procedures. Hence, active training requiring discriminated behavioral responses does not appear to be necessary for changes in psychoacoustic measures. This observation aligns with experiments in normal hearing animals and humans which found that passive exposure to sound can be sufficient to remodel auditory representations.
6. Animal data and the neural synchrony model of tinnitus imply that training for sounds that cover the tinnitus frequency region is likely to be most effective in modifying tinnitus, provided that residual hearing is present in this region. The results on this point are, however, conflicting. With a few exceptions [27, 37, 51], systematic manipulation of the...
frequency content of the trained sounds has not been attempted in auditory training studies. Loudness growth curves are rescaled in normal hearing individuals by augmenting or reducing background sound [49], and rescaling occurs in hyperacusis patients exposed to high-frequency complex sounds [39], in both situations with broad frequency selectivity. However, applications of these procedures to tinnitus remain largely untested. Because the measurement of brain correlates often involves presenting sounds, effects of sound therapy on loudness recruitment are potential contributing factors to such measurements in tinnitus.

While these conclusions are less than satisfying, they do give guidance for continuing study. Future research should emphasize psychoacoustic measures, particularly MML and LDL, as well as standardized measures of tinnitus spectra [14, 20] which can obviate clearly MML and LDL, as well as standardized measures of tinnitus spectra [14, 20] which can obviate loudness recruitment are potential contributing factors to such measurements in tinnitus. Systematic variation of trained frequencies between groups or within individuals is highly desirable, including untreated control conditions. Such evidence is needed to determine whether auditory training induces specific changes in tinnitus, or whether it instead amplifies the nonspecific effect of procedures common to all tinnitus therapies. Neural correlates offer the advantage of comparative immunity to response bias. Finally, care should be taken to specify clearly how many participants of the total recruited contributed to a data analysis, and why and when exclusions occurred. Progress toward an optimal auditory training treatment will be limited until replications are reported involving substantial sample sizes.

We also suggest that applications of auditory training will be enriched when we know more about how neural plasticity works in normal hearing individuals and in individuals with tinnitus. Current findings showing that passive exposure to sound is sufficient to remodel auditory representations in people with normal hearing could be good news for tinnitus, since compliance with treatment procedures may improve when performance requirements are minimal. The results reflect the propensity of the human auditory system to extract and represent the features of salient environmental sounds, regardless of behavioral response requirements. However, that passive exposure is sufficient does not preclude the possibility that active processing may yield more long-lasting outcomes [51].

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References


Keypoints

1. Tinnitus Retraining Therapy (TRT) is strictly based on the neurophysiological model of tinnitus.
2. Tinnitus is a phantom auditory perception, i.e. perception of tinnitus is not linked to any vibratory activity within the cochlea.
3. The model postulates that it is necessary to include interconnections within a network of systems in the brain in the study and treatment of tinnitus.
4. The auditory system, while needed for perception of tinnitus, is secondary for clinically relevant tinnitus (i.e. tinnitus which is bothersome to the extent of requiring treatment).
5. The limbic and autonomic nervous systems are the main systems responsible for negative tinnitus-evoked reactions.
6. Tinnitus is frequently accompanied by a decreased sound tolerance, consisting of hyperacusis and misophonia.
7. Hyperacusis results from an increased gain within the auditory pathways and is determined solely by physical characteristics of sound (i.e. its intensity and spectrum).
8. Misophonia results from enhanced functional connections between the auditory and the limbic and autonomic nervous systems, and reactions occur to specific patterns of sound, with the total spectral energy being secondary or irrelevant.
9. In misophonia, the meaning of sound and an individual’s past history of encountering it is crucial, with the auditory characteristics of the sound playing a secondary role.
10. There are two loops in network processing tinnitus signal:
   a. High loop, which involves cognitive processing of the signal and which is dominant at the initial stages of tinnitus.
   b. Low, subconscious loop, which appears to become dominant in chronic tinnitus.

Connections within the neural networks that are involved in the adverse effects of tinnitus are governed by the principles of conditioned reflexes.

11. The primary goal of TRT is habituation of reactions evoked by tinnitus.
12. Habituation is initiated and further facilitated using the method of modified passive extinction of the conditioned reflexes and involves:
   a. Teaching/counseling aimed at reclassification of the tinnitus signal to the category of neutral stimuli.
   b. Sound therapy, which decreases the strength of the tinnitus signal by increasing the level of background neuronal activity in the auditory system achieved by providing an enhanced sound background.
13. Habituation of perception happens automatically once sufficient level of habituation of reactions is achieved.
14. Decreased sound tolerance must be treated concurrently with tinnitus.
15. Different protocols must be used for hyperacusis than for misophonia.
16. A specific variant of treatment, related to classifying a patient to one of 5 categories, is determined by the following factors:

a. Impact of tinnitus on patients’ lives and/or duration of clinically significant tinnitus.

b. The presence of hyperacusis.

c. The presence and significance of hearing loss.

d. Prolonged exacerbation of tinnitus/hyperacusis by sound.

17. Misophonia is treated independently by specific protocols concurrently with tinnitus, hyperacusis, and hearing loss.

18. Results from many centers have confirmed the effectiveness of the tinnitus retraining therapy (TRT) for tinnitus, reporting a success rate of more than 80%.

19. Specific studies performed to assess the stability of improvement after 3 and 5 years revealed that improvement continues to be present with patients over time. These studies show a trend of continuing improvement, even after ending the treatment.

20. Prevention of clinically significant tinnitus or potential worsening of already existing tinnitus could be achieved by:

a. Avoidance of silence and providing enriched sound environment.

b. Avoidance of negative counseling and providing proper information in advance.

21. Certain populations are at high risk of developing clinically significant tinnitus, such as military personnel, police officers and firefighters, and patients who are going to have ear-related surgery. Providing them with a proper short informational session about tinnitus would significantly decrease the risk of developing bothersome tinnitus. People will still hear tinnitus, but it will not be a problem for them.

**Keywords**  Tinnitus • Habituation of reaction • Habituation of perception • Conditioned reflexes • Phantom perception • Retraining • Neurophysiological model of tinnitus • Sound therapy • Counseling • Teaching

**Abbreviations**

TRT  Tinnitus retraining therapy

LDL  Loudness discomfort level

**Outline of the Concepts Presented in this Chapter**

The theoretical ideas and the description of treatments presented in this chapter propose a different view on the phenomenon of tinnitus and its treatment than those found in the majority of other published hypotheses about tinnitus and descriptions of treatments. Therefore, definitions of tinnitus and decreased sound tolerance (hyperacusis and misophonia), as used in this chapter, will be provided first, followed by a brief outline of the main concepts of the neurophysiological model of tinnitus [1]. Tinnitus Retraining Therapy (TRT) is strictly based on this model and is one of several potential implementations of therapies aimed at habituation of tinnitus. Furthermore, as it is argued in this chapter, tinnitus should not be treated alone, but as one of the components of a more general dysfunction of the auditory system (including hearing loss and decreased sound tolerance), which needs concurrent treatment. Furthermore, the emphasis is on dynamic interaction of the auditory system with other systems in the brain, which is governed by principles of conditioned reflexes, and the role of subconscious pathways is stressed as well. The main goal of TRT is habituation of negative reactions evoked by tinnitus, with habituation of perception occurring as the subsequent, but inevitable, process.

**Definitions of Tinnitus and Decreased Sound Tolerance**

Tinnitus is defined as a phantom auditory perception, namely perception of sound without corresponding vibratory, mechanical activity in the cochlea [1, 2]. This perception is absolutely real and can be compared to phantom pain (see Chaps. 14 and 15) and the phantom limb phenomena. There is a tinnitus signal in the form of neural activity somewhere in the brain that is perceived as a sound, thus tinnitus. It is not known exactly where in the brain this occurs, but some studies indicate
that the secondary auditory cortex plays an important role in this respect. Understanding the phantom aspect of tinnitus is fundamental for understanding the interaction of tinnitus with external sounds and thus is the basis for the different forms of sound therapies currently in use. Problems can arise from misunderstanding of the role of external sounds on tinnitus, such as when the suppression of tinnitus perception by external sound is called “masking” [3]. Masking represents interaction of two traveling waves at the basilar membrane of the cochlea and therefore exhibits a “V-shaped” masking curve and depends on the phenomenon of the critical band (i.e. it is impossible to mask one sound by a second if there is a sufficient frequency difference between the two sounds). None of these phenomena exists in connection with tinnitus, which can be equally easily suppressed by sounds from wide range of frequencies [4]. Obviously, it is possible to interact with the tinnitus signal, including suppression of the perception that it leads to, but the mechanism is one of interactions between sound-evoked neural activity and the tinnitus-related neuronal activity. Furthermore, it is possible to decrease the strength of the tinnitus signal by increasing the general level of sound-evoked neuronal activity, and thus by decreasing the difference between the tinnitus signal and the background neuronal activity.

Consequently, the author is against classification of tinnitus into “subjective” and “objective” tinnitus and instead supports using the term “somatosound” in place of “objective tinnitus” as well as reserving the term “tinnitus” for what other authors have referred to as “subjective tinnitus.” This terminology is used in this chapter.

Tinnitus is frequently accompanied by decreased sound tolerance [5–7]. It is possible to identify two components of decreased sound tolerance, hyperacusis (see also Chap. 3) and misophonia (see also Chap. 4) [7–9]. Results from several centers show that about 25–30% of individuals with tinnitus also have hyperacusis. Our results from the Emory Tinnitus & Hyperacusis Center showed that out of 149 consecutive patients, 66% required treatment for decreased sound tolerance and 33% required treatment for hyperacusis while 57% required treatment for misophonia [8]. Most patients with decreased sound tolerance have both hyperacusis and misophonia, contributing to a decreased sound tolerance to different degrees. While these two components evoke a similar extent of behavioral reaction, there are significant differences in the categories of sound which trigger hyperacusis compared with misophonia, and the physiological mechanisms and treatments of hyperacusis and misophonia are distinctively different. Treatments that are effective for hyperacusis are not effective for misophonia, and treatments for misophonia have only limited impact on hyperacusis.

It is characteristic for hyperacusis that the reaction depends exclusively on the physical characteristics of a bothersome sound, such as its energy and frequency spectrum. The meaning of a sound and an individual’s past history are irrelevant. For example, a person may have a strong negative reaction to speech sounds. If this speech signal is recorded, the spectrum is determined, and then the sound is re-synthesized from individual frequencies with randomly assigned phases, this procedure will yield a sound with the identical energy spectrum as the original speech sound; however, this sound will be perceived as a noise without any meaning. A person with hyperacusis will react in the same manner to both such sounds; it is irrelevant whether this sound is familiar to the person or being encountered for the first time. The environment in which this sound is presented (e.g. doctor’s office, home, part of a sound track of a favorite movie) will not affect how a person with hyperacusis reacts to the sound. People with hyperacusis have a tendency to react stronger (or have lower sound tolerance levels) to sound of higher frequencies (e.g. sound of a metal spoon hitting china, washing machines with clicking plates), reflecting the general tendency of high-frequency sound being more bothersome even for individuals who do not have hyperacusis.

It is proposed that the neural mechanisms of hyperacusis involve abnormally high amplification within the auditory system, with only secondary activation of other centers of the brain responsible for negative reactions (i.e. the limbic and autonomic nervous systems). In other words, the activity that occurs within the auditory pathways after stimulation by 80 dB HL sound in a person with hyperacusis would be similar to that occurring in an individual who does not have hyperacusis and is exposed to a much louder sound such as a sound of 120 dB HL. Studies in animals support proposed mechanism [10, 11]; however, lack of an animal model of hyperacusis hinders researchers from performing more specific studies.

It is characteristic for misophonia that the adverse reactions occur due to specific patterns of sound, with the sound’s spectrum being secondary or irrelevant. The meaning of a sound and the past history of an individual encountering it is crucial. Sounds, which in the past have been associated with something negative (e.g. discomfort, pain, or other situations associated with
strong negative emotions), will trigger misophonic negative reactions. Basically, the mechanism of misophonia involves the creation of a conditioned reflex linking specific patterns of a sound to negative reinforcement. Significant hyperacusis, even present for a short period of time, will automatically create misophonia, because exposure to the sound will create discomfort/pain and it will consequently provide the negative reinforcement associated with the sound. Once this reflex is created, it will persist, even when hyperacusis ceases to exist.

For example, in a situation such as the one described above, a person with misophonia may react very strongly to normal speech but show no reaction to re-synthesized speech sounds that seem like meaningless noise. People who have misophonia may exhibit strong reactions to soft sounds (e.g. sounds of eating or speech of certain people) while not having problems with even very loud sounds. Many individuals with misophonia react to sounds “louder than” a certain level, yielding Loudness Discomfort Levels (LDL) determined for pure tones following the shape of audiogram and being better for frequencies where hearing loss exists [8, 12] (see Chap. 4).

The auditory system is perfectly normal in persons with pure misophonia; however, selective connections from the auditory system to the limbic and autonomic nervous systems for specific patterns of sound are abnormally activated or enhanced. Functional properties of these connections are governed by principles of conditioned reflexes. Consequently, the strength of the reactions they cause depends on the strength of the reinforcement, and the sound level plays a secondary role.

Sounds-evoking misophonic reactions do not have to be unpleasant on their own, but it will be sufficient that the individual who has misophonia identify sounds, exposure to which enhances tinnitus for some time. These sounds will be associated with an increased emotionally negative status, caused by enhancement of tinnitus which will be sufficient to create a conditioned reflex arc evoking misophonic reaction to these sounds, even at lower levels than that needed to increase tinnitus loudness.

**Physiological Basis for Tinnitus-Induced Negative Reactions**

It is crucial to distinguish between mechanisms involved in the generation of tinnitus perception and mechanisms involved in tinnitus-evoked negative reactions. Most individuals who have tinnitus are just experiencing a sound sensation, without any problems related to it. Only about 20% of people with tinnitus have negative reactions evoked by tinnitus (see Chap. 5). It is interesting that the psychoacoustic characteristics of tinnitus in these two subpopulations are undistinguishable and not related to the severity of the tinnitus as it is experienced by the people who have the bothersome tinnitus.

These observations indicate that there are different mechanisms involved in the generation of the neural signal that causes tinnitus perception, and other mechanisms responsible for evoking negative reactions to this signal. Recognition of this distinction is important, and by aiming treatments at the mechanism of tinnitus-induced negative reactions, it should be possible to remove the problems of the tinnitus without trying to remove tinnitus perception. In the past, most research and treatment attempts were aimed at removing, or at least decreasing tinnitus perception. These approaches were not particularly successful and so far we do not have any reliable method that would make it possible to achieve this goal. Notably, decrease of tinnitus perception does not automatically translate into decrease of tinnitus severity, and actually there is no relation between tinnitus loudness match and the perceived severity of the tinnitus [13].

Analysis of the negative effects of tinnitus on individuals provides information about which system in the brain may be involved in this process. It is possible to distinguish between two main categories of negative effects: (1) physiological responses to tinnitus (e.g. anxiety, depression, sleep problems, increased stress level) and (2) behavioral responses and consequences (e.g. attention and concentration problems, decreased joy of life, and affected life activities such as social interactions, work impairment, family problems). Two major systems in the brain are involved in generating the negative effects of tinnitus, namely the limbic and autonomic nervous systems, which interact with many other systems such as the prefrontal cortex, thalamus, reticular formation, and cerebellum playing some role as well. It has been proposed that it is necessary to include these systems in the analysis of the generation of tinnitus and a person’s reaction to the tinnitus (see Chaps. 20 and 21) and its treatment [1, 12].
The Neurophysiological Model of Tinnitus

The basic concept of the neurophysiological model of tinnitus is that it is necessary to include variety of systems in the brain in study and treatment of tinnitus [1, 12] (see also Chaps. 20 and 21). The auditory system, while needed for perception of tinnitus, plays a secondary role for clinically relevant tinnitus (i.e. tinnitus which is bothersome to the extent of requiring treatment). In the past, studies and treatments of tinnitus have tended to be cochleocentric. The neurophysiological model of tinnitus proposed earlier [1] and outlined here shifts the attention away not only from the cochlea but also from the auditory nervous system. The main focus of the model can be envisioned in a form of a diagram, first published in late 1990s [14] (Fig. 73.1). This concept is currently generally accepted and it is believed that any valid neurophysiological model of tinnitus must include several different systems in the brain to represent mechanisms of tinnitus and to be useful in the treatment of tinnitus (see also Chaps. 20 and 21).

According to their model, the tinnitus signal – the generation of which is typically linked to the periphery of the auditory system – is detected and processed by subconscious centers of the auditory pathways and finally interpreted at the highest level of the auditory system (probably the secondary auditory cortices). If a person just perceives tinnitus without having a negative reaction induced by it, the tinnitus signal may be constrained within the auditory pathways. If, however, this activity spreads to the limbic and autonomic nervous systems by activation of specifically the sympathetic part of the autonomic system, it evokes several negative reactions such as annoyance, anxiety, and panic and triggers survival reflexes resulting in a decreased ability to enjoy life activities. This last mentioned effect has a profound impact on a person’s life by depriving an individual of positive aspects of life which may push a person into depression [12] (and see Chaps. 62 and 63). The model shown in Fig. 73.1 has been described in detail already [15–23] and only the main aspects are outlined in this text.

Two Loops

All the systems in the brain are interconnected and work in the dynamic balance scenario, i.e. if a connection is frequently active it becomes stronger, if it is not activated it gradually becomes weaker (see also Chaps. 12, 13, and 20). This feature is reflected in the diagram of the model (Fig. 73.1), with the main systems mutually interconnected. It was postulated that tinnitus as a problem results mainly from over-activation of the

![Fig. 73.1 The neurophysiological model of tinnitus](image-url)
sympathetic part of the autonomic system [1, 19, 22]. It is, therefore, important to analyze pathways involved in tinnitus-related activation of these systems. Firstly, it should be noted that continuous activation of the connections illustrated in Fig. 73.1 causes their strengthening and yields stronger activation of the limbic and autonomic nervous systems by the same tinnitus signal according to the general rules of neural plasticity. Secondly, increased activation of the limbic and autonomic nervous systems occurs via reciprocal connections (feedback) and causes increased activity in the system from which the initial signal was coming. For example, autonomic system activation via a backward feedback can increase the activity in the limbic system, in cognitive brain areas, as well as auditory system. Therefore, the term “loop” is used instead of the term “connections” to emphasize the feedback aspects of the interaction between the different systems.

The tinnitus signal activates the limbic and autonomic nervous systems via two such loops. The upper one (“high loop”) (named the “high route” by LeDoux [24], see Chap. 8) involves conscious areas of the cerebral cortex, involves perception, evaluation, verbalization, conscious associations, and fears. This loop is crucial in the initial stage of developing clinically significant tinnitus. The second, lower (“low loop”, named the “low route” by LeDoux [24], see Chap. 14), involves subconscious centers in the brain. It branches from the auditory system at the level of extralemniscal subnuclei of the medial geniculate body, reaching the lateral nucleus of amygdala and, via other parts of the limbic system, reaches centers of the autonomic nervous system (see Chaps. 8, 12 and 21). Documented connections link the amygdala with the inferior colliculus, and therefore both connections (from auditory system to limbic system and back) are included in the diagram. The importance of these connections described in the model has been documented [25–27].

Both high and low loops contribute to the final activation of the autonomic nervous system and the negative reactions evoked by tinnitus. High loop is dominant in the acute stage of tinnitus development, but once tinnitus reaches a chronic stage, the subconscious becomes more important or even dominant (see Chap. 8). The analysis of results from over 300 patients with chronic tinnitus revealed that the proportion of time when patients are aware of tinnitus and subjectively ranked tinnitus loudness does not contribute significantly to tinnitus severity [28]. These results argue strongly against the dominant role of the conscious, high loop, because then tinnitus awareness and tinnitus loudness would be expected to be highly significant factors. These findings have a profound implication on tinnitus treatment.

**Conditioned Reflexes**

The connections between the brain systems involved in processing the tinnitus signal are governed by principles of conditioned reflexes. The tinnitus signal in the auditory pathways acts as a conditioning stimulus, which, via one or more reflex arc, activates the limbic and autonomic nervous systems and thereby evokes negative reactions. Several different scenarios may create these conditioned reflexes. The most common is the situation of “negative counseling,” i.e. a person is told something which links tinnitus with a threatening, unpleasant, or dangerous situation such as “nothing can be done, you will have tinnitus up to the end of your life, you need to learn to cope with it, and we need to do a brain scan just to eliminate the possibility of a brain tumor.” An extreme example of such negative counseling is when a patient is told that he/she has tinnitus because “he/she has a bad brain.” The negative counseling provides a reinforcement which creates a conditioned reflex arc causing the tinnitus signal to subsequently evoke strong reactions of the limbic and autonomic nervous systems, causing physiological and behavioral reactions.

Another common scenario occurs when a person with tinnitus is under strong emotionally negative stress, such as during retirement, divorce, or from non-related health problems. Indeed, a study ranking ordered factors present when a person’s tinnitus became a clinical problem revealed that there was no auditory-related factor in the top 10 most frequent situations [29]. While noise exposure is regarded a frequent cause of the appearance of tinnitus perception, it is not the case for emergence of tinnitus as a problem. Non-bothersome tinnitus may be present for years, and only when it becomes associated with something negative does it become a problem. It should be noted that no causal link is necessary for the creation of a conditioned reflex of any kind, but a close temporal association of a conditioning signal and reinforcement is sufficient to create the reflex.
The neurophysiological model described above and presented in simplified form in Fig. 73.1 predicts that a combination of rapid appearance of tinnitus together with high-level emotional stress is particularly effective in evoking clinically significant tinnitus. Indeed, bothersome tinnitus is evoked typically as a result of sudden hearing loss or when tinnitus starts at a specific time when a person is in the state of highly negative emotions due to sudden hearing loss. Consequently, clinically significant tinnitus can be expected to be more prevalent in professions where there is a combination of a high level of noise, particularly impulsive noise (e.g. gun fire) with a high level of negative emotional stress. Policeman, firefighters, and soldiers are typical examples of members of such professions.

This prediction has been confirmed by the fact that tinnitus occurs in a high rate (49%) of soldiers returning from Iraq and Afghanistan who were exposed to blast noise, the occurrence of which is even higher than the reported proportion of soldiers with blast-induced hearing loss (25%) (see Chap. 67). This unfortunate issue has significant financial connotations, as the American Veterans Administration spent $1.1 billion in 2009 (doubling from $540 million spent in 2006) on compensation for tinnitus alone, with the expected compensation for tinnitus reaching $2.3 billion by 2014 [30]. Since impulse noise can evoke hyperacusis and misophonia, in addition to tinnitus, adding significant problems for the Veterans health care system that must be taken care of in the near future and which will persist for many years to come (untreated clinically significant tinnitus tends to be stable for many years and untreated misophonia tends to worsen with time).

Once the reflex is established, a negative reaction can be evoked without a negative reinforcement, which means that while general health may improve and work problems may be resolved, a person’s tinnitus will keep evoking negative reactions. One of the reasons is that a tinnitus-evoked negative reaction acts as the reinforcement to the reflex arc that has been created and which causes these negative reactions. This aspect of tinnitus explains the low rate of spontaneous recovery, since clinically significant tinnitus is constantly present and that it evokes constant negative reactions, passive extinction of this reflex will not occur and it may actually cause further reinforcement of the reflex arc that causes the negative reactions.

**Tinnitus Treatments**

There are different methods in use for treatment of tinnitus, and before discussing TRT, it may be useful to briefly discuss some of these other main treatments for tinnitus. Traditionally, the goal has been to eliminate the tinnitus source and tinnitus perception, thus, aiming at achieving a cure of the tinnitus. So far, however, this goal is rarely achieved. Many treatments, typically aimed at the cochlea by delivering drugs directly to the cochlea or through the middle ear, have been tried, and some studies of the outcome of such treatments are currently in progress. Another traditional approach for treatment of tinnitus has been aimed at eliminating tinnitus perception. Suppression of tinnitus perception by external sound, labeled “masking,” has been widely promoted. This approach has not been as successful as hoped, with reported effectiveness from zero [31] to 60% [32]. Recently, “masking” has been re-defined as use of any sound which provides some immediate relief for tinnitus [33]. This approach has shown some effectiveness [34, 35], but it is not clear if it is better than any other type of sound therapy (see Chaps. 72, 74 and 75).

Different investigators have used the term “masking” in different ways to describe tinnitus suppression. Auditory masking results from interaction between two traveling waves on the basilar membrane of the cochlea, and as such exhibits phenomena of “critical band” and “V-shaped suppression curve.” None of this is true for tinnitus, it is possible to suppress tinnitus perception equally easy by sound of any frequency, and there is lack of significant dependence on the intensity of the sounds needed to suppress tinnitus from a frequency of the tone [4]. These findings support the hypothesis that tinnitus is a phantom auditory perception without any correspondence to the vibratory activity within the cochlea.

Another approach to suppress tinnitus perception that has been described makes use of electrical stimulation of the cochlea/auditory nerve (see Chap. 77) or, recently, electrical stimulation of the auditory cortex [36–39] (see Chap. 90). In the case of the auditory cortex, in addition to direct electrical stimulation, Transcranial Magnetic Stimulation (TMS) has been used [40–42] (see Chap. 88). In TMS, impulses of a very strong magnetic field are applied locally to the skull and the induced electrical current stimulates the cerebral cortex. All these attempts to treat patients with
Tinnitus were partially successful, with an average rate of about 50%. These methods are now under further investigation.

Different classes of treatment have been aimed at decreasing tinnitus-evoked reactions by improving coping strategies, modifying an individual’s thinking about tinnitus, or by using psychotropic drugs to attenuate activity of the limbic system [12, 43–46] (see Chaps. 78 and 79). Psychological approaches have shown effectiveness in the range of 50% (see Chaps. 71 and 72), while so far none of the drugs tested have shown significantly positive effects.

Last, but not least, a variety of sound therapies based on the concept of attenuating tinnitus or making it less noticeable have been described [33, 47–51] (see Chaps. 72, 74, and 75). These treatments have shown some effectiveness, but for most of these methods lack of systematic, independent studies have made it impossible to accurately assess their efficacy. Recently, the concept of using sounds where the energy at frequencies around the pitch of a person’s tinnitus were eliminated has been reintroduced [52]. The use of such sounds is based on the hypothesis that utilizing the mechanism of lateral inhibition in the auditory cortex would suppress tinnitus. Lateral inhibition, which occurs commonly in the brain and reflects situation that stimulation of one neuron, is frequently accompanied by inhibition of nearby neurons [12, 53]. In the case of the auditory system, which exhibits tonotopic organization, stimulation with a given frequency can inhibit neurons that respond best to nearby frequencies. Specifically, in case of tinnitus, it has been postulated that by removing the music’s frequencies around a person’s tinnitus pitch, the neurons in this range will be inhibited due to activation of neurons which respond best to nearby frequencies (see Chap. 75).

Treatments Aimed at Habituation of Reactions to Tinnitus and Its Perception

Above Outlined treatments aim at removing or attenuating source of tinnitus signal, or at alleviation reactions evoked by tinnitus. The neurophysiological model of tinnitus suggests another possible direction for treatment, namely the possibility of blocking the spread of the tinnitus signal to other than auditory regions of the brain, particularly to the limbic and autonomic nervous systems. If such treatment is successful, a person may still perceive their tinnitus, but tinnitus will not bother her/him. This process is called habituation of tinnitus-evoked reactions. Notably, once sufficient level of habituation of reactions is achieved, habituation of perception automatically follows and a person is aware of tinnitus for smaller and smaller proportions of time as the brain automatically habituates all stimuli that are not important [12] (see Chap. 20). As a result, an individual with tinnitus changes from being a sufferer to becoming a member of the population of people with tinnitus who experience it, but are not bothered by it. It is important to note that this treatment will not work when attempts have been made to first induce habituation of perception. Any method yielding habituation of tinnitus may be labeled Tinnitus Habituation Therapy [54].

According to the model outlined in Fig. 73.1, habituation of reactions will occur when all connections carrying the tinnitus signal to the limbic and autonomic nervous systems are attenuated and preferably blocked (Fig. 73.2). Proper counseling can relatively easily modify the functional connections from the cognitive areas down to the limbic and autonomic nervous systems. Retraining subconscious connections between the auditory system and the limbic and autonomic nervous systems, however, is more complex and difficult to accomplish. Counseling alone will not work, and it is necessary to utilize methods appropriate for retraining the conditioned reflexes.

From the time of Pavlov, it is well known that a conditioned reflex created by exposing a person many times to a sensory stimulus spontaneously undergoes extinction if reinforcement is not given (e.g. using the classical example of the Pavlovian dog, the bell keeps ringing but food is no longer given). This process is known as passive extinction of conditioned reflexes or habituation of reaction [12, 55, 56] (see also Chap. 87). While effective in many situations, this technique cannot be applied in its original form to tinnitus, because the tinnitus signal and its perception are constant and cannot be eliminated. Reinforcement is provided by reactions of the limbic and autonomic nervous systems and, consequently, is constant and cannot be blocked. To solve the problem, the author proposed a modified version of passive extinction of conditioned reflexes using a simultaneous decrease of the sensory signal and reinforcement with these changes maintained for some time (corresponding to the ringing of the Pavlov bell being softer and less food given). This process
should be effective, but it requires more time than classical passive extinction. In the case of tinnitus, it requires that the strength of the tinnitus signal is decreased, with a decrease of the strength of negative reactions happening at the same time. These interventions must be carried on for some time to obtain good results.

**Tinnitus Retraining Therapy and Its Clinical Goals**

TRT is a specific implementation of general Tinnitus Habituation Therapy, which utilizes counseling to decrease tinnitus-evoked reactions and sound to decrease the strength of the tinnitus signal. The primary goal of TRT is to achieve habituation of tinnitus-evoked negative reactions and remove the effect of tinnitus on patients’ lives. As pointed out above, once habituation of reactions has been at least partially achieved, habituation of perception automatically occurs without the need of any additional action. At the end of a successful treatment, people are not bothered by tinnitus (or bothered very little), even when perceiving it, and tinnitus has no impact on their lives. Additionally, their awareness of tinnitus typically drops to 5 or 10% of their waking time.

Counseling: Habituation cannot be achieved to stimuli indicating danger or are threatening and is achieved with difficulty to stimuli that evoke strong emotional reactions (negative or positive). Therefore, the primary role of counseling in TRT is to achieve reclassification of tinnitus to a category of neutral stimuli. This is achieved by intensive teaching about mechanisms of the tinnitus origin and its benign nature (as perception), which nevertheless may evoke strong negative reactions affecting patients’ lives. Patients typically have many incorrect concepts about tinnitus and, at the same time, tinnitus remains a mystery for them. Therefore, demystification of tinnitus and providing patients with solid knowledge is important.

The modified method of passive extinction requires a decrease of the strengths of both the activation of the limbic and the autonomic nervous systems and the tinnitus signal. It is impossible to modify the activity of the subconscious low loop that connects information from the thalamus to the limbic system directly [12] (see Chap. 8), but it is possible to attenuate and finally remove the contribution of cognitive components (the high loop). As both the high and low loops contribute to the final activation of the limbic and autonomic nervous systems, it is possible to achieve a decrease of activation of the limbic system by removing or at least decreasing the transmission and processing of tinnitus signals in the high loop, thus removing its contribution.
By reclassifying tinnitus to the category of neutral stimuli, showing its benign character, providing explanation about its origin and mechanisms, pointing out that the patients have “a proper reaction but to an improper stimulus,” and answering questions, etc., it is possible to eliminate transmission of the tinnitus signal in the high loop in a relatively short time.

Another, more general mechanism of attenuation of the effects of the high loop in tinnitus is based on the fact that people react stronger to unknown dangers than to even significant dangers which are known. Therefore, once patients are able to predict the behavior of their tinnitus (e.g. typical increase of the tinnitus when in quiet places or in a stressful situation) and their own reactions to it, they may still be annoyed and bothered by their tinnitus, but to a smaller extent.

Even complete abolishment of tinnitus signal processing and transmission in the high loop is not sufficient to remove negative reactions evoked by tinnitus because the low loop remains fully active and unchanged.

The negative reaction to a person’s tinnitus will undergo gradual extinction as the result of this modified method of passive extinction, but the process will be slower than eliminating processing and transmission of tinnitus signals in the high loop. Nevertheless, on average, patients show clear improvement after just one month of TRT treatment. This observation has been recently collaborated by reports from other centers [57, 58].

The counseling/teaching session is crucial for achieving high effectiveness of the TRT treatment. Without it, sound therapy will have some positive effect, as well as using some general counseling, but the effectiveness of such treatments is clearly lower [34] because they will not eliminate processing and transmission of the tinnitus signal in the high loop. Nevertheless, on average, patients show clear improvement after just one month of TRT treatment. This observation has been recently collaborated by reports from other centers [57, 58].

Sound therapy: In principle, many different methods may be used to decrease the tinnitus signal (e.g. drugs, electrical stimulation, TMS), but in practice, the use of sound is simple and can be easily controlled and adapted to the needs of an individual patient (see Chap. 74). Sound therapy utilizes the principle that the strength of the neuronal signal within the brain is based on the contrast principle, thus the difference of the signal is from background sound or background neuronal activity. Therefore, the strength of the tinnitus signal can be decreased by systematically increasing the background neuronal activity within the auditory pathways. This can be achieved by enhancing background sound levels to which patients are exposed, thereby affecting the perception of the tinnitus.

Specifics regarding the implementation of sound therapy, including use of sound generators, combination instruments, and hearing aids, have been described in detail elsewhere [5, 8, 15, 59]. It is crucial to remember the basic rule for its successful implementation: “Never use sound as a part of sound therapy, which would create annoyance or discomfort for any reason.” Use of a sound which would evoke any negative reactions would activate the limbic and autonomic nervous systems, worsening the situation and making habituation more difficult to achieve. Other recommendations, which are helpful while less critical, include the use of sound enrichment preferably all the time, 24/7, use more than one type of sound source (e.g. sound generators and tabletop sound machines) and preferably use nature sounds in the background. Music is typically used in the basic protocol for misophonia [8, 12] (see also Chap. 75).

**Decreased Sound Tolerance**

Tinnitus is typically accompanied by decreased sound tolerance, both hyperacusis and misophonia (see also Chap. 3). Results from many centers indicate that hyperacusis coexists with tinnitus in 25–30% of people who seek help for their tinnitus. Our own data indicate that misophonia is present in about 60% of the patients we treat [8]. Proper diagnosis and treatment of decreased sound tolerance and its components is crucial for successful outcome of tinnitus treatment. Hyperacusis is relatively easy to treat with the desensitization protocol [12], and typically it can be attenuated or eliminated within a couple of months of treatment. Treatment of misophonia is much more complex and lengthy and requires specific protocols. This reflects the fact that the same neuronal networks are involved in tinnitus and misophonia, and consequently, treatment of misophonia takes a similar amount of time as treatment of tinnitus (see also Chap. 20). Methods used for successful treatment of hyperacusis are not effective for treatment of misophonia.

The situation is further worsened by the fact that the presence of tinnitus frequently induces or enhances
misophonia, as patients dislike and start to avoid sounds in general, which makes their tinnitus worse (or patients think that this is happening). Last, but not least, misophonia tends to trigger the tensor tympani syndrome (fullness in the ears, pain, feeling of pulsation, vestibular problems, headaches, etc.) [60], which may become a significant, or even a dominant problem.

Severe decreased sound tolerance is more debilitating than severe or even catastrophic tinnitus and can totally disable people. Without adequate treatment of decreased sound tolerance, and particularly misophonia, the effectiveness of tinnitus treatment becomes substantially decreased.

On the positive side, TRT is very effective for treatment of both hyperacusis and misophonia, and it is possible to achieve a cure in most patients, which means total elimination of hyperacusis and misophonia. Another positive aspect is that after successful treatment of misophonia, the tensor tympani syndrome disappears as well, and that tinnitus, if still bothersome, typically also improves.

Outline of Treatment by TRT

Patient Evaluation

In evaluation of patients with tinnitus and decreased sound tolerance, the detailed initial interview is crucial for the diagnosis. We are using a structured interview for initial and follow-up visits conducted with help of specific forms [61, 62] (see Appendix A). While information provided by this interview gives good insight into many aspects of tinnitus including its severity, the Tinnitus Handicap Inventory (THI) is used as well to assess tinnitus severity in a more formal manner [63, 64].

An audiological evaluation includes a pure-tone audiogram (up to 12 kHz), determination of pure-tone Loudness Discomfort Levels (LDL) (measured for all frequencies evaluated in the audiogram), evaluation of speech discrimination, and high-frequency resolution Distortion Product Otoacoustic Emission (DPOAE) (10 points per octave, frequency range of f2 from 1 to 10 kHz), all are tests that are needed for evaluation of patients for treatment using TRT. Audiogram and speech discrimination scores show the patient’s hearing ability. LDLs are crucial in assessing decreased sound tolerance, and DPOAE is extremely helpful during counseling, particularly when Discordant Dysfunction Theory (DDT) is used [1, 12, 65]. This theory postulates that the tinnitus signal originates from the regions of the basilar membrane where there is decreased activity of Outer Hair Cells (OHC) while Inner Hair Cells (IHC) are functional. Measurements of tinnitus pitch and loudness match are performed as well, but these results have no impact on diagnosis or treatment. Determination of Minimal Masking Levels (MML) is done for research purposes.

The acoustic reflexes and reflex decay are not a part of routine evaluation. They are not necessary or sufficient to determine presence of vestibular schwannoma (if it is suspected). Since most patients with tinnitus have decreased sound tolerance, the exposure to loud sounds (which are necessary for testing the acoustic middle-ear reflexes) may cause worsening of the symptoms and make subsequent interaction with patients more difficult.

Diagnosis

The following information is used for diagnosis and categorizing the patients’ category and variant of TRT treatment in TRT which should be used.

Impact of Tinnitus on Patients’ Lives and/or the Duration of Clinically Significant Tinnitus

This information provides insight regarding the strength of neuronal connections in the network processing the tinnitus signal. If tinnitus has a low severity, the connections are likely to be weak because it has been shown that tinnitus severity does not depend on the loudness to which a patient matches his/her tinnitus, and it is as well established that the strength of reactions that occurs through activation of conditioned reflexes depends primarily on the strength of the reinforcement; the strength of the conditioned stimulus plays a limited role. Therefore, weak reactions indicate weak strength of the connections in the neural networks involved in causing the tinnitus-evoked reactions. If an individual’s tinnitus is very recent (a few weeks), these connections have not had enough time to become permanent, which would make it easier to modify them
(see chapter neural plasticity). In this connection, it should be noted that the duration of the symptoms regards clinically significant tinnitus and not the duration of the tinnitus perception as such. Tinnitus may be perceived for many years, without bothering a person, before suddenly becoming a problem.

Hyperacusis

Presence of hyperacusis imposes some constraints on the use of sound therapy, including the allowed category of protocols for misophonia. The differentiation between hyperacusis and misophonia is complex, but a detailed interview, together with a comparison of the shape of a person’s audiogram and that of the LDLs, typically provides enough information to make this distinction [8]. A positive diagnosis of hyperacusis requires that the average LDL is less than about 90 dB. A low value of the LDL does not prove the presence of hyperacusis, however, because a low value of LDLs may be due to misophonia. The behavioral reactions evoked by hyperacusis and misophonia are identical, thus they cannot be used for differentiation either. Detailed interview with attention paid to sounds evoking negative reactions is necessary for differentiation of hyperacusis form misophonia and assign their relative contribution to decreased sound tolerance. If both hyperacusis and misophonia are present together and hyperacusis is treated successfully, but misophonia is not treated and disappears, misophonia typically may be enhanced, and at the behavioral level, no improvement in the patient’s condition is achieved.

The Presence and Significance of Hearing Loss

Approximately 70–80% of individuals with tinnitus have some degree of hearing loss, which is typically less than 60 dB HL in the frequency range up to 8 kHz. It is important to consider whether such hearing loss has any impact on a person’s everyday life (see Chap. 5). The same degree of hearing loss can be of large significance to one person (e.g. professions requiring communication in noise or musicians) while having no significance to another (e.g. farmers). A factor to consider when evaluating a person’s hearing is whether the hearing loss is accompanied by a “strain to hear” in everyday life because this increases the severity of tinnitus. Only after all these factors have been taken into consideration can a patient be classified as having hearing loss of significance for treatment of tinnitus. It should be noted that a patient’s subjective awareness of her/his hearing loss has little relevance because patients often do not acknowledge mild or even significant hearing loss.

Prolonged Exacerbation of Tinnitus/Hyperacusis by Sound

Over 50% of patients report that their tinnitus becomes worse for some time after exposure to sound (loud, moderate, or even soft). This time is in the range of minutes to hours for most individuals with tinnitus and typically affects hyperacusis or misophonia more than tinnitus. Some patients report that the effect persists through the next day, even after a good night’s sleep, or it may even last several days. This observation can be due to two scenarios and has profound impact on the diagnosis and treatment: (1) it may involve functional plastic changes in the nervous system that occurs as the “kindling” or “wind up” phenomenon (see also Chaps. 10 and 12) or (2) strong misophonia (see also Chap. 4). “Kindling” is a term from the field of epilepsy that describes the phenomenon that may occur when a weak stimulus that initially does not evoke a seizure evokes epileptic seizures after being presented repeatedly over several weeks.

The “Wind-up” phenomenon is a term from the field of chronic pain and describes the situation when the second presentation of a painful stimuli causes a stronger reaction than caused by the first presentation or when a stimulus presented for a limited time (e.g. a few minutes) is not inducing pain; when its duration is longer, it becomes painful (see also Chaps. 14 and 15). This is similar to what occurs when a sound, which initially is without any effect causes a worsening of tinnitus or hyperacusis after the sound has been presented for a longer period of time. These phenomena are particularly observed in people with certain medical problems, such as after head injury, brain surgery, Lyme disease, or symptoms associated with hormonal changes (for instance, during menopause).

In the past, it has not been appreciated that perception of prolonged sounds can often cause misophonia, causing elements of phonophobia to become
worse. People with tinnitus may become afraid that exposure to sound may cause their hyperacusis or tinnitus to become permanently worse and by paying extra attention to such problems and avoiding sounds (using ear protection) can enhance tinnitus and hyperacusis/misophonia or cause prolongation of an initial worsening. The experience of treating patients with tinnitus has shown that most incidences of worsening were due to the development of misophonia; only a few people have had indications that the worsening had a medical basis. Nevertheless, when medical reasons are reported by a patient, it should be considered.

Categories of Tinnitus

Five categories of patients are proposed on the basis of the factors listed above \[12, 22, 66\], and specific variant of TRT treatment has been associated with each one of these categories. These categories, listed below, provide general directions, and the borders between them are not sharp. While the categorization forms a continuum that provides some general guidelines that can help in avoiding some mistakes in the treatment, there is a certain degree of overlap between neighboring categories. For example, patients can be categorized as C 2/3 or 3/2 in case of coexisting hearing loss (category 2) and hyperacusis (category 3), depending on which problem is dominant. If ear-level instrumentation is used, it should be aimed at preserving or restoring symmetrical stimulation of the auditory pathways. Consequently, most of those who use instruments use these bilaterally, with exception being for cases with no hearing in one ear.

Misophonia may be present or absent in all these categories since treatment of misophonia is different from treatment of hyperacusis and tinnitus and can be conducted simultaneously with treatments aimed at the patient’s tinnitus and their hyperacusis. Consequently, misophonia is not included as a discriminating factor in the categorization. A detailed description of the categories and associated variants of treatments has been presented earlier \[7, 19\].

In the following, each one of the 5 categories is described as well as the methods in which patients with each category are treated using proper variant of TRT method.

Category 0

This category of patients is characterized by a low degree of severity (or hyperacusis) or a short duration of the problem. An abbreviated version of counseling is conducted providing basic information aimed at reclassification of tinnitus into a category of neutral stimuli, furthermore, following the principle that the problem should not be presented, so that it gives an impression that it could be worse than the patient reports (e.g., patients are not told that tinnitus or decreased sound tolerance can be debilitating and push them into depression or suicide!). All the remaining 4 categories have higher severity tinnitus and/or hyperacusis as well as potentially having tinnitus of a longer duration.

Treatment of patients with this category of tinnitus consists of providing basic information about sound therapy, with a short discussion about enrichment from environmental sounds by using sound machines producing nature sounds, or by using other sound sources. The benefit from the principle “Avoid Silence” during the treatment is pointed out. Ear-level instrumentations (e.g., sound generators) are typically not needed in this category, but such devices may anyhow be beneficial. However, as they are not essential for a successful outcome of the treatment and due to financial reasons, they are not recommended.

While an initial visit is typically sufficient to achieve noticeable improvement in this category of patients, a short follow-up visit or telephone call at 1, 3, and 6 months after treatment is worthwhile. Patients who are not improving should be reassessed and, if needed, have more extensive counseling, and recommendation of an ear-level instrumentation should be considered.

Category 1

Patients with this category of tinnitus have significant tinnitus, without hyperacusis, but misophonia may also be present. There is no significant hearing loss, and there is no sound-induced prolonged worsening of tinnitus (except when induced by misophonia). The treatment involves full counseling focused on the patient’s tinnitus, with omitted elements related to hyperacusis.

Sound generators are recommended as part of the therapy, providing well-controlled sound delivery. The sound level is typically determined by the level that evokes annoyance, and only in some patients it is
possible to reach the sound level where the patient can perceive their tinnitus and the external sounds as separate entities, but with both sounds start to mix or blend together (the “mixing point”).

Formally, this is the level where partial suppression (“partial masking”) starts to occur. Reaching the mixing point is not important for successful outcome of the treatment. In fact, pushing patients toward reaching the mixing point at the expense of going above the level evoking annoyance or discomfort is counterproductive and works against facilitation of habituation. Sound levels which are low should be avoided, however, because of the effect of stochastic resonance will enhance a person’s tinnitus and work against its habituation [67, 68]. Real Ear Measurements (REM) are highly recommended as a part of fitting an in-the-ear device and are repeated at follow-up visits.

**Category 2**

The characteristic feature of this category of patients is the presence of significant hearing loss as defined above. Full counseling is performed with stressing matters that are related to hearing loss. Combination instruments (a combination of an independently controlled sound generator and hearing aid in one shell) are preferable for sound therapy to be used in conjunction with enrichment of environmental sounds, as recommended for other categories. If such devices cannot be used due to technical or financial issues, the focus should be on achieving sufficient enrichment of background sounds, typically including the use of tabletop sound machines, with increased stimulation of the auditory system further enhanced by hearing aids. Fitting and use of hearing aids is specific for individuals with tinnitus and different than for people without tinnitus [59, 69].

Sound generators alone are not used for this category of patients, as they would make the understanding of speech even more difficult. Such devices would make tinnitus worse due to an increase in the strain to hear and understand speech.

**Category 3**

The characteristic feature of this category of tinnitus is a presence of significant hyperacusis that must be treated first. Full counseling is performed, stressing issues related to hyperacusis (e.g. both peripheral and central mechanisms controlling amplification within the auditory pathways are explained). Sound generators are always recommended for sound therapy in patients without hearing loss. Combination instruments are used in patients with hearing loss with stress in the initial stage on sound generators (and low amplification of hearing aid part) followed by second stage when amplification is increased. In both stages, sound generator and hearing aids parts are used concurrently. When combination instruments cannot be utilized, a two-stage procedure may be considered: the first stage with use of sound generators and in the second stage, they are replaced by hearing aids. In this scenario, patients need to be counseled properly to ensure that they expect and accept increased impairment of understanding of speech while using sound generators during the first stage of treatment. Patients should not use these devices when speech communication is essential. Another option is to make use of enrichment of sound background for treatment of hyperacusis before proceeding to the stage in the treatment where hearing aids are used. As hyperacusis is relatively easy to treat and the treatment is fast, this approach may be considered as well.

Combination instruments are the most versatile ear-level instruments for tinnitus treatment and, in theory, they can be used in about 80% of all patients. Technical limitations of currently available instruments as well as their high costs hinder their general use. Another significant aspect is that they require a specific fitting, and lack of proper theoretical and technical know-how by the people who do the fitting results in a high return rate of such instruments. In our experience, we have excellent results with the use of combination instruments and very few were returned.

It should be noted that the presence or absence of misophonia is irrelevant for treatment of patients with this category, which has hyperacusis present. If misophonia is present, it may be treated in the same way as described for the other categories of tinnitus, and for example patients with tinnitus and misophonia will be classified as category 1 with misophonia. Therefore, the presence of decreased sound tolerance is not sufficient to classify patient as belonging to category 3; presence of hyperacusis is required. If misophonia is present in this category, due to presence of hyperacusis, certain restrictions are imposed on the type of protocol that can be used for the treatment of misophonia. Note that the
basic features of the protocol for misophonia can be used even for patients with severe hyperacusis [8, 19].

**Category 4**

The characteristic feature of the tinnitus in this category is a prolonged exacerbation of the patients’ worst problem; typically, hyperacusis that may last at least after a good night’s sleep and does not result from misophonia. If there are medical problems involved that cannot be treated medically, such as from the effect of brain injury after car accidents, blast injuries from military operations, or brain operations, the treatment is highly individualized and difficult. Checking for the presence of Lyme disease may be worthwhile, because it has been reported that hyperacusis is present in 48% of Lyme disease cases [70]. If Lyme disease is the base of the problem, treatment for Lyme disease with antibiotics could be helpful even regarding the tinnitus.

**Results of TRT**

TRT works independent of the cause of the tinnitus, and the habituation of the reaction to the tinnitus occurs outside the central auditory pathways. Therefore, the etiology of tinnitus is irrelevant, and TRT can be successfully used for any type of tinnitus, e.g. bilateral, unilateral, continuous, or intermittent, as well as for somatosounds. This prediction from the neurophysiological model of tinnitus has been confirmed by results of clinical studies. The results of our past studies showed significant improvement in over 80% of the patients with noteworthy improvement as observed after about 12 months after the beginning of the therapy disregarding the etiology of tinnitus [23, 71]. Recent results of studies of over 300 patients treated in the Emory Tinnitus & Hyperacusis Center showed statistically significant improvement after only 3 months of TRT treatment, with further improvement occurring when the treatment was continued [15, 72].

Results of open studies reported from various centers using TRT also consistently showed significant improvement in over 80% of the patients who were treated [57, 73–87].

The results of a 5-year follow-up study showed that TRT had a high degree of effectiveness in treatment of patients with tinnitus and hyperacusis and that the improvement is persistent [88]. A recently published study evaluated the effects of 18 months of TRT treatment (and the following 18 months without continuing the treatment) [89]. Results immediately following the treatment show a high level of statistically significant improvement which persisted for the 18 months after the study’s treatment completion. Moreover, the proportion of patients reporting disappearance of their tinnitus-evoked difficulties while attempting to relax and concentrate and reporting problems with sleep, social interaction, and work increased continuously after treatment completion [89].

Of interest are also reports presenting results of a systematic randomized study, which showed that TRT is not only highly effective in general but is also effective for patients with severe symptoms typically reported to be particularly difficult to treat using other approaches [34, 35]. Interestingly, there was no indication of the results reaching a plateau after the 18 months the treatment, suggesting the possibility for achieving even better results with further continuation of the treatment.

**Prevention and Early Treatment**

Tinnitus and decreased sound tolerance present a big problem once they are established. Obviously, prevention of the occurrence of bothersome tinnitus, or treatment at the very early stage of tinnitus, would have a significant impact on the extent of problems created by tinnitus in the general population (see also Chap. 69). Unfortunately, tinnitus prevention is an area that has been largely ignored. The neurophysiological model as described earlier [1, 17, 19, 90–92] and outlined in this chapter provides guidelines for prevention of the appearance of clinically significant tinnitus and indicates how to achieve relief of problems related to tinnitus shortly after they appear [19].

**Avoidance of Silence and Providing an Enriched Sound Environment**

The vast majority of our patients at the Emory Tinnitus & Hyperacusis Center and at University of Maryland
in Baltimore describe the initial appearance of their tinnitus occurred during a period where they were in a silent environment. Experiments have documented that it is possible to evoke tinnitus perception in most people after a few minutes in a quiet environment [93–95]. If a person is in a negative emotional state while perceiving tinnitus, it may lead to development of conditioned reflexes producing tinnitus-evoked negative reactions. Exposure to sound, particularly by being in an environment enriched by natural sounds or sounds generated by tabletop machines, music players, etc., decreases the probability of bothersome tinnitus materializing.

The neurophysiological model of tinnitus developed by the author and described earlier [1, 17, 19, 90–92] predicts that if a person is exposed to an enriched sound environment shortly after the occurrence of tinnitus perception due to any reason (e.g. explosion or exposure to another damaging sound), it will decrease the probability of development of clinically significant tinnitus or, if it does occur, exposure to sound will increase the likelihood of habituation of the tinnitus. Neuronal connections responsible for the tinnitus-evoked reactions will then not have enough time to become permanent, and it is easier to retrain them if tinnitus is treated early after its occurrence (see Hebb’s principle in Chap. 12). This prediction has support from the results of both clinical studies and recent experiments on animals, showing that it is possible to prevent or reverse the reorganization of cortical maps that have been induced by sound overexposure by providing animals with an enriched sound environment [96, 97].

It is obvious that avoidance of excessive noise, which may cause damage to the cochlea or other changes in the auditory system (see Chap. 37), is recommended; however, recommendations given to patients with tinnitus frequently result in overprotection and have the opposite effect. The phenomenon of auditory toughening (i.e. increased resistance of the cochlea to damage and protection against loud, damaging sound offered by pre-exposure to moderate to loud sound) [98–100] (see Chap. 37), is not appreciated in general and even more in case of hyperacusis and tinnitus.

Sound exposure is necessary for keeping normal gain in the auditory nervous system; if it does not receive enough sound input, the gain increases and contributes to the development of tinnitus and/or hyperacusis. The experience from treating patients with tinnitus and animal experiments is that both overexposure to sound and overprotection from sound can be harmful. This message needs to be strongly emphasized in both public and professional health education. Exposure to appropriate forms of sound should be promoted as an integral part of our life that is essential for personal well-being.

Avoidance of Negative Counseling and Providing Proper Information in Advance

Negative counseling frequently provided by health care professionals, patient support groups, and the Internet can trigger mechanism that can create clinically significant tinnitus and make existing tinnitus worse. Health professionals should be alerted to the danger of such negative counseling that is offered to people with tinnitus. Instead, the general population should be educated with correct and basic knowledge about tinnitus, pointing out that there is much that can be done to alleviate the harm from tinnitus and that there is a high likelihood of successful treatment and of decreased sound tolerance (hyperacusis and misophonia). The most frequent issues related to negative counseling (e.g. “nothing can be done, let’s take a brain scan to exclude a brain tumor”) should be properly presented, so that the effect of negative counseling a person may receive is eliminated or at least prevented from having any profound effect.

Such education is particularly important for individuals who have a high risk of acquiring tinnitus, such as soldiers. For example, a huge amount of suffering and money would be spared if all soldiers going into combat situations undergo just one thoughtfully prepared 1–2-h educational session. Identification of other high-risk populations (e.g. police officers, firefighters, construction workers, and patients before any type of ear operation) and providing them with proper education would be highly beneficial as well.

Conclusions

TRT is a specific implementation of the Tinnitus Habituation Therapy, which utilizes teaching/counseling to reclassify tinnitus into the category of neutral
stimuli, and sound therapy to decrease the tinnitus-related neuronal activity (tinnitus signal) within the brain. As a result of TRT, habituation of both a person’s reactions evoked by the tinnitus and its perception occurs. TRT is strictly based on the neurophysiological model of tinnitus developed by the author [1, 19, 22] (outlined in Fig. 73.1), which stresses the necessity of including a network of interaction between many different systems in the brain in models of tinnitus and hyperacusis. From the beginning, the model stressed that the auditory system plays a secondary role [19] (see also Chap. 20). Emphasis instead is placed on structures involved in evoking tinnitus-induced negative reactions, mainly but not exclusively, the limbic and autonomic nervous systems.

TRT has been used clinically for treatment of tinnitus and decreased sound tolerance since 1988. The method of TRT underwent many modifications since its first description, and the method does not have a stagnant protocol but continues to evolve on the basis of information gathered from both treatment of patients and animal research findings. While the main features and assumptions of TRT remained the same, implementation of TRT has changed substantially regarding both the counseling part and sound therapy. In counseling, main changes included introduction of the concept of misophonia, the emphasis on conditional reflexes, subconscious processing of information, and on direct teaching about using a modified version of passive extinction of conditioned reflexes. The use of sound therapy has also undergone changes from the introduction of specific protocols for misophonia, changing the parameters for the sound stimulation from the use of levels, which could evoke annoyance or discomfort, to the use of lower levels. These modifications resulted in a significant reduction of the time needed to achieve improvement in the patients’ problems, from 1 year to 1 month.

Results from many tinnitus treatment centers show that TRT causes noticeable improvements or cures in and above 80% of patients with any type of tinnitus. Notably, refined counseling and sound therapy increased the effectiveness of TRT in treatment of decreased sound tolerance, so that it now becomes possible to achieve complete elimination (cure) in most patients.
# Appendix A: Forms for Structured Initial and Follow-up Interviews

## Form 1: Tinnitus/Hyperacusis initial interview form

<table>
<thead>
<tr>
<th><strong>TINNITUS / HYPERACUSIS</strong></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>INITIAL INTERVIEW FORM</strong></td>
<td></td>
</tr>
</tbody>
</table>

- **T&HC#:** 
- **tel:** 
- **e-mail:** 

<table>
<thead>
<tr>
<th><strong>Description of T sound(s)</strong></th>
<th><strong>Onset:</strong> Gradual / Sudden</th>
<th><strong>When</strong></th>
<th><strong>Frequency</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>RE / LE / Both / Head = &gt;</strong></td>
<td><strong>Intermittent / Constant</strong></td>
<td><strong>Fluctuations in volume Y / N</strong></td>
<td><strong>&quot;Bad days&quot; Y / N</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Activities prevented or affected:</strong></th>
<th><strong>Effect of sound:</strong></th>
<th><strong>How long:</strong></th>
<th><strong>Ear overprotection Y / N</strong></th>
<th><strong>% of time</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Concentration</strong></td>
<td>None / Louder / Softer</td>
<td>min / hours / days</td>
<td>in quiet Y / N</td>
<td>% of time</td>
</tr>
<tr>
<td><strong>Sleep</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>QRA</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Work</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Restaurants</strong></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td><strong>Sports</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Social</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Other</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>% of time when:</strong></td>
<td><strong>Aware</strong></td>
<td><strong>Annoyed</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Severity:</strong></td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Annoyance:</strong></td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Effect on Life:</strong></td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Comments:**

<table>
<thead>
<tr>
<th><strong>OVERSENSITIVITY:</strong></th>
<th><strong>Frequency</strong></th>
<th><strong>&quot;Bad days&quot; Y / N</strong></th>
<th><strong>Description of troublesome sounds</strong></th>
<th><strong>Effects of sound:</strong></th>
<th><strong>How long:</strong></th>
<th><strong>Ear overprotection Y / N</strong></th>
<th><strong>% of time</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Y / N</strong></td>
<td><strong>Physical discomfort? Y / N</strong></td>
<td><strong>Effected sound:</strong></td>
<td><strong>None / Stronger / Weaker</strong></td>
<td><strong>min / hours / days</strong></td>
<td><strong>in quiet Y / N</strong></td>
<td><strong>Any other T specific treatments</strong></td>
<td><strong>% of time</strong></td>
</tr>
<tr>
<td><strong>Activities prevented or affected:</strong></td>
<td></td>
<td><strong>Effect of sound:</strong></td>
<td><strong>None / Stronger / Weaker</strong></td>
<td><strong>How long:</strong></td>
<td><strong>Ear overprotection Y / N</strong></td>
<td><strong>% of time</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Concerts</strong></td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
<td><strong>Effected sound:</strong></td>
<td><strong>None / Stronger / Weaker</strong></td>
<td><strong>How long:</strong></td>
<td><strong>Ear overprotection Y / N</strong></td>
<td><strong>% of time</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Church</strong></td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
<td><strong>Effected sound:</strong></td>
<td><strong>None / Stronger / Weaker</strong></td>
<td><strong>How long:</strong></td>
<td><strong>Ear overprotection Y / N</strong></td>
<td><strong>% of time</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Dining</strong></td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
<td><strong>Effected sound:</strong></td>
<td><strong>None / Stronger / Weaker</strong></td>
<td><strong>How long:</strong></td>
<td><strong>Ear overprotection Y / N</strong></td>
<td><strong>% of time</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Driving</strong></td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
<td><strong>Effected sound:</strong></td>
<td><strong>None / Stronger / Weaker</strong></td>
<td><strong>How long:</strong></td>
<td><strong>Ear overprotection Y / N</strong></td>
<td><strong>% of time</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Housekeeping</strong></td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
<td><strong>Effected sound:</strong></td>
<td><strong>None / Stronger / Weaker</strong></td>
<td><strong>How long:</strong></td>
<td><strong>Ear overprotection Y / N</strong></td>
<td><strong>% of time</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Other</strong></td>
<td>0 1 2 3 4 5 6 7 8 9 10</td>
<td><strong>Effected sound:</strong></td>
<td><strong>None / Stronger / Weaker</strong></td>
<td><strong>How long:</strong></td>
<td><strong>Ear overprotection Y / N</strong></td>
<td><strong>% of time</strong></td>
<td></td>
</tr>
</tbody>
</table>

**Comments:**

<table>
<thead>
<tr>
<th><strong>HEARING:</strong></th>
<th><strong>Type:</strong></th>
<th><strong>Recommended:</strong></th>
<th><strong>Category:</strong></th>
<th><strong>Recommendation:</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Y / N</strong></td>
<td><strong>Hearing aid(s):</strong> Y / N</td>
<td><strong>Ever recommended:</strong> Y / N</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>RANKING problems:</strong></th>
<th><strong>Tinnitus:</strong></th>
<th><strong>Sound tolerance:</strong></th>
<th><strong>Hearing:</strong></th>
<th><strong>Pln decision:</strong></th>
<th><strong>Next visit:</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>0 1 2 3 4 5 6 7 8 9 10</strong></td>
<td><strong>0 1 2 3 4 5 6 7 8 9 10</strong></td>
<td><strong>0 1 2 3 4 5 6 7 8 9 10</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>T - tinnitus</strong></th>
<th><strong>ST - sound tolerance (hyperacusis + phonophobia)</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Is you T preventing or affecting any activities in your life?</td>
<td></td>
</tr>
<tr>
<td>QRA - quiet recreational activities: Is your T interfering with QRA such as reading or meditating.</td>
<td></td>
</tr>
<tr>
<td>% of time: Aware - What % of time were you aware of your T over last month?</td>
<td></td>
</tr>
<tr>
<td>Annoyed - What % of the time over last months T bothered you?</td>
<td></td>
</tr>
<tr>
<td>Severity - How strong or loud was your T on average over last month? 0 - no T, 10 - as strong as you can imagine.</td>
<td></td>
</tr>
<tr>
<td>Annoyance - How much was T annoying you on average over last month 0 - no effect, 10 - as much as you can imagine.</td>
<td></td>
</tr>
<tr>
<td>Effect on life - How much was T affecting your life on average over last month. 0 - no effect, 10 - as much as you can imagine.</td>
<td></td>
</tr>
<tr>
<td>Any other T specific treatments - Are you using any other treatments for your T.</td>
<td></td>
</tr>
<tr>
<td>Sound tolerance - Is your tolerance to louder sounds the same as people around you?</td>
<td></td>
</tr>
<tr>
<td>Hearing - Do you think you have a hearing problem?</td>
<td></td>
</tr>
<tr>
<td>Ranking - rank importance of your problems with 0 - no problem, 10 - as large as you can imagine</td>
<td></td>
</tr>
</tbody>
</table>

P.J. Jastreboff, 1999
Form 2: Tinnitus/Hyperacusis follow-up interview form

<table>
<thead>
<tr>
<th>Tinnitus Hyperacusis Follow-Up Interview Form</th>
</tr>
</thead>
<tbody>
<tr>
<td>CATEGORY:</td>
</tr>
<tr>
<td>Date of init. couns.</td>
</tr>
<tr>
<td>Date of instr. fitt.</td>
</tr>
<tr>
<td>SG:</td>
</tr>
<tr>
<td>HA:</td>
</tr>
<tr>
<td>FUQ #:</td>
</tr>
<tr>
<td>Month #:</td>
</tr>
<tr>
<td>Type of visit:</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Activities prevented or affected:</th>
<th>Changes: Y / N</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concentration</td>
<td>QRA</td>
</tr>
<tr>
<td>Sports</td>
<td>Work</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>% of time when: Aware Annoyed</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1st)</td>
</tr>
<tr>
<td>(1st)</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Effect of sound:</th>
</tr>
</thead>
<tbody>
<tr>
<td>None / Louder / Softer</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Effect on:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Life:</td>
</tr>
<tr>
<td>Work:</td>
</tr>
<tr>
<td>Sports:</td>
</tr>
<tr>
<td>Social:</td>
</tr>
<tr>
<td>Other:</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Comments:</th>
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</thead>
</table>

<table>
<thead>
<tr>
<th>Tinnitus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Severity</td>
</tr>
<tr>
<td>Annoyance</td>
</tr>
<tr>
<td>Effect on Life</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sound Tolerance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Activities</td>
</tr>
<tr>
<td>Changes: Y / N</td>
</tr>
<tr>
<td>Concerts</td>
</tr>
<tr>
<td>Shopping</td>
</tr>
<tr>
<td>Movies</td>
</tr>
<tr>
<td>Restaurants</td>
</tr>
<tr>
<td>Driving</td>
</tr>
<tr>
<td>Sports</td>
</tr>
<tr>
<td>Church</td>
</tr>
<tr>
<td>Housekeeping</td>
</tr>
<tr>
<td>Social</td>
</tr>
<tr>
<td>Other</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Effect of sound:</th>
</tr>
</thead>
<tbody>
<tr>
<td>None / Stronger / Weaker</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Effect on:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quiet:</td>
</tr>
<tr>
<td>Work:</td>
</tr>
<tr>
<td>Social:</td>
</tr>
<tr>
<td>Other:</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Comments:</th>
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</table>

<table>
<thead>
<tr>
<th>Hearing problem</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>The problem in general:</th>
<th>Same / Better / Worse</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tinnitus: 0 1 2 3 4 5 6 7 8 9 10</td>
<td></td>
</tr>
<tr>
<td>Sound tolerance: 0 1 2 3 4 5 6 7 8 9 10</td>
<td></td>
</tr>
<tr>
<td>Hearing: 0 1 2 3 4 5 6 7 8 9 10</td>
<td></td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Recommendation:</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>How would you feel if you had to give back your instruments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Are you glad you started this program? Y / N / NS</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Main problems discussed:</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>T - tinnitus</th>
</tr>
</thead>
<tbody>
<tr>
<td>ST - sound tolerance (hyperacusis - phonophobia)</td>
</tr>
<tr>
<td>Type of visit:</td>
</tr>
<tr>
<td>Phone, fax, e-mail, office visit</td>
</tr>
<tr>
<td>Is you T preventing or affecting any activities in your life?</td>
</tr>
<tr>
<td>QRA - quiet recreational activities:</td>
</tr>
<tr>
<td>% of time when:</td>
</tr>
<tr>
<td>Aware - What % of time were you aware of your T over last month?</td>
</tr>
<tr>
<td>Annoyed - What % of the time over last months T bothered you?</td>
</tr>
<tr>
<td>Severity - How strong or loud was your T on average over last month?</td>
</tr>
<tr>
<td>Annoyance - How much T was T annoying you on average over last month?</td>
</tr>
<tr>
<td>Effect on life - How much T was affecting your life on average over last month?</td>
</tr>
<tr>
<td>Any other treatments - Are you using any other treatments for your T?</td>
</tr>
<tr>
<td>Sound tolerance - Is your tolerance to louder sounds the same as people around you?</td>
</tr>
<tr>
<td>Hearing - Do you think you have a hearing problem?</td>
</tr>
<tr>
<td>Ranking - rank importance of your problems with 0 - no problem, 10 - as large as you can imagine</td>
</tr>
</tbody>
</table>

MM & PJ Jastreboff, 1999
References

Tinnitus Retraining Therapy

60. Impedance fluctuation and a “Tensor Tympani Syndrome”. 79 Sep 25; Lisbon: Universidad Nova de Lisboa Ed Penha & Pizarro; 1979, Ref ID: 8065
Keypoints

1. There is considerable evidence that many forms of tinnitus are caused by central changes that may occur after peripheral lesions.
2. Auditory stimulation is one of the most employed therapeutic methods for tinnitus, and one of the most beneficial.
3. Sound generators that emulate environmental sounds are small devices that allow a person to select the favorite kind of sound at the most comfortable volume.
4. Custom sound generators, for normal hearing persons, are similar to hearing aids, very light, and to be worn behind the ear. They generate a wide-band sound that can be adjusted to the user’s needs.
5. Hearing aids designed for people with tinnitus and hearing loss provide amplification that facilitates auditory stimulation to ameliorate tinnitus.
6. Implantable hearing aids are now used by many people, which made it possible to assess their efficacy in tinnitus treatment.
7. Other devices can be used for tinnitus management for immediate relief before a more complete sound therapy can be initiated.
8. Prosthesis and “open-ear” hearing aids are preferred for treatment of tinnitus. These devices provide amplification in narrow frequency bands which can be adjusted to coincide with the frequencies of the patient’s hearing loss.
9. Sound stimulation has its beneficial effect on most forms of tinnitus by activating neural plasticity, which requires time to develop. The time it takes for sound stimulation to reduce an individual’s tinnitus varies and may require a 6- to 8-month time frame.
10. The selection of hearing aids must be tailored to individual patients, based on the patient’s clinical picture.
11. The specific guidelines on hearing aid device adaptation are crucial for an effective auditory stimulation of tinnitus-affected patients.

Keywords  Tinnitus • Hearing aid • Sound enrichment • Hearing loss • Auditory stimulation

Abbreviations

Combi  Combination hearing aid
dB  Decibel
Hz  Hertz
SA  Spontaneous activity
TRT  Tinnitus retraining therapy

Introduction

Effective treatment of tinnitus depends on understanding the cause of tinnitus. Especially regarding treatment with sound, it is important to know if tinnitus is caused by pathology of the ear or the auditory nervous system.

The past years have witnessed a change in the understanding of the cause of tinnitus. Previously, tinnitus was believed to originate from the peripheral auditory
system [1, 2]. There is now considerable evidence that most forms of tinnitus are caused by changes in the central nervous system after peripheral lesions [3, 4]. For treatment of tinnitus, it is important to distinguish between these two models, as they imply different therapeutic strategies. In fact, the peripheral model suggests that the aberrant neural activity is responsible for tinnitus perception. This hypothesis has been inspired by the results of an animal study [5], by showing an increase in the spontaneous activity (SA) in the cochlear nerve after the administration of a high dose of salicylate (400 mg/kg in cats) assumed to cause tinnitus. A recent study [6] has shown that salicylate-induced tinnitus may be caused by activation of NMDA receptors expressed in the synapses of cochlear hair cells and dendrites of spiral ganglion neurons. If tinnitus was normally caused by increased activation of NMDA receptors, a possible therapeutic approach that could suppress such “peripheral tinnitus” would be inactivating NMDA receptors [6, 7]. However, NMDA receptor blockage has not been shown as effective treatment of tinnitus.

High doses of salicylate are also known to cause nonspecific (toxic) effects, especially in cats, which lack the enzyme necessary to metabolize salicylate (glucuronyltransferase). Such nonspecific effects could account for the increase in SA in the cochlear nerve after the administration of high doses of salicylate (see above). More recent studies have shown that salicylate, at a dose of approximately 200 mg/kg, known to induce tinnitus in animals [8], does not increase SA in the cochlear nerve [9, 10], but increases neural activity in auditory centers [11–13]. These studies then question the peripheral origin of salicylate-induced tinnitus. It is also worth noting that recent studies suggest that salicylate has strong effects on the central auditory nervous system [14–17]. These findings indicate that salicylate may induce tinnitus through central mechanisms.

The most frequent causes of tinnitus seem to be cochlear damage, as almost all individuals with tinnitus have hearing loss. Importantly, cochlear damages – induced after noise trauma, for instance – cause a dramatic decrease of SA in the cochlear nerve [18, 19]. Damages to the inner hair cells (or their stereocils) have been shown to decrease the spontaneous release of glutamate from the inner hair cells (cochlear nerve synapses), thereby causing the decrease in SA. This strongly argues against a peripheral origin of tinnitus encountered in human subjects (related to peripheral damage). If the neural activity is decreased in the cochlear nerve, there should be a kind of compensatory mechanism, which could generate an aberrant neural activity in the auditory centers. In this context, it has been shown that cochlear damage decreases the inhibitory neurotransmission in the auditory centers [20–23]. This decrease in central inhibition is supposed to account for the changes in the evoked and SA after cochlear damage. First, hearing loss of a sufficient extent induces a reorganization of the tonotopic map, i.e., neurons with their characteristic frequency corresponding to the hearing loss region change their frequency tuning toward the cut-off frequency of hearing loss [24, 25]. In addition, a strong neural hyperactivity has been observed in the auditory cortex after a noise trauma [26]. This hyperactivity could be a neural correlate of hyperacusis, i.e., overestimation of loudness, sometimes reported by subjects presenting a hearing loss. Finally, changes in the pattern of spontaneous discharge (increase in firing rate and synchrony), consistent with the psychoacoustic properties of tinnitus [4], have been observed after acoustic trauma [3, 27]. These neural changes of the SA could then be neural correlates of tinnitus.

Rationale for Stimulating the Frequency Range of Hearing Loss: Reversing Central Changes Induced by a Decrease in Afferent Inputs

In summary, the decrease in afferent input caused by peripheral lesions could trigger dramatic central changes, such as a release from central inhibition. These central changes could ultimately result in the emergence of an aberrant neural activity that could induce tinnitus. In this context, we have suggested an approach consisting of preventing/compensating the decrease in afferent input related to hearing loss. This could reverse the central changes normally associated to it and, as a consequence, decrease/suppress tinnitus [3, 4, 28, 29]. The aim of this approach is to normalize the SA over frequency (in patients with high-frequency hearing loss, the approach consists of
increasing sensory inputs in this frequency band) and/or increase the overall level of sensory input (in patients with flat and severe hearing loss). Central inhibition could control a kind of central gain [28, 30], increasing central inhibition, by providing the auditory system with augmented input that is supposed to decrease neural hyperactivity induced after hearing loss. In animals, we have shown that an acoustic environment enriched in high frequencies could prevent the central changes normally induced after a noise-induced hearing loss [24, 31]. Moreover, we could induce a dramatic decrease of hypersensitivity in human subjects reporting hyperacusis, after these subjects were stimulated a few hours a day for several weeks with a customized stimulus (the long-term spectrum of the stimulus corresponded to the hearing loss of each subject [28]).

Auditory Stimulation Delivery

Auditory stimulation is one of the most employed therapeutic methods and one of the most beneficial for patients suffering from tinnitus [32] (see also Chap. 75). Such therapy has no noticeable side effects and may be administered through simple devices [33]. Sounds used may resemble environment sounds, which enrich the atmosphere in the room they are used. In case sound enrichment should be required all day long (and tinnitus is not associated to hearing loss), “custom” ear level sound generators may be suitable. These are small electronic devices fitted to the ear. For individuals with hearing loss, open-ear hearing aids are suitable [34, 35], as well as tinnitus control combination instruments (Combi), which combine a prosthesis and a sound generator. These devices both amplify environmental sounds and generate sound enrichment.

Sound Environment Generators

Sound environment generators are contained in a small case, in which batteries and speakers are also housed. The volume can be regulated by means of a small roller on the side of the device. Different buttons may be pushed to select different sounds such as sea waves, creeks, waterfalls, rain, the woodlands, and white noise. For most users, these sounds are relaxing, as they are monotonous and repetitive without interruption. Once a given sound has been selected and the volume has been regulated, the user can use the environmental sound as background noise. For this reason, such sound generators are particularly useful during night rest (Fig. 74.1).

Custom Sound Generators

Custom sound generators look like regular hearing aids; they are light and designed to be worn behind the ear. A thin wire connects the generator to the speaker placed at the entrance of the ear canal.

Unlike the sound generated by environmental sound machines, the sound generated by custom sound generators can only be heard by the person wearing the device. These devices generate a wide band sound that can be adjusted by the audiologist to meet the final user’s needs by means of high-pass or low-pass filters and may even be modulated in width.

The size of the mini speaker placed at the entrance of the auditory canal is such that it does not affect normal hearing. Custom sound generators are beneficial for individuals with normal hearing.

The small size of these sound generators makes them easy to wear during everyday activity. Once they have been worn and the volume regulated, the person may
“forget” they are wearing them for the rest of the day. Their maintenance is limited to periodically replacing the battery which can be done by the user. Custom sound generators are both useful for total masking therapy [36] and for partial masking therapy, according to tinnitus retraining therapy (TRT) [37] (Fig. 74.2).

**Hearing Aid Devices**

The most suitable hearing aids for sound therapy are the open-ear hearing aids [38, 39], which have a mini speaker placed at the entrance of the ear canal. The size of the small case housing the battery resembles that of a bean. Like the custom sound generators, their ease is such that wearers often do not even feel them.

Hearing aids are designed to compensate for hearing loss and lack of auditory stimulation. Unfortunately, hearing aids currently available are not able to amplify sounds with a frequency above 6–7 kHz, a range of hearing that is often impaired in individuals with tinnitus; for this reason, ordinary hearing aids may be less efficient in compensating for lost auditory stimulation. Besides hearing aids, the new generation Combi (combination hearing aids) now available, combine common prostheses with the ability to generate an enrichment sound, similar to what custom sound generators provide. The Combi devices represent the most innovative and efficient therapeutic tools for tinnitus and hearing loss, because they can combine auditory stimulation in impaired hearing areas with either partial or total tinnitus masking [40, 41].

**Implantable Hearing Aids**

Traditional acoustic prostheses and Combi hearing aids are not generally recommended for patients with conductive hearing loss caused by external and middle-ear malformations or in patients with chronic middle-ear infection. Such individuals may benefit from the bone-anchored hearing aids, which transmit sound vibrations to the inner ear through a titanium rod implanted into the bone. The increase in use of implantable hearing aids during recent years has made it possible to assess their efficacy for treatment of tinnitus. Implantable middle-ear prostheses provide better sound therapy for some patients with tinnitus than traditional hearing aids [42], probably because they provide amplification in a wider frequency range and because of the “naturalness of the amplification”. The cost, as well as the required surgery, limits the use of these devices. Cochlear implants can provide input to the auditory nervous system that can reduce tinnitus in many individuals, both in those with severe hearing loss and in individuals with good hearing on one ear who have severe tinnitus referred to that side [43] (see also Chaps. 76 and 77).

**Other Sound Therapy Devices**

Besides sound generators and acoustic prostheses, other devices that are not specifically designed for treatment of tinnitus can be used for tinnitus manage-
ment. In fact, a simple fan or fish tank can be used as first-aid treatment of tinnitus. Music players, such as MP3 players with headphones, are often employed to reduce tinnitus. Recorded nature sounds played through home stereo systems are used for this purpose as well.

Can such devices replace custom sound generators or acoustic prostheses? We do not believe they can for the following reasons: MP3 players and headphones can hardly be worn by individuals carrying out non-sedentary activities; they partially occlude the ear; and they may become intrusive and cannot be worn (and forgotten) for 6–8 h a day. Conversely, these devices may be useful for immediate relief before a more complete sound therapy is started.

Auditory Stimulation from Theory to Practice

Clinical studies [29, 34, 35] have shown that not only do hearing aids improve hearing ability, but they can also reduce or suppress tinnitus.

For instance, in a study carried out in 1999 [44], 50% of hearing aid wearers experienced relief from tinnitus, with a median improvement of 10% after only 6 weeks from the first application. These results were confirmed by subsequent studies, which extended the investigation to individuals who had tinnitus and mild hearing loss [34, 45].

Prosthesis and open-ear hearing aids are important for proper treatment of tinnitus. Modern hearing aids can provide amplification at the frequencies where hearing loss occurs, without uncomfortable side effects, such as over amplification or rumbling, which were typical in the old generation devices.

Individuals with hearing loss that is limited to mild damage of hair cells not affecting the subjective hearing sensitivity benefit from custom sound generators or sound environment generators [46]. Our experience from daily clinical practice, as well as the experience of others reported in published studies [47], has shown that hearing aids and sound generators can achieve the following goals:

- Improving communication and reducing the discomfort often reported by patients as sounds and voices covered by tinnitus.
- Stimulating the auditory nervous system in a normal way and not only with tinnitus (phantom sounds).

The Approach to Sound Therapy

The role of the therapist should not be limited to the technical aspects of hearing aids and their application, but should aim at developing an empathic and confident relationship with the individual patient. Only a comprehensive evaluation may allow the therapist to have an accurate picture, in order to tailor the most appropriate and effective therapeutic plan. Hearing device application and control for adaptation may require a series of scheduled visits every 3–4 months, although in some cases a stricter follow-up schedule may be necessary. The results of long-term treatment may be assessed through visual analog questionnaires and the use of different kinds of scales [37] to allow tracking treatment progress. Audiometric test results do not usually reflect variations in tinnitus and thus, are not valid measures of relief [48]; tests, therefore, do not need to be periodically repeated. Cerebral plasticity requires some time to develop, and the needed duration of therapy may, therefore, vary from patient to patient [49]. Optimal relief from tinnitus may require a 6- to 8-month therapy using hearing aids and sound generators [50, 51].

Hearing Aid Selection

The selection of the most appropriate hearing aid device should be based on the individual patient’s needs. For example, sound environment generators are mostly indicated during night rest in patients affected by mild tinnitus. However, patients with disturbing tinnitus and without subjective hearing impairments benefit from custom sound generators, which should be worn at least 8 h during the daytime, in combination with an environment generator during night rest. Combi-type devices are suitable for patients with mild hearing loss. These can also provide environmental sound enrichment during night rest.
Hearing Aid Device Adaptation

In order to achieve an optimal auditory stimulation, specific guidelines on hearing aid device adaptation should be followed, for custom sound generators, Combi devices, or prostheses [29, 34]. The parameters are crucial for auditory stimulation achieving maximal benefits on tinnitus.

The best results are achieved when the external auditory canal is left as accessible as possible. In fact, even partial occlusion of the auditory canal may cause unease of use and may even increase tinnitus perception. It may also affect the natural acoustic properties of the external ear, with further negative side effects causing a loss of the natural acoustic resonance, which is important for naturalness of hearing. Occlusion of the ear canal also causes over-emphasis of low frequencies with rumbling sensations resulting together with diminished perception of sound in the most important frequency range of hearing. It is also important not to underestimate the hearing of one’s own voice which often causes difficulties in the understanding of speech, as well as being unpleasant for the individual and may cause a sensation of “closure” that can worsen tinnitus. The introduction of the so-called open-ear hearing aids helped overcome some of these problems, allowing application of hearing aids to individuals with mild hearing loss, such as many individuals with tinnitus have. Open-ear hearing aids also provide a stimulation mainly in the frequency region of the tinnitus pitch. The open-ear hearing aids, thus, provide important advantages, such as sound enrichment, that reduce tinnitus by activating the neural plasticity. Open-ear prostheses can also be employed in patients with severe hearing loss; acoustic feedback is reduced (or eliminated) by computer programs in modern digital hearing aids. Hearing naturalness and ease of use are important factors or advantages of digital hearing aids. In the selection of hearing aids, all elements that can cause a patient’s discomfort and increase the perception of tinnitus must be taken into account, including cosmetic aspects. Hearing aids and sound generators should ideally be forgotten after they have been applied. In other words, people should become unaware of wearing a hearing aid device.

Hearing aid devices should simultaneously be worn in both ears, in order to favor a complete and simultaneous stimulation of the entire auditory nervous system. This is also important for unilateral tinnitus. Moreover, the frequency band of hearing aids should be adjusted to mostly amplifying the frequency range that is most important for hearing. Sound generators should be adjusted to the frequency of the tinnitus in order to activate the auditory nerve close to tinnitus frequency.

Our clinical experience supports the use of prescription formulas of gain/output suggested by device manufacturers, although major modifications are very often necessary. In fact, many tinnitus patients are sensitive to amplification, which sometimes requires less gain and maximum output than in patients who do not have tinnitus. Patients with moderate to severe hearing loss often benefit from amplifications that are 50–70% lower than traditional prescription formulas. The large variability of the requirements for tinnitus patients regarding amplification has prevented adaptation of an uniform formula that is suitable for all tinnitus patients. Individuals with tinnitus often benefit from having the option of noise reduction switched off or turned down.

Patients must be properly instructed in how to adjust the volume on their devices. Patients are generally able to fully understand the volume regulation procedure and to safely carry it out, but often more than one round of counseling is necessary and analog scales should be used to track the intensity of both tinnitus and therapeutic sound. During TRT therapy, the correct balance between sound stimulation and amplification can be determined with in situ instruments after some weeks of use [52]. The intensity of auditory stimulation should be 5–6 dB higher than the threshold level in order to prevent stochastic resonance phenomena [37].

Optimal results in management of tinnitus are not only obtained with the application of technologically advanced hearing aid devices but, most of all, with their adjustment to the individual person’s needs and through patient counseling. Each single patient must be listened to, counseled, and informed throughout therapy planning and during follow-up. This enables therapists to fully understand their patient’s problems and to solve them to the greatest extent through a proper selection of prosthetic devices and finding the optimal settings.

References


Rehabilitation of Tinnitus Patients Using the Neuromonics Tinnitus Treatment

Dayse Távora-Vieira and Paul B. Davis

Chapter 75

Keypoints

1. Tinnitus, a phantom perception of sound, is a frequent clinical condition that may cause significant debilitation.
2. Tinnitus treatments can focus on the condition itself or on patients’ reaction to their tinnitus.
3. There is a growing acceptance that sound stimulation incites a neuroplastic change in the auditory pathways.
4. Neuromonics utilizes highly tailored music and broad frequency wave sounds, in the context of a structured counseling, support, and monitoring program, with the intention of reversing the neurological, psychological, and audiological processes that caused the disturbance.
5. Neuromonics also aims to facilitate a relaxation response and shift of attention, leading to a desensitization effect.
6. Clinical trial and private clinic outcomes show that the neuromonics tinnitus treatment can consistently provide clinically significant levels of desensitization to tinnitus perception over a relatively short period of time.

Keywords Neuromonics • Tinnitus • Hyperacusis • Desensitization • Acoustic stimulation

Abbreviations

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<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>HL</td>
<td>Hearing level</td>
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<td>LDL</td>
<td>Loudness discomfort levels</td>
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<td>MML</td>
<td>Minimum masking levels</td>
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<td>NTT</td>
<td>Neuromonics tinnitus treatment</td>
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<td>RI</td>
<td>Residual inhibition</td>
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<td>THQ</td>
<td>Tinnitus history questionnaire</td>
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<td>TRQ</td>
<td>Tinnitus reaction questionnaire</td>
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<td>TRT</td>
<td>Tinnitus retraining therapy</td>
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Introduction

Tyler [1] categorizes tinnitus treatment in two ways: one focused on tinnitus reduction or elimination (e.g., medications, electrical suppression) and the other focused on a patient’s reaction to the condition. Current thinking on the underlying mechanism of tinnitus emphasizes changes in the auditory and neural systems that can be broadly related to the aspects of perception, attentional, and emotional reaction to tinnitus [1–4] (see Chap. 73). Recent animal studies have shown that tinnitus may be linked to cortical reorganization [5]. In addition, brain imaging studies have shown altered levels of activation in several areas of the brain in tinnitus sufferers. These areas include, for instance, the left temporal lobe (Brodmann area 21 & 41), left hippocampus and posterior thalamic region [6], right middle frontal and right middle temporal gyri [7], amygdala, parahippocampal gyrus and hippocampus [8], inferior colliculus [9], subcallosal area [10], right inferior colliculus in the left hippocampus [11], and the Heschl’s gyrus [12].

The neuromonics tinnitus treatment (NTT) is a structured tinnitus rehabilitation program consistent with these current models, incorporating structured...
counseling and individually customized broad frequency sounds including music for relaxation. The sounds are spectrally shaped to individually correct for each patient’s hearing loss configuration and it aims to address the auditory deprivation element of tinnitus pathogenesis, providing most the broadest stimulation of the auditory pathways. The purpose is to decrease the limbic system/amygdala’s involvement in the patient’s perception and reaction to tinnitus, thus promoting relaxation and relief [13–16]. This chapter provides an overview of the underlying principles of NTT, describes the standard protocol, and outlines candidacy for NTT in the context of published clinical trial and private practice data. The importance of counseling and the main challenges faced by patients and clinicians will be discussed.

**Theoretical Basis for Counseling and Sound Treatment**

Most hypotheses about the pathology of tinnitus agree that the abnormal neural activity in the brain that is perceived as a sound is a result of neuroplastic processes (e.g., [17, 5]). There is a general agreement that the limbic system and autonomic nervous system are involved in generating the awareness and annoyance from tinnitus [1–3, 17, 18]. This can explain why an individual’s reaction to tinnitus seems to be linked to the person’s emotional state.

The cognitive and emotional reaction to tinnitus is the target for tinnitus treatment that uses counseling. Some commonly used forms of tinnitus treatment combine counseling with acoustic stimulation: tinnitus retraining therapy (Chap. 73) and a hearing aid program (Chap. 74) are among these approaches.

NTT is based on correcting the abnormal neural activity that causes tinnitus by inducing neural modification within areas of the brain related to audition, attention, and emotion [15]. NTT involves counseling and sound stimulation consisting of music and broad frequency sound (shower noise-like sound) [13–18].

A unique aspect of NTT is that the spectrum of the sound is individually modified to account for each patient’s hearing loss; this enables the intensity to be set to a comfortably relaxing level. In a study of 35 patients, the average reduction in loudness of their tinnitus was 16.11 dB in a paired comparison between the original source music and the corresponding customized music, which the patient had set to their minimum masking level (MML) [13]. The mean level of the source music was 72 dBA, which would make sleep onset and concentration difficult. The mean customized signal was 56 dB. This 16 dB difference is quite clinically significant when one recalls that 6 dB constitutes a perceptual halving of loudness. This ability to provide a high degree of relief at a comfortable level tends to greatly facilitate a sense of control over the tinnitus. As a result, it greatly reduces the significance of the tinnitus [15].

The purpose of the “shower noise”, combined with music, is to restore normal activation of the auditory system that may have been deprived of stimulation due to hearing loss or other forms of auditory dysfunction. The rationale is this maximizes the efficiency of sound stimulation to induce a neuroplastic change in the auditory pathways [18]. This has been shown in numerous studies by the marked improvement over treatment in neuroplastic-mediated processes that are reflected by MML and loudness discomfort levels (LDL) [13, 14, 16].

Asymmetric hearing loss is compensated for to allow a balanced loudness perception. Consequently, a true stereo sound is afforded by the system’s ability to phase-lock the left and right channels. Considerable peak compression is also applied to fit within the typically narrow dynamic ranges of those with disturbing tinnitus. Further details of the algorithms for the customization of the sound stimuli are described in Hanley and Davis [15].

If the tinnitus is considered to be important, it will become part of patients’ consciousness and has been hypothesized to trigger a strong negative emotional reaction [19, 20]. In tinnitus retraining therapy (TRT), the use of noise was proposed because it may be considered “neutral” to the limbic system. Some patients may find noise to be unpleasant, thus possibly contributing to reasons why some patients have rejected the use of TRT [21, 22]. NTT uses relaxation music as the predominant signal in order to activate the limbic system with positive associations. The sound is presented within the context of a counseling program with a more collaborative, patient-centered orientation.

Music has long been recognized to provide a therapeutic effect and has been empirically used for facilitating a major relaxation response, a welcomed shift of attention [23]. Tyler [1] suggests the use of music
to provide distraction from tinnitus and notes that it has otherwise been highly underutilized. The use of music in NTT enables tinnitus sufferers to have a pleasant and relaxing sound to listen to while being treated. This approach facilitates patients’ compliance to treatment and provides gradual desensitization to the tinnitus signal [13, 18].

In summary, NTT aims to facilitate the process of gradual desensitization of tinnitus disturbance. Together with a structured counseling program, NTT uses acoustic stimulation in a way that is effective, yet enjoyable (Fig. 75.1).

What follows is the standard NTT protocol aiming to assist clinicians in delivering the program to the most highly suitable “Tier 1” patients, who comprise around 43% of a typical tinnitus clinic population [16].

**The NTT Protocol and Patient Selection**

Based on clinical trials [13, 14] and outcomes of the treatment reported by private clinics [15], the most suitable candidates are as follows:

- Patients with four-frequency average hearing thresholds better than 50 dB in at least one ear
- Clinically significant tinnitus disturbance reported in the tinnitus reaction questionnaire with score of at least 17

- Normal or decreased sound tolerance
- Tinnitus is neither pulsatile nor multi-tone
- Tinnitus is not exacerbated by normal level of acoustic stimulation (i.e., not highly reactive)
- No active Ménière’s disease or other causes of wide fluctuations in hearing levels
- Patient is well protected when exposed to a noisy environment.

**Assessment**

A comprehensive tinnitus history questionnaire (THQ) is mailed to the patient prior to his/her first visit to the clinic. The THQ involves data on the nature of the tinnitus, tinnitus history, general hearing difficulties, effect of tinnitus, and general patient’s health. It helps the patient to recall all relevant factors and information relevant to the rehabilitation process. It also helps to determine the cause and influencing/exacerbating factors of the tinnitus and assess the patient’s candidacy [18]. The patient also completes the tinnitus reaction questionnaire (TRQ). The score on the TRQ is used for counseling purposes, rehabilitation planning, and to monitor the patient’s progress throughout treatment [13]. It has five response options that relate to the previous week of treatment, so it is more sensitive than most other tinnitus questionnaires [24].
The following baseline audiological measures are obtained: tympanometry, but no acoustic reflexes in patients with decreased sound tolerance; pure-tone audiometry up to at least 12.5 kHz; tinnitus pitch match; tinnitus loudness balance; MML; LDL at 0.5, 1, and 4 kHz; and residual inhibition (RI) (Fig. 75.2).

**Practical Use of NTT**

Prior to commencing the treatment, patients are taught how to manage the device (Fig. 75.2), and discussion on the realistic outcomes during the first phase and neural changes related to the desensitization process over the second phase is considered of high priority for NTT. A good understanding of tinnitus and the proposed treatment are likely to make the patient more compliant and positive about the therapy [1]. Patients are provided with take-home reading material about their assessment results, how NTT works, what to expect at each phase of the treatment, advice on factors that can exacerbate tinnitus perception, and a list of suggested activities that can be done while using the device. The first stage of NTT lasts for approximately 2 months after the fitting date. Patients are instructed to initially use the device for at least 2–4 h a day, and especially during the times that his/her tinnitus is the most disturbing. Patients are also instructed to set the volume at the beginning of each session to a comfortable level that provides a high level of interaction with his/her tinnitus. The high level of interaction aims to increase the amount of neurostimulation (representing the deprived sounds at the auditory cortex) and to provide maximal relief and relaxation for the tinnitus sufferers. This approach would facilitate the desensitization process of the tinnitus signal at a later stage by enabling them to relax despite their tinnitus and overcome prior problems like concentration disturbance and sleep onset/maintenance [15]. As observed by the authors and other clinicians in private clinics, the patients’ monitoring of their tinnitus is contra-productive, potentially delaying, or even stopping the benefit of the treatment. Therefore, patients are encouraged to undertake another quiet activity while using the customized device to avoid monitoring the music and consequently risk monitoring the tinnitus. The second phase of the treatment aims to gradually promote desensitization to the tinnitus signal as a more permanent effect. Patients are re-instructed to set the volume to a level that only covers up their tinnitus around half of the time. Hence, the patient will experience an intermittent interaction with their tinnitus, which happens during the intensity troughs of the music. This fleeting exposure to the tinnitus whilst in a relaxed state is intended to help the brain develop a capability of placing the tinnitus in the background. This phase usually lasts for approximately 4 months, and the progress is
reviewed at weeks 16 and 24. The MML and LDL are measured again at 2, 4, and 6 months after beginning the program to evaluate the patient’s progress. The TRQ is also re-administered and the patient needs to estimate how often they have been aware of (and also how often they have been disturbed by) their tinnitus over the past week. The measures are used to determine the current progress and link back toward the stage-specific pretreatment goals. The patient’s response to the second stage of treatment is checked at the tenth week, as there are a few cases where the patient feels too anxious about not having the extra “shower sound” to fill in the quieter parts of the dynamic signal, and they can worry that their tinnitus is increasing. Few patients may benefit from extra time on phase one. For instance, recent data from the authors suggest that patients with higher levels of hearing loss or protected exposure to noise may benefit from a few extra weeks of high level of interaction [25].

The clinical trials have shown that once the systematic desensitization starts to occur, patients typically report that their tinnitus annoyance gradually decreases, and often their current data suggest improvement from their TRQ, their audiometric MML, LDL, and even their estimates of the percentage of time they are aware of their tinnitus (e.g., [16]). Desensitization is deemed to have occurred when the patient reports that they do not need the device to distract them from the tinnitus. That represents the end of the second stage. A few patients will occasionally use the device in certain situations that trigger their tinnitus, for instance, high levels of stress, noise exposure, middle-ear pathology. [18].

Once tinnitus is no longer a problem, the third phase begins. The patients are encouraged to use the device at least once a week to maintain their improvement because complete withdrawal from treatment may cause re-emergence of tinnitus or decreased sound tolerance. This is consistent with the notion that the subsequent lack of stimulation had caused the auditory deprivation process to reappear. Rebound has not been found to be a factor when patients have kept using their devices, suggesting that the constant tonotopic representation of the hearing loss-frequencies at the cortex has been successful at maintaining their gains [18].

The first step for adequate counseling is the patient’s understanding of hearing and tinnitus mechanisms [1]. Patient should be aware of the factors or illnesses that influence on the habituation process. They should also be advised about specific habits and behaviors that can interfere with the treatment, such as diet, alcohol, excess caffeine intake, insufficient exercise, noise exposure, stress levels.

Motivation and realistic expectations are very important matters for tinnitus treatment, particularly in private practice. Clinicians need to certify that the patient acknowledges the success of the treatment demands their active participation in the rehabilitation program. A number of patients continue to find it hard to understand the difference between tinnitus and tinnitus consequences, and that the NTT can only work on consequences such as the level of awareness and disturbance. If inadequately prepared, patients can display significant decreases on their TRQ score and percentage of disturbance, but show disappointment because they still have tinnitus and often may need revisiting if the patients begin to “move the goalposts”.

**Special Considerations**

Based on clinical outcomes for the first 400 patients treated with NTT [16], patients are assigned to one of three categories: Tier 1 candidacy includes the most suitable patients for NTT; Tier 2 suitability includes patients with TRQ score of less than 17, high psychological disturbance, and four-frequency average hearing thresholds worse than 50 dB; and Tier 3 suitability includes patients with reactive, pulsatile or multi-tone tinnitus, ongoing noise exposure, Ménière’s disease, or hearing loss greater than 50 dB in both ears. The standard (Tier 1) patient group represents the largest cohort in NTT populations found in regular private practice (48%); Tier 2 represents 37% and Tier 3 represents 15%. Whilst Tiers 2 and 3 can still be treated with NTT, they need to understand and acknowledge that their progress may be slower and more modest than usual [16].

Patients with very decreased sound tolerance (mean LDLs <85 dBHL) are common, comprising 66% of participants in Trial 3 [13]. These patients were found to respond consistently well to NTT, with a > 5 dB improvement in LDLs found in 85% of those patients using the two-stage stimuli (mean change was 11 dB). A hyperacusis protocol has been subsequently formulated [26] (neuromonics, ND). This hyperacusis protocol constitutes a variation on the usual protocol. They are not encouraged to strive for a high level of interaction in the first phase, but instead just keep it as high as it can be whilst remaining comfortable. In case
of history of even every day sounds making their tinnitus flare up, they should begin on the second phase of treatment immediately, again emphasizing comfort in setting the device intensity levels.

Patients with significant hearing loss across the speech range should be considered for hearing aid fitting in conjunction with NTT, but only if the patients have sufficient dynamic range to make the hearing aids tolerable (see Chap. 74). Many hearing impaired patients have tried or are wearing hearing aids, but report that tinnitus is most disturbing during quiet times, mainly when trying to sleep at night [26]. Some may find difficult to tolerate amplified sounds. These patients have found that NTT can improve their LDLs and reduce their reactivity, so that they become progressively better candidates for amplification [16].

Patients with high levels of hearing loss are likely to need to be kept in phase one of the treatment longer, as they can progress slower. They should be advised to wear the treatment for longer each day, and that the full program will also be extended for a longer period of time [25].

Patients with unilateral severe to profound hearing loss and those with asymmetric hearing loss (difference greater than 45 dB between the ears) are still considered candidates for NTT, applying a variation of the standard protocol with contra-lateral stimulation, which has been found to be effective in a cohort of 40 participants with severe unilateral hearing loss [27]. The thresholds of the better ear are used to customize the device, and the stimulation is provided only in that ear.

Patients with a level of tinnitus disturbance that is not having a significant effect on their quality of life (TRQ composite score less than 17) [24] have been found to be poor responders to NTT [16], perhaps because there is little central gain to reduce. These patients are more likely to benefit from counseling and an educational training. The clinician should provide information on the hearing and tinnitus mechanism and offer a follow-up appointment if the patient feels it is needed. Patients can still be good candidates for NTT when they have low TRQ, but a high level of specific distress (e.g., major concentration disturbance). As the TRQ is more relevant to tinnitus than hyperacusis, when the hyperacusis is the main problem, the TRQ can be low. However, these patients are in great need of treatment and are usually excellent NTT candidates.

Finally, patients pursuing compensation related to his/her tinnitus are encouraged to complete the case prior to treatment, given their responses have been found to be reduced and far less consistent [28]. The stress of the legal process and the continual reminders of the significance tends to be very counterproductive to a desensitization program such as neuromonics. However, if they are prepared to pay for the program themselves, it may be an indication that they may have enough intrinsic motivation to start sooner (see also Chap. 70).

Clinical Trials

NTT has been the subject of three major clinical trials. The first clinical trial strongly supported the idea that customization of music according to a patient’s hearing profile is clinically more effective in tinnitus masking than customized noise [29]. The second clinical trial was a randomized controlled study of 50 tinnitus patients [14]. The exclusion criteria for this study were severe hearing loss in the better ear (four-frequency average hearing threshold greater than 70 dB), tinnitus-related compensation claim, ongoing noise exposure, major psychological disturbance (such as depression or psychosis), cognitive impairment, TRQ score of less than 17, and another simultaneous tinnitus treatment.

The second clinical trial compared clinical outcomes obtained with NTT, counseling alone, and counseling plus broadband noise set at the TRT mixing point. All groups had equal amounts of clinician time, and the in-depth counseling was reinforced by use of the self-help book, “Living with Tinnitus” [30]. Patients were evaluated at 3, 6, and 12 months after commencing treatment. They were asked to report their current tinnitus disturbance using TRQ. After 6 months of treatment, the results demonstrated that NTT was significantly more effective in improving tinnitus symptoms than the two reference groups. Patients receiving NTT presented a mean TRQ improvement of 66%, compared with mean of 22% for patients receiving noise plus counseling, and a mean of 15% for those receiving only counseling [14]. By 12 months, the mean gains of the neuromonics group had improved significantly more, but not for the other two groups. In terms of mechanisms, the study demonstrated that counseling was helpful (at a similar level to other studies),
and the use of one-size-fits-all-type noise made a further improvement, but the highly tailored intermittent acoustic stimulation was far more effective. In terms of consistency of response, the proportion of patients who had a clinically significant improvement (TRQ improvement > 40%) was 86% of the neuromonics group, 47% of the noise and counseling group, and 23% of the counseling alone group.

The third clinical trial included 35 individuals with clinically significant tinnitus distress [13]. Participants were randomly allocated into two groups: the first group, called the one-stage group, received intermittent tinnitus interaction throughout the 6-months program, while the second group, or two-stage group, received high interaction for 2 months. Then it moved to intermittent interaction for the last 4 months. Both groups had the same structured support program and followed the suitability criteria discussed earlier in this chapter.

All the audiometric and psychometric measures were performed at pretreatment, 2, 4, and 6 months after commencing the program. The results suggested that patients from both groups improved significantly, both statistically and clinically. At the conclusion of the 6-month program, the TRQ scores showed that 91% of the participants displayed an improvement of their tinnitus disturbance greater than 40%, with a mean improvement of 65%. The two-stage group was found to have a faster response to treatment. In a questionnaire administered after 12 months of commencing the treatment, more than three-quarters of the patients reported that the treatment had provided relief and increased their general well-being by a moderate or large amount [13]. This Ear and Hearing article compared the neuromonics results to a similarly randomized and controlled trial of TRT and masking [31]. It found that at 6 months, the neuromonics group had a higher proportion of patients having a clinically significant response (91%) compared to the TRT group (29%); it took the TRT group 18 months to achieve a more comparable outcome of 74%. Similarly, the TRT group required 15 h of clinician time per patient, whilst the neuromonics group required less than half the total amount of clinician’s time per patient.

A team of tinnitus specialists over eight centers across the US, led by the Cleveland clinic, has recently reported the 6-month post-therapy preliminary results of their study of the effectiveness of NTT [32]. Their dataset of 45 patients displayed this level of improvement of their tinnitus disturbance) to the prior Australian clinical trials and private practice results [16], and independently replicated those results. Another independent study reported on the long-term outcome of NTT [33]. The results of this report revealed that more than 85% of tinnitus patients treated with NTT sustained the full benefits of treatment 6–24 months after concluding the program.

Conclusion

Based on results from the clinical trials and outcomes showed by the private clinics, NTT can promote a major desensitization of tinnitus perception in a high proportion of patients. Among its advantages are that it is non-invasive, easy and pleasant to use, suitable for patients with a wide range of hearing and tinnitus characteristics, and it is not relatively time consuming for the clinician.

References


Chapter 76
Middle Ear Implantable Devices in Tinnitus Treatment

Eberhard Biesinger and Manuela Mazzoli

Keypoints

1. Tinnitus is often associated with high-frequency hearing loss.
2. Rehabilitation with hearing aids has shown effectiveness in reducing tinnitus. However, in some individuals with severe high-frequency hearing loss, classical hearing aids are not always able to amplify the high frequencies sufficiently and provide enough power.
3. Active middle ear implants are an alternative to conventional hearing aids that allow more power delivered to the cochlea, especially at high frequencies, and can also be used when middle ear ossicles are damaged.
4. A study supported by the Tinnitus Research Initiative (TRI) and MED-EL of the effect of a middle ear implant showed that individuals with severe tinnitus and high-frequency hearing loss achieved relief of their tinnitus after implantation.
5. Some patients had complete relief of their tinnitus after activation of the middle ear implant. Similar effects cannot be achieved by conventional hearing aids.
6. Individuals who have significant residual inhibition of their tinnitus and high-frequency hearing loss seem to be the best candidates for implantations.

Keywords Tinnitus • Middle ear implants • High-frequency hearing loss

Abbreviations

CE Conformity mark in the European economic market
DACS Direct acoustical cochlear stimulation
FDA Food and Drug Administration
FMT Floating mass transducer
MET Middle ear transducer
SR Stochastic resonance
TBF Tinnitus Beinträchtigungs Fragebogen (engl: tinnitus handicap inventory)
THI Tinnitus handicap inventory
TRI Tinnitus Research Initiative

Introduction

Individuals with tinnitus, who also suffer from hearing loss, often benefit from amplification. The use of hearing aids in tinnitus patients may make the patient less aware of the tinnitus as well as improve communication by reducing masking by the tinnitus. Hearing aids may also reduce the tinnitus, because they provide input to the nervous system that may reverse some of the plastic changes from deprivation of sound that has caused tinnitus and may counteract the deprivation of sound that causes some forms of tinnitus.

It has been reported that up to 67% of individuals who received unilateral hearing aids and 69% of individuals who received bilateral hearing aids report improvement in their tinnitus [1].

The quality of sound contributes many aspects. The effect of the pinna and the resonance in the external auditory canal contribute to optimize gain at higher frequencies [2].
Hearing aids often increase the perceived quality (color, crispness, clarity, pureness) of sounds, which could be important in reducing tinnitus annoyance, but unfortunately is not measured in routine clinical practice and is difficult to define.

Today’s digital hearing instruments are very advanced, offering maximum performance and reducing many of the difficulties encountered in earlier designs. In past years, improvements in hearing devices have substantially helped control feedback, widening the frequency range, and, to some degree, have improved sound quality. However, some individuals still experience the stigma and practical problems of using these devices.

Traditional hearing aids lack amplification of high frequencies (above 6,000 Hz) and fail to provide sufficient power. This is a problem in connection with suppression of tinnitus, which requires that high-frequency sounds are delivered to the ear at sufficient intensity.

Good reproduction of high-frequency sounds is also necessary for directional hearing and hearing when background noise is present.

Using a conventional “loudspeaker” at the end of the amplification chain seems to be the limiting factor for a sophisticated development of these devices.

Relocating the loudspeaker to the outer ear canal increased the performance of amplification in the high-frequency range.

Recognizing these problems and the fact that sound quality will always be an issue for those who use traditional hearing instruments and individuals with tinnitus, promoted the development of active middle ear implants. This has solved many of the problems of traditional hearing aids. It was therefore of great advantage in the treatment of some forms of tinnitus, occurring together with hearing loss, when devices that provide sound delivered directly to the middle ear bones or directly into the cochlea were developed. The amplification and the power that can be delivered to the cochlea using such devices exceed those of conventional hearing aids. Particularly, amplification is achieved in a larger frequency range than what is possible using traditional hearing aids.

With customized active middle ear implants, there is no need for a “loudspeaker” (receiver), thus reducing the distortion and reduction in the quality of sounds that occurs in traditional hearing aids. Furthermore, the ear canal is never occluded when implantable hearing aids are used.

**History of Implantable Hearing Aids**

Middle ear implants started in 1935 when Wilska [3] experimented with iron particles placed on the tympanic membrane. Wilska generated a magnetic field from an electromagnetic coil inside an earphone, which caused the iron filings to vibrate in synchrony with the magnetic field. This vibration in turn caused the eardrum to vibrate and allowed sound to be transduced to the cochlea in normal fashion. Later, Rutschmann (1959) [4] successfully stimulated the ossicles by gluing 10-mg magnets onto the umbo. An electromagnetic coil created a magnetic field that caused the ossicles to vibrate. Devices actually placed into the middle ear did not appear until 1970s [5].

Today, three general types of transducers are used in middle ear implants, each with advantages and disadvantages related to power, performance, frequency range, and reliability. The types of transducers used in middle ear implants consist of piezoelectric, electromagnetic, and electromechanical transducers.

Yanagihara and his colleagues [6] described an implantable piezoelectric device attached to the head of the stapes and performed the earliest human trials using these devices [7–12]. Their device was intended for patients with conductive and sensorineural loss.

A totally implantable piezoelectric device, known as the Esteem Hearing implant [13], was developed by St. Croix Medical, Inc. (now Envoy Medical Corporation) (Fig. 76.1).

Electromagnetic transduction devices consist of a magnet and an energizing coil. The magnet is attached to the ossicular chain, tympanic membrane, or the inner ear (round window or oval window). Specific experiences with regard to the influence on tinnitus have not been published.

Another implantable middle ear device known as Carina™ is shown in Fig. 76.2.

**Figure 76.3:** Otologics MET fully implantable middle ear device (Carina™, Photo: Otologics). Reproduced with permission from OTOlogics GmbH, Heidelberg, Germany.

**Fig. 76.3:** Otologics MET fully implantable middle ear device (Carina™, Photo: Otologics). Reproduced with permission from Med-EL, Innsbruck.

**The Soundbridge**

Soundbridge is the middle ear implant with the longest clinical experiences, 3,000 patients so far (2009). It was first marked by Symphonix Devices in San Jose, California, as the Vibrant Soundbridge. It has received both European CE-mark in March 1998 and FDA approval in the U.S. in August 2000 [14–16]. However, the company went out of business in 2002 only to return in March 2003 as the Med-EL Vibrant Soundbridge.
The semi-implantable device consists of an outward audio processor which is placed over the implanted coil and magnet. The coil is linked by a golden wire to the floating mass transducer (FMT) (Fig. 76.3). The frequency range is 1,000–8,000 Hz, but technically amplification up to 16,000 Hz is possible.

In the last 2 years, the Vibrant Soundbridge has assumed particular importance through the fact that the FMT can also be implanted in the round window [17] (Fig. 76.4). The indication here refers to a destroyed middle ear, such as after removal of the petrosal bone, malformations, cholesteatoma, sclerosis of the footplate, etc. The FMT provides a better way to induce sound energy into the cochlea than using the ossicular chain.

A special form of implanting the FMT was achieved by Hüttenbrink with “TORP-Y-Vibroplasty” [18].

Tinnitus-Related Clinical Observations and Studies

In the ENT Clinic in Traunstein, 52 patients have been equipped with the implant since 1998 (four of them bilaterally). All patients were provided with conventional hearing aids before and, for different reasons, were not content with these devices. All patients continue to use their middle ear implant as of August 2009 without any technical problems. It was surprising that most patients, who simultaneously suffered from tinnitus, reported that the middle ear implant largely reduced their tinnitus which could not have been achieved by traditional hearing aids.

In 2000, a patient implanted on both sides with middle ear implants reported that this tinnitus disappeared completely after activating the implant. Six years later, inspired by the results obtained by the middle ear implant in this person, the Tinnitus Research Initiative
E. Biesinger and M. Mazzoli (http://www.tinnitusresearch.org) offered a grant to study the effect of the middle ear implants on tinnitus. The participants had sensorineural hearing loss at high frequencies and tinnitus and had been given middle ear implants. All patients received the Vibrant soundbridge. They were studied for 1 year using a visual analogue scale (VAS), Goebel–Hiller score [19], and the Tinnitus Handicap Inventory (the German TBF-12 [20, 21]).

The first patient was implanted in January 2007 on the left side. His audiogram showed severe hearing loss on the left side and minor hearing loss on the right side (Fig. 76.5). The combination, with a good dynamic range assessed by the level of discomfort, gave a good indication for implantation.

After the operation, the audio processor was activated, and the reaction of his tinnitus was surprising: the tinnitus shifted from his left side to his right side. Figure 76.6 shows the result: because of the remaining tinnitus, there was no improvement regarding annoyance after 2 months, and the person did not develop any habituation. Facing the fact that this individual now had tinnitus on the right side, we also implanted the right side with the Soundbridge 12 weeks after the original implantation. With the activation of both audio processors, the annoyance due to tinnitus diminished, and the quality of life improved.

Figure 76.7 shows the functional gain (green line) after the implantation of the Soundbridge in both ears. The patient describes that his tinnitus decreases already by switching on the device, although he is not able to hear the receiver noise. This indicates that in addition to a masking effect, there might be other effects from the implanted device.

At the 1-year follow-up exam (patient “M.A.”), this patient is no longer annoyed by tinnitus when the audio processor is activated.

The first patient of a new study, sponsored by the company Med-El® with five participants with unilateral tinnitus and reproducible residual inhibition, received a Soundbridge implantation in June 2008. The device
Fig. 76.5  Audiogram participant “B.A.”
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Fig. 76.6  One-year follow-up of participant “B.A.” with tinnitus questionnaires and VAS shows his improvement with tinnitus after implantation on both ears. Reproduced with permission from the authors

Fig. 76.7  Functional gains of both ears of participant “B.A.” Reproduced with permission from the authors
was activated and fitted in August 2008. Immediately after the fitting process, the patient’s tinnitus disappeared completely after switching on the device. Although they were once greatly bothered and annoyed by the tinnitus, the activation of the device gave complete relief.

Three other participants in this study reported similar effects.

**Conclusion**

Implantable hearing aids have shown to be effective in reducing tinnitus in individuals with severe hearing loss and tinnitus, where the hearing loss was caused by middle ear or cochlear pathologies. The reason the middle ear implantable devices provide relief of tinnitus may be masking, but it seems more likely that the benefit is caused because these devices provide effective activation of the auditory nervous system, and thereby counteract the effect of deprivation of sound input that had activated neural plasticity causing the tinnitus. This means that the effect of the implanted hearing aids on tinnitus is similar to that of cochlear implants (see Chap. 77).

One reason for the success of implantation might be that it facilitates residual inhibition.

**References**

Chapter 77
Cochlear Implants and Tinnitus

Andrea Kleine Punte, Olivier Meeus, and Paul Van de Heyning

Keypoints

1. Many forms of tinnitus are caused by deprivation of sounds, and electrical stimulation has been applied to the promontory for treatment of tinnitus, providing significant relief from tinnitus by supplying input to the auditory nervous system.

2. Immediate relief of tinnitus has been reported in approximately 82% of the patients and longer term tinnitus suppression in 45% of such treatment.

3. Cochlear implants, therefore, may offer long-term tinnitus suppression in patients with severe sensorineural hearing loss by providing input to the auditory nervous system.

4. This chapter provides evidence of tinnitus relief in up to 90% of individuals with severe tinnitus following cochlear implantation.

5. An indication for the use of cochlear implants in individuals who are deaf in one ear while having incapacitating tinnitus on that side is provided in this chapter.

6. Research in the field of cochlear implants and tinnitus is discussed, and suggestions for future research are made.

Keywords Tinnitus • Cochlear implants • Promontory stimulation • Treatment

Abbreviations

- EPS Electrical Promontory Stimulation
- SNHL Sensorineural Hearing Loss
- SSD Single-sided Deafness
- VAS Visual Analogue Scale

Introduction

Tinnitus is one of the most common otological complaints, affecting 10–15% of the adult population (see Chap. 5). Various treatments have been developed to suppress or reduce tinnitus (see chapters in Sect. 5). Many forms of tinnitus are now thought to be caused by auditory deprivation (see Chap. 11), and hearing aids can, therefore, provide relief from tinnitus in some individuals. Reduction of tinnitus following the use of a hearing aid was first reported in 1947 [1]. Later, several studies confirmed the beneficial effect of hearing aids for tinnitus relief. In 1981, a significant improvement of the value of binaural aids compared to monaural hearing aids in reduction of tinnitus and associated problems was reported. The improvement was present in almost half of the individuals surveyed [2]. Similar conclusions were drawn from another study by Surr et al. [3], in which approximately half of the respondents with tinnitus reported that their hearing aids provided either partial or total relief from tinnitus. Individuals rating their tinnitus as being severe reported partial relief of tinnitus rather than total relief, but other studies showed no effect of hearing aids on tinnitus [4].
Hearing aids are not useful for treatment of individuals with severe sensorineural hearing loss (SNHL) and tinnitus, but if the auditory nerve is preserved, electrical stimulation of the inner ear can supply necessary auditory input in deaf individuals.

Electrical promontory stimulation (EPS) seems to be a promising tinnitus treatment, providing significant relief. Research on EPS shows at least temporary and partial tinnitus suppression. Immediate relief of tinnitus has been reported in approximately 82% of patients and longer term tinnitus suppression in 45% of these patients [5]. Rubinstein et al. [6] also described the effect of high-frequency EPS on tinnitus, and the authors advocated that the effect should be investigated with an implantable device. There are indications that cochlear implants may provide long-term tinnitus suppression in individuals with severe sensorineural hearing loss. Cochlear implants have been reported to provide tinnitus relief in up to 90% of patients. There is evidence that deafferentation of the auditory pathway plays an important role in causing tinnitus (see Chaps. 10 and 11), and that the effect can be reversed by electrical stimulation of the auditory system via EPS or through cochlear implants. A particularly new indication for cochlear implants is single-sided deafness (SSD) with concomitant incapacitating tinnitus [7]. In this chapter, results of studies of the use of cochlear implants for treatment of tinnitus are discussed and suggestions for future studies are made.

**Electrical Promontory Stimulation and Tinnitus**

Nearly 200 years ago, electrical stimulation was first described as possible tinnitus treatment. Only in the 1960s and 1970s, the potential beneficial effect of electrical stimulation on tinnitus was rediscovered. Feldmann [8] reported suppressed tinnitus by Volta’s platinum–zinc cell. Since then, electrical stimulation as treatment for profound SNHL and tinnitus has been widely investigated. Originally, electrical stimulation of the cochlea was used to assess the integrity of the neural structure in the cochlear prior to cochlea implantation. A side effect of this test in some cases was a suppression of the accompanied tinnitus [9–13].

Electrical stimulation of the cochlea is possible with EPS or round window stimulation. In EPS, a needle electrode is placed on the promontory in order to stimulate the cochlea. This technique has been investigated thoroughly and is used pre-operatively to predict speech reception results with a cochlear implant (CI) [14–16]. An overview of the literature of tinnitus suppression with EPS [17–19] is given in Fig. 77.1.

Portmann et al. [20] suggested that the effectiveness of electrical stimulation depends on the electrode placement and electrical stimulation at the round window was better than promontory stimulation. Also, temporary tinnitus suppression was most effective

**Fig. 77.1** Tinnitus modulation after electrical stimulation of the inner ear
when using positive electrical pulses. In the reported studies, the efficacy of EPS for suppressing tinnitus was done using stimulation for only a very short time in acute experimental set-ups. Repeatability of the tinnitus suppression remains unclear, and the long-term effects of EPS on the cochlea and acoustic thresholds have not been thoroughly investigated [6].

Cochlear Implant for Bilateral Profound Hearing Loss and Tinnitus

Many people who have bilateral profound sensorineural hearing loss have severe tinnitus. The first report showing that suppression of tinnitus could occur after cochlear implantation was published in 1976 by House [21]. Baguley [22] and Quaranta [23] reviewed the results of studies of suppression (or modulation) of tinnitus after cochlear implantation. Recent studies provided additional support of these findings [24, 25].

Figure 77.2 summarizes the results obtained concerning tinnitus modulation after cochlear implantation [24–32]. It is clear that tinnitus decreased after cochlear implantation in most of the people receiving these implants as treatment. For these individuals, the tinnitus was a secondary complaint to the main problem of deafness.

Cochlear Implant for Single-Sided Deafness and Incapacitating Tinnitus

A small group of people have suffered from SSD, and due to this deafness, incapacitating tinnitus developed. The tinnitus was referred to the deaf ear, with the other ear having normal hearing or showing only moderate hearing loss and no tinnitus. At the Antwerp University Hospital, such individuals received a cochlear implant in the deaf ear in order to reduce tinnitus and also to restore some hearing (Medel Combi 40+ with an M-electrode or Pulsar CI100 with Flexsoft electrode). We studied these individuals in a prospective clinical study to assess the long-term effects of cochlear implantation on tinnitus in people with SSD and ipsilateral incapacitating tinnitus [7]. Twenty-one individuals who received a cochlear implant and suffered from severe incapacitating tinnitus that was unresponsive to other treatments participated in this study. Tinnitus loudness was measured using a Visual Analogue Scale (VAS); loudness perception of tinnitus was recorded with the CI both activated and deactivated. Tinnitus distress was measured using the Tinnitus Questionnaire (TQ) pre- and post-operatively.

All 21 patients reported a subjective benefit when the cochlear implant was activated. Tinnitus loudness was reduced significantly after cochlear implantation.

Fig. 77.2 Tinnitus modulation after cochlear implantation
At the 12-month follow-up exam, the loudness of their tinnitus had decreased from an average of 8.5 to 2.5 on the VAS (of 0–10). Also, the tinnitus questionnaire (TQ) total score decreased significantly. Figure 77.3 shows the average tinnitus loudness as a function of time.

Cochlear Implantation seems to be a successful treatment of severe tinnitus in patients with SSD. A significant suppression of tinnitus occurred already after 1 month of cochlear implantation. All patients but one had a tinnitus score lower than 5/10 on a VAS. The tinnitus largely recurred when their cochlear implant was deactivated.

The results of this study show that after a long period, tinnitus does not reoccur, and there is no adaptation of the tinnitus to the electrical stimulation presented by the cochlear implants. Long-term results up to 48 months after cochlear implantation also suggest cochlear implantation provides durable tinnitus relief in these individuals (Kleine Punte et al.) [33]. It must, however, be emphasized that other causes of tinnitus have to be excluded before cochlear implantation for tinnitus is recommended and severe depression is a contra-indication.

Experience from the use of cochlear implants to treat patients with tinnitus indicates that electrical stimulation of the auditory nerve can reverse the reorganization associated with peripheral deafferentation that causes tinnitus and thus, reverse plastic changes that may have caused the tinnitus (see Chap. 12). Also, the increase in activation of the auditory nerve may provide inhibitory influence on the cells in the auditory nervous system, which may play a role in its effect on tinnitus. Enhanced attentiveness to environmental sounds could contribute to the observed suppression of tinnitus. The results from this study suggest that inhibition of tinnitus by cochlear implants is stable, and tinnitus does not return over time. This long-term stability suggests that cochlear implants may permanently suppress tinnitus in these patients. Besides providing significant tinnitus relief, patients with SSD also experienced an improvement in their hearing capabilities after cochlear implantation [34].

It should be taken into account that other factors may also be responsible for the tinnitus relief obtained after cochlear implantation. Psychological factors may have an influence on tinnitus loudness and tinnitus annoyance: the long inclusion procedure before the implantation includes thorough psychological assistance, which may increase the well-being of the patient. However, in the months before cochlear implantation, no attenuation of tinnitus occurred in our study group, which is consistent with the Blue Mountain follow-up study [35]. Finally, an increased assurance after a recuperation of the auditory function may also have contributed to diminishing the tinnitus annoyance.

![Figure 77.3](image-url)

**Fig. 77.3** Average tinnitus loudness and standard deviation with CI activated as function of time after first CI fitting.
Development of Tinnitus After Cochlear Implantation

Although in the majority of cases, cochlear implantation results in an abolishment or suppression of tinnitus, a small percentage of individuals have been reported to develop tinnitus or experience an increase of their tinnitus after cochlear implantation. The reported incidence ranges from 0 to 9% [22, 30, 36–38]. Akdogan et al. [39] investigated the tinnitus properties due to cochlear implantation and found that 4 out of 17 patients (23.5%) developed tinnitus after cochlear implantation. The mean tinnitus loudness was 17.5 dB SL. Quaranta et al. [24] reported that 7 out of 41 individuals (17%) developed tinnitus immediately after the insertion of the electrode array for a cochlear implant. However, 1 month later, only two of these individuals still perceived the tinnitus they acquired at the time of the implantation and their tinnitus became mild. Although the risk of getting tinnitus from cochlear implantation is minimal, it is important to counsel candidates about the risk of cochlear implants.

Suppression of Bilateral Tinnitus After Unilateral Cochlear Implantation

A few reports about bilateral tinnitus suppression after unilateral cochlear implantation have been published. A study by Di Nardo et al. [25] reported complete tinnitus suppression bilaterally in 4 of 9 (44%) individuals who received a cochlear implant in one ear, while four others (44%) experienced bilateral attenuation of tinnitus. Another study performed in 14 individuals who had bilateral tinnitus before implantation reported bilateral suppression of their tinnitus or attenuation of the tinnitus in 12 (86%), while the tinnitus increased bilaterally in 2 (14%) of the participants in the study [32].

The effect of cochlear implantation on bilateral tinnitus was more extensively described in a study performed in 41 individuals with bilateral tinnitus [24]. When their cochlear implant was turned on, the tinnitus was abolished bilaterally in 23 participants of the study (56.1%). Tinnitus was completely suppressed in the implanted ear only in 4 (9.7%) individuals and contralaterally in 4 (9.7%) of the participants. That means tinnitus suppression occurred in 31 (76.5%) participants in this study.

Patients with bilateral tinnitus sometimes experience tinnitus suppression after sequential bilateral cochlear implantation [40]. Increase of tinnitus was also described after bilateral cochlear implantation, similar to tinnitus aggravation after unilateral cochlear implantation.

Future Research

Although many reports of tinnitus suppression after cochlear implantation have been published, more detailed studies are needed to assess the advantages of cochlear implants for treatment of tinnitus. Since double-blind studies are impossible in the field of cochlear implantation, the emphasis should lie on conducting randomized controlled studies on the effects of cochlear implantation on tinnitus. Also, the working mechanisms of tinnitus suppression after cochlear implantation are not totally understood and need to be further explored in studies of humans.

Conclusion

Tinnitus can be influenced by electrical stimulation of the inner ear, at least when the tinnitus occurs in connection with sensorineural hearing loss. Transtympanic electrical promontory stimulation or round window stimulation can provide temporary tinnitus relief. In individuals with profound hearing loss, cochlear implantation can provide more permanent tinnitus suppression.

Table 77.1 Inclusion and exclusion criteria of cochlear implantation as tinnitus treatment in patients with SSD

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Severe tinnitus: tinnitus loudness 6–10 on VAS for &gt;6 months</td>
<td>Major depression</td>
</tr>
<tr>
<td>Subjective tinnitus due to ipsilateral profound SNHL</td>
<td>Not willing to attend regular follow-up and rehabilitation</td>
</tr>
<tr>
<td>Tinnitus as primary complaint</td>
<td>Duration of tinnitus &gt;10 years</td>
</tr>
<tr>
<td>Standard treatments of tinnitus have no effect</td>
<td>Realistic expectations</td>
</tr>
<tr>
<td>Normal hearing to moderate hearing loss contralaterally</td>
<td>Patent Scala tympani</td>
</tr>
</tbody>
</table>
References

27. Itu J, Sakakihara J. Tinnitus suppression by electrical stimulation of the cochlear wall and by cochlear implantation. Laryngoscope 1994;104(6 Pt 1):752–4
33. Kleine Punta A, Vermeire K, Hofkens A Van de Heyning P. Durable tinnitus reduction and bimodal hearing after cochlear implantation in single-sided deafness; results up to 48 months post-implantation 8th Wullstein Symposium, Treatment of Unilateral Deafness Wurzburg, 2009
Chapter 78
Pharmacological Approaches to Tinnitus Treatment

Ana Belén Elgoyhen and Berthold Langguth

Keypoints

1. Available treatments for the management of tinnitus are diverse.
2. Although most patients benefit from treatment to some degree a large percentage of them are left untreated and in despair with the notion that “they have to learn to live with their tinnitus”.
3. Currently there is no drug that is approved by the Food and Drug Administration (FDA) or the European Medicines Agency (EMEA) for the treatment of tinnitus. Thus, tinnitus is still a clinically unmet need, and most patients would welcome a drug that abolishes their phantom sound once and for all.
4. There are different forms of tinnitus which probably differ in their response to pharmacological treatment. Thus, even if a specific drug has failed to demonstrate efficacy in controlled clinical trials in a large sample, a beneficial effect in a subgroup of tinnitus patients should not be precluded. At present, most evidence-based pharmacological treatments treat specific comorbidities rather than the core of the disorder itself.
5. There is an urgent need for effective treatment approaches. Since in some individuals, tinnitus causes irritability agitation, stress, depression, insomnia, and interferes with normal life – leading to suicidal attempts in severe cases – even a drug that produces a small but significant effect would have an enormous therapeutic impact. This review describes strategies currently available for tinnitus pharmacotherapy.

Keywords Tinnitus • Phantom sound • Lidocaine • Neramexane • Hearing • Noise trauma

Abbreviations

EMEA European Medicines Agency
RNID Royal National Institute for Deaf People (UK)
SSRI Selective serotonin reuptake inhibitors
THQ Tinnitus handicap questionnaire
SNRI Serotonin-norepinephrine reuptake inhibitors
GABA Gamma amino butyric acid
PDE5 Phosphodiesterase type 5
NMDA N-methyl-D-aspartate

Introduction

Available treatments for the management of tinnitus are diverse. These include counseling and cognitive behavioral therapies; different forms of sound therapies; methods that attempt to increase input to the auditory system, such as hearing aids and cochlear implants (for use in patients whose tinnitus is caused by deprivation of signals to the auditory nervous system); neurobiofeedback; and various forms of electrical stimulation of brain structures, either through implanted electrodes or by inducing electrical current in the brain with transcranial magnetic stimulation and drug treatments [1–9]. Although most patients benefit to some degree from the above-mentioned therapies, a big fraction of them are
left untreated and in despair with the notion that “they have to learn to live with their tinnitus”. Thus, tinnitus is still a clinically unmet need. Most patients would welcome a drug that abolishes their phantom sound once and for all.

The market for a drug specifically for tinnitus relief is huge. There have been numerous patents filed worldwide on drugs with the potential to offer relief. Furthermore, tinnitus can be found attached to many more patents filed on molecules aimed at a range of diverse therapeutic classes. The Royal National Institute for Deaf People (RNID) in the UK estimates that a novel tinnitus drug could have a product value of US $689 million in its first year of launch [10]. However, we are still left without a single FDA-approved drug on the market targeting tinnitus relief. For the majority of tinnitus sufferers who seek medical advice, the treatment goals are aimed at symptomatic relief of their phantom sound and/or the management of the associated distress. This approach is usually justified, as serious underlying pathologies are rare. Over four million prescriptions are written each year for tinnitus relief in Europe and the US, but these are all off-label prescriptions from a wide variety of therapeutic drugs (Table 78.1) [10]. Most clinicians who treat tinnitus patients urge for an effective drug therapy targeted at tinnitus. Thus, there is a tremendous need both from patients and medical doctors to develop a drug targeting tinnitus relief. Since in some individuals, tinnitus causes irritability, agitation, stress, depression, insomnia, and interferes with normal life — leading to suicidal attempts in severe cases [11] — even a drug that produces a small but significant effect would have an enormous therapeutic impact. However, ideally a disappearance of tinnitus should be the ultimate goal of any research and development platform toward designing a tinnitus drug.

The lidocaine experience: tinnitus can be pharmacologically targeted.

Tinnitus is a symptom that is associated with virtually all diseases and disorders affecting the auditory system and can arise from a lesion in any part of the auditory pathway. Some causes that trigger tinnitus are well known. In particular, noise traumas, administration of ototoxic drugs, and head and neck injuries have been associated with the development of subjective tinnitus. Interestingly, while the initial lesion might affect the peripheral organ of the auditory system, the neural correlate of the sound perceived is most likely in the central auditory circuitry [12, 13] and involves non-auditory brain areas [11, 14, 15]. A central origin of the tinnitus percept is demonstrated by the fact that the phantom sound sensation persists after deprivation of input from the periphery via sectioning of the auditory nerve [16]. Although the mechanisms of the production of tinnitus are far from being fully understood, there is growing evidence that changes in neuronal activity, neuronal synchrony, disruption of the balance between excitation and inhibition, and rearrangements of the tonotopic organization in different

<table>
<thead>
<tr>
<th>Substance</th>
<th>Number of prescriptions in 2001 (in thousands)</th>
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<tr>
<td>Ginkgo biloba</td>
<td>782</td>
</tr>
<tr>
<td>Trimetazidine</td>
<td>650</td>
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<tr>
<td>Betahistine</td>
<td>314</td>
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<td>Pentoxifylline</td>
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<tr>
<td>Piracetam</td>
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<td>Naftidrofuryl</td>
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<td>Buflomedil</td>
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<td>Cinnarizine</td>
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<td>Clonazepam</td>
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<td>Nicergoline</td>
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<td>Flunarizine</td>
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<td>Nimodipine</td>
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<td>Acetylsalicylic acid</td>
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<td>Hetastarch</td>
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<td>Ajmalicine</td>
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<td>Moxaverine</td>
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<td>Caffeine</td>
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<td>Cyclandelate</td>
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<td>Gold</td>
<td>10</td>
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<td>Viscum album</td>
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<tr>
<td>Hypericum perforatum</td>
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</table>
parts of the auditory pathway, including the dorsal cochlear nucleus, inferior colliculus, thalamus, and/or auditory cortex underlie tinnitus pathology [12, 15, 17–20]. Neuronal excitability can be modulated by different neurotransmitters, neuremodulators, and voltage-gated channel acting compounds [21–25]. Thus, there is no reason to believe that tinnitus cannot be pharmacologically approached. The fact that the voltage-gated sodium channel blocker lidocaine that is used as a local anesthetic and an antiarrythmic, given intravenously, leads to the temporary disappearance of tinnitus or a major change in the nature of the tinnitus in 70% of patients [26–28], indicates that activity-driven changes underlying tinnitus can be pharmacologically targeted.

Although intravenous lidocaine seems to be effective in a great number of tinnitus patients, the effect is short lasting and the route of administration is not a practical one in a clinical setting of a chronic condition. Moreover, side effects are considerable and include cardiac arrhythmia, drowsiness, dizziness, confusion, and restlessness [29]. Several other oral antiarrythmic drugs like tocainide, flecainide, and mexiletine have been studied for tinnitus. None of these compounds have been demonstrated to be particularly useful. Almost all of these exhibit severe side effects and are now in disuse [6–8].

Given the positive results with lidocaine, the possibility of novel local anesthetics like tonicaine and sameridine with longer duration of action [30] has been suggested [31]. Tonicaine is a quaternary ammonium and most likely does not efficiently pass the blood–brain barrier. Thus, its use in tinnitus is probably limited. Sameridine has additional μ-opioid antagonist properties [30], a biological system that has not been targeted so far as a treatment for tinnitus, and a possible avenue to explore. Moreover, alternative delivery systems with prolonged duration of action, as has been suggested for the treatment of pain with a liposomal bupivacaine formulation and intradermal injection [32], could be tested. A preliminary report has shown some positive results in tinnitus patients with an intradermal lidocaine injection [33]. The pharmaceutical company Epicept (http://www.epicept.com) has a lidocaine patch formulation currently at Phase II for tinnitus. However, it has to be considered that the systemic concentrations that can be reached with the currently available lidocaine patches are probably much lower than the concentrations needed to suppress tinnitus.

**Pharmacological Treatment of Tinnitus**

The management of tinnitus sufferers is a pressing need faced by medical doctors in their daily practice. Drug therapy is one approach to the problem, and a large variety of different drugs are used (Table 78.1), even if supporting evidence is scarce. Since tinnitus is a symptom that might be the manifest of different underlying pathologies and has several etiologies and comorbidities – which can include various degrees of affective disorders – heterogeneity within tinnitus patients is expected [11, 34]. Thus, the pharmacological treatment of tinnitus faces the “one drug won’t fit all” scenario. Once (and if) the different forms of tinnitus are established, different treatment approaches will be devised. Up front, there is a consensus among clinicians treating tinnitus that the pathophysiology underlying the onset of tinnitus may differ from that of chronic tinnitus. Therefore, treatment approaches will probably vary with the duration of the disease. Immediately after tinnitus onset, more casually oriented treatment approaches might be possible, and involvement of the cochlea might still be important. In case of abrupt onset tinnitus associated with sudden hearing loss or noise trauma, treatment strategies which restore hearing function are expected to have beneficial effects on tinnitus. There is no clear boarder between “acute” and “chronic” tinnitus. The currently used distinction is arbitrary and varies between 3 and 6 months. Furthermore, recent data suggest that the neural correlate of tinnitus might change even after a duration of several years [15].

**“Acute State”**

One form of acute tinnitus that deserves special attention is associated to sudden hearing loss. Sudden hearing loss is characterized by the abrupt loss of hearing, typically unilateral [35–37]. A proportion of patients also experience dizziness and tinnitus with the hearing loss. Most individuals are able to pinpoint the precise moment of hearing reduction, such as awakening with the symptom. No specific sound frequency region in the cochlea appears to be preferentially affected, and the severity of hearing loss ranges from mild to profound. The spontaneous recovery rate is high; up to
65% of patients may experience recovery of pre-loss hearing. The high variability of the spontaneous course probably reflects the fact that there are different forms of sudden hearing loss with different etiologies. Among others, vascular, inflammatory, and infectious mechanisms are probably involved. However, in most cases, the exact etiology remains unknown. Many potential prognostic factors have been identified. Vertigo, persistent or profound hearing loss, and prolonged duration of hearing loss are negative prognostic factors. Subjects with good recovery usually experience that recovery within 2 weeks of the onset of hearing loss. Proposed treatments have included systemic or intratympanic steroids [36–38], vasodilators [39], antiviral agents [40, 41], and hyperbaric oxygen [42, 43]. Although some studies report positive results in subset of patients, none have proven effective in well-controlled studies [35, 41, 42].

Acute tinnitus associated to noise-induced hearing loss of abrupt onset such as that produced by exposure to a blast or after a rock concert also deserves special consideration. Noise is the greatest causative factor among the defined etiologies of tinnitus. Since the industrial revolution, an increasing number of people are being exposed to extreme levels of noise. Noise at levels of 85 dBA and higher can lead to both mechanical and metabolic damage of the cochlea [44]. Single, repeated, or continuous exposure to high levels of noise can cause noise-induced hearing loss and tinnitus. In developed countries, the appetite for leisure noise among the young (like attending rock concerts or discos or the use of MP3 players) is expected to have a substantial, deleterious impact on hearing loss and tinnitus incidence in older generations in the near future [10, 45, 46]. In a retrospective study of 3,466 claimants who sought compensation for occupational noise-induced hearing loss, the prevalence of those reporting tinnitus as a function of hearing loss at 4 kHz, ranged from 41.7 to 56.5%, regardless of the amount of hearing loss sustained [47]. Excessive noise can cause structural damage to the hair cell bundles and can generate excitotoxic effects on the sensory nerve terminals [48]. Compared to the millions of photoreceptors in the eye or the number of olfactory neurons in the nose, the number of sensory hair cells in the cochlea is extremely modest (~15,000). Hair cells die by apoptosis and unlike supporting cells in non-mammalian epithelia, mammalian supporting cells in the organ of Corti do not proliferate to replace lost hair cells, and they do not naturally change their phenotype [49]. Loss of hair cells leads to loss of spiral ganglion neurons, which depend on hair cells for the production of survival factors such as the neurotrophin NT-3 and brain-derived neurotrophic factor. Accumulation of free radicals, excitotoxicity mediated by glutamate receptors, and activation of apoptosis, are predictable players in the loss of cells [46]. Animal experiments show that growth factors and drugs directed against apoptosis, excitotoxicity, and oxidative stress can provide valuable protection from hearing loss and tinnitus if applied during exposure [50] and also probably immediately after exposure.

Otoprotectants at clinical development to prevent noise-induced hearing loss and associated tinnitus are various. In a double-blind placebo-controlled study involving 300 young and healthy military recruits, those supplemented daily with 4 g of oral Mg granulate (6.7 mmol Mg aspartate) showed significantly less permanent threshold shift 1 week post noise than those in the placebo control group (11.2% vs. 21.5%) [51]. In a recent study, normal-hearing adults were dosed orally with placebo or 900 mg of the glutathione prodrug N-acetylcysteine 30 min before entering a nightclub where they were exposed to 2 h of loud music. Personal dosimeters recorded a mean noise level of 98.1 dB (A-weighted). An average of 14 dB temporal threshold shift at 4 kHz was reported in subjects immediately

| Table 78.2 Off-label drugs investigated for the treatment of tinnitus |
|-----------------|-----------------|-----------------|
| • Antiarythmics | • Antidepressants |
| Lidocaine       | Amitriptyline   |
| Tocainide       | Trimipramine    |
| Flecaainide     | Nortriptyline   |
| Mexiletine      | Paroxetine      |
| • Anticonvulsants | Sertraline |
| Carbamazepine   | Fluoxetine      |
| Gabapentine     | • Dopaminergic  |
| Lamotrigine     | Sulpiride       |
| Valproic Acid   | Piribedil       |
| • Anxiolytics   | • Others        |
| Alprazolam      | Atorvastatin    |
| Clonazepam      | Cyclandelate    |
| Diazepam        | Furosemide      |
| • Glutamate receptor antagonists | Herbal products |
| Acamprosate     | Misoprostol     |
| Caroverine      | Melatonin       |
| Memantine       | Nimodipine      |
|                 | Vardenafil      |
|                 | Zinc            |
after exposure (within 15 min). No significant differences between groups were identified [52]. This observation might be related to the requirement of a high dose of N-acetylcysteine to effectively prevent noise-induced hearing loss in animal models, or possibly to the limited ability of the compound to prevent temporary threshold shifts. Several ongoing trials are being performed to test the efficacy of this compound [53]. Sound Pharmaceuticals is developing SPI-1005 (ebselen), an antioxidant. In Phase I studies, it has been shown to have a favorable toxicity and pharmacokinetic profile. The company is now testing the compound in Phase II trials with the Navy/Marine Corps. Auris Medical has a Phase II trial to test AM-101, an NMDA antagonist for the treatment of tinnitus derived from excitotoxicity in the cochlea due to noise trauma, and AM-111 (a JNK MAPK-mediated apoptosis blocker) at pre-clinical studies for the treatment of acute sensorineural hearing loss from acute acoustic trauma, sudden deafness, and inner-ear surgery.

Summarizing, there is consensus among clinicians that acute tinnitus deserves specific attention and that there might be a short therapeutic window for specific pharmacologic interventions. However, there are no treatments available, which have shown repeated efficacy in controlled trials. This might be due to etiologic heterogeneity of acute hearing loss and a high rate of spontaneous recovery. Probably, the most widely used treatment strategy is systemic and intratympanic steroid administration. Further clinical trials to validate treatments of acute tinnitus are urgently needed.

“Chronic Form”

Antidepressants

Antidepressants are commonly used in pharmacological protocols for the management of chronic tinnitus [6–8] (Table 78.2). The reason for such a large use of antidepressants can be found in the well-described comorbidity between depressive disorders and tinnitus. Among all antidepressants that have been investigated for tinnitus, a particular interest has been paid to tricyclic antidepressants, mainly because of their beneficial effects on chronic pain syndromes [54]. This property of several tricyclic drugs is interesting in view of the proposed etiological similarities between tinnitus and neuropathic pain (see Chaps. 14, 15 and 94) [55]. Among the tricyclic antidepressants analyzed (amitriptyline, trimipramine, and nortriptyline), nortriptyline is worth mentioning. In a small-scale, single-blind placebo-washout study involving patients with severe tinnitus and major depression, nortriptyline significantly reduced depression and tinnitus loudness (10 dB reduction) [56]. In a follow-up double blind-placebo-controlled study involving subjects with severe tinnitus and severe depression or depressive symptoms, nortriptyline significantly reduced depression scores, tinnitus disability scores, and tinnitus loudness (6.4 dB reduction) relative to placebo [57]. There was a significant correlation between the reduction in tinnitus disability scores and depression scores, suggesting that nortriptyline is effective in reducing tinnitus loudness and severity in severely depressed tinnitus patients, but has less benefit in non-depressed individuals [58]. One study has compared amitriptyline with placebo and found after 6 weeks of 100 mg amitriptyline a significant reduction of tinnitus complaints and tinnitus loudness compared to the placebo group [59]. In another study, where amitriptyline was compared with biofeedback, 27.5% of patients reported improvement. However, this was less effective than biofeedback per se [60]. Trimipramine has been evaluated in a small double-blind placebo cross-over study, which did not demonstrate a difference between trimipramine and placebo treatment [61]. It should be noted that the induction of tinnitus with tricyclic antidepressants has been described [62–64].

Selective serotonin reuptake inhibitors (SSRI) such as paroxetine or sertraline have been tested. In a randomized double-blind placebo-controlled study of patients without severe hearing loss, but with depression, anxiety, and a high risk for developing severe tinnitus, sertraline was significantly more effective than placebo in reducing tinnitus loudness and tinnitus severity [65]. In a double-blind, placebo-controlled study involving chronic tinnitus patients, few of whom suffered from depression, the paroxetine group showed little difference from placebo on tinnitus loudness matching, tinnitus handicap questionnaire (THQ) scores, and other measures; however, the paroxetine group showed a significant improvement on tinnitus aggravation compared to the control group [66]. The combination of paroxetine with vestipitant and
vestipitant alone is currently undergoing a phase II clinical trial for the treatment of tinnitus (http://clinicaltrials.gov/ct2/show/NCT00394056) by GlaxoSmithKline. Vestipitant is a novel neurokinin-1 substance P receptor antagonist. Substance P receptor antagonists have been shown to be effective with pain [67]. Information on the clinical effectiveness of these drugs is currently unavailable. Very little has been reported for serotonin norepinephrine reuptake inhibitors (SNRI), such as duloxetine and venlafaxine, or for the dual acting drug mirtazapine. Since activity on norepinephrine reuptake is considered necessary for an antidepressant to be effective on neuropathic pain [54], it might be worthwhile to investigate this group of drugs for its use in tinnitus treatment.

It has to be considered that the scales used for the measurement of tinnitus correlate highly with depression scales. Thus, the observed reduction of tinnitus severity under antidepressant treatment might, at least to some extent, be a pure consequence of the antidepressant effect of the investigated drugs. Nevertheless, available data provide converging evidence that tinnitus patients with depression and anxiety may gain benefit from antidepressant treatment and clearly suggest that the use of an antidepressant in this patient group is highly indicated. However, available results do not allow for determining whether one specific compound is superior to others [66]. Therefore, in clinical practice, selection of the antidepressant drug should be guided by the patient’s comorbidities and the side effect profile of the specific drug. For example, in tinnitus patients with insomnia, the use of a sedating antidepressant such as amitriptyline might be preferable. Available studies and clinical experience suggest that the dose of antidepressants for the treatment of tinnitus is in a similar range as that used in the treatment of depression. In general, a low starting dose and slow increase of the dosage reduce side effects. Since beneficial effects do not occur immediately, minimum treatment duration of 6–12 weeks at the effective dose is recommended. If treatment effects are unsatisfactory and the decision is made to discontinue or change treatment, dosage should be reduced slowly. If a patient experiences beneficial effects, treatment should be continued at a stable effective dose for about 6 months, then the dose can be reduced over the course of weeks to months. Should the tinnitus get worse during a reduction of the dose, it is recommended to keep the dosage at the minimum providing relief.

**Benzodiazepines**

Severe tinnitus can be an extremely stressful condition, heavily influencing every aspect of the patient’s life. Since benzodiazepines are allosteric potentiators of the GABA<sub>A</sub> receptor [68] and tinnitus is thought to be the result of an imbalance between excitatory and inhibitory neurotransmission toward the former [12], benzodiazepines should have a positive effect on tinnitus by increasing inhibitory neurotransmission. Furthermore, due to their anxiolytic and sleep-inducing properties, benzodiazepines should have beneficial effects on comorbid anxiety and insomnia, and thus may help patients cope with their tinnitus.

In a prospective double-blind placebo-controlled study, 12 weeks of alprazolam administration at an individually adjusted dosage reduced tinnitus loudness in 76% of subjects – measured with a tinnitus synthesizer and a visual analog scale – whereas only 5% showed a reduction in tinnitus loudness in the control group [69]. Although the strong positive effects of alprazolam are encouraging, the study has criticized because of the small sample size, drug dosing method, and failure to assess emotional effect [7]. On the other hand, diazepam, evaluated in a double-blind triple cross-over trial involving 21 tinnitus patients, had no effect on tinnitus loudness [70]. In a retrospective study of medical records from over 3,000 patients taking clonazepam (0.5–1 mg/day, 60–180 days) for vestibular or cochleovestibular disorders, 32% reported an improvement in their tinnitus [71]. However, the lack of a control group makes it difficult to evaluate the significance of these findings. In a prospective, randomized, single-blind clinical trial involving ten patients per group, clonazepam significantly reduced tinnitus loudness and annoyance (visual analog scale) relative to the control group [72]. Summarizing, available results seem to suggest a beneficial effect of benzodiazepines on tinnitus. However, additional studies are needed in order to evaluate the efficacy of benzodiazepines on tinnitus.

Due to their immediate effects, short-acting benzodiazepines such as lorazepam or alprazolam are widely used for acute treatment of anxiety, agitation, and insomnia – symptoms that frequently occur with tinnitus. The longer acting clonazepam provides some relief in a considerable subgroup of patients. The use of benzodiazepines should be restricted to short periods of time due to the risk of drug dependency. Moreover,
caution is warranted since protracted tinnitus has been reported after discontinuation of benzodiazepines [73, 74].

**Non-Benzodiazepine Anticonvulsants**

Anticonvulsants are increasingly used in the treatment of several non-epileptic conditions, including various psychiatric disorders and pain syndromes [75]. Some of them have also been investigated for the treatment of tinnitus Table 78.2. Diverse pharmacological mechanisms of action are responsible for the therapeutic effects of antiepileptics; among them effects on voltage-gated sodium and calcium channels, and on synaptic transmission – mainly mediated by gamma amino butyric acid type A (GABA_A) receptors [76]. Since antiepileptics reduce neuronal excitability, in principle, they should be beneficial for the treatment of tinnitus.

The anticonvulsant carbamazepine, which binds to voltage-gated sodium channels and stabilizes the sodium inactivation state, thereby reducing neural firing [77, 78], has been investigated for tinnitus with mixed results. Based on the assumption that carbamazepine resembles lidocaine in its mechanism of action, three studies investigated the effect of carbamazepine in tinnitus patients who previously had responded to intravenous lidocaine [79–81]. About half of these patients had a positive response to carbamazepine (600–1,000 mg daily). However, controlled studies have not demonstrated benefits of the drug compared to placebo [82–84]. A significant benefit from carbamazepine has been reported for a rare group of patients who have intermittent tinnitus that sounds like a typewriter, popcorn, or ear clicking, and which is caused by a neurovascular conflict [85, 86].

The anticonvulsant gabapentin acts on voltage-gated calcium channels and is also used for the treatment of seizures, neuropathic pain, and migraine [87–89]. The results with gabapentin for the treatment of tinnitus are contradictory. One controlled trial has shown a significant improvement in tinnitus annoyance and loudness for a subgroup of participants with tinnitus related to acoustic trauma [90]. A second study did not detect any improvement in tinnitus handicap, but did report a significant improvement in tinnitus annoyance when compared to placebo [91]. However, further controlled trials did not report any benefit of the compound on tinnitus annoyance or loudness [92, 93].

Thus, although the effects of gabapentin are limited, it might benefit a subpopulation of patients in which tinnitus is associated with acoustic trauma [94].

Pregabalin, which resembles gabapentin in its mechanisms of action, is indicated not only for the treatment of partial seizures but also for neuropathic pain, fibromyalgia, and anxiety [95–97]. Beneficial effects on sleep have also been reported [98]. There are no data available for its use in tinnitus, but based on available data and clinical experience, pregabalin seems to be a promising option for the treatment of tinnitus-related anxiety and insomnia.

Lamotrigine, which stabilizes neuronal membranes by inhibiting voltage-sensitive sodium channels, has been investigated in a double-blind placebo-controlled cross-over clinical trial on 33 patients where it failed to demonstrate a beneficial effect [99]. Valproic acid, which is one of the most frequently prescribed antiepileptic drugs and which acts by multiple mechanisms has not been systematically investigated and only been reported in case reports as useful in tinnitus [100, 101].

**Antiglutamatergic Compounds**

Glutamate receptor antagonists have been tried in tinnitus sufferers. The rationale behind it is that imbalance between inhibitory vs. excitatory neurotransmission is observed in several regions of the auditory pathway in tinnitus [12, 13]. Moreover, blocking glutamatergic neurotransmission could also exert neuroprotective effects, as it is known that noise overexposure is followed by an excitotoxic injury of the hair cells [102]. The putative non-selective NMDA receptor antagonist acomprosate has been tried in a double-blind study [103]. Patients received placebo or acamprosate (333 mg, three times per day) and rated the loudness and annoyance of their tinnitus before and at monthly intervals of treatment. Acomprosate did not show any significant effects after 30 days of treatment, but a modest benefit at 60 days, and a significant effect at 90 days. Approximately 87% of the subjects in the acamprosate group showed some improvement, including three subjects in which tinnitus disappeared, compared to 44% in the placebo group. A larger clinical trial is currently underway to analyze the encouraging results from this preliminary study (http://clinicaltrials.gov/ ct2/show/NCT00596531). Treatment with i.v. caroverine, an antagonist of non-NMDA and NMDA receptors,
has been analyzed with contradictory results [104, 105]. In a prospective randomized double-blind crossover study using the tinnitus handicap inventory to assess efficacy, 90-day treatment with the non-selective NMDA antagonist memantine was no more effective than placebo [106]. The memantine analogue nera-"x80, which blocks both NMDA [107] and a9a10 nicotinic cholinergic receptors [108] is at phase III of a clinical trial setting (http://clinicaltrials.gov/ct2/show/ NCT00405886).

### Dopaminergic–Antidopaminergic Drugs

Both dopaminergic and antidopaminergic drugs have been proposed for treating tinnitus. Dopaminergic pathways in limbic and prefrontal areas may be involved in mediating emotional aspects of tinnitus [109, 110]. In one double-blind placebo-controlled study, sulpiride significantly reduced subjective ratings of tinnitus and tinnitus visual analogue scores. Effects were more pronounced when sulpiride was combined with either hydroxyzine (an antihistamine and anxiolytic) or melatonin [111, 112]. The dopamine agonist piribedil was investigated recently in a double-blind placebo-controlled cross-over study. In this study, piribedil was not superior to placebo; however, a post hoc analysis suggested that a subgroup of patients with specific findings in the electrocochleography may benefit from piribedil [113]. Although these results are preliminary and need further studies, they are encouraging and indicate that the dopaminergic pathway might be a promising target for tinnitus relief. A clinical trial is under way to assess the efficacy of flupentixol (a type of thioxanthene drug that acts by antagonism of D1 and D2 dopamine, and serotonin type 2A receptors [114]) plus clonazepam, compared to clonazepam alone (http://clinicaltrials.gov/ct2/show/NCT00841230).

### Other

Some other miscellaneous drugs have been tested with limited efficacy or that require further controlled trials Table 78.2. These include the HMG-CoA reductase atorvastatin, the vasodilator cyclandelate, the loop diuretic furosemide, some herbal products like *Ginkgo biloba*, melatonin, the prostagaldin E1 analogue misoprostol, the L-type calcium blocker nimodipine, the phosphodiesterase type 5 inhibitor vardenafil, and minerals including zinc.

Atorvastatin reduces the synthesis of cholesterol by inhibiting HMG-CoA reductase [115]. In a randomized double-blind placebo-controlled study over 13 months involving elderly patients with elevated cholesterol, atorvastatin failed to slow the progression of age-related hearing loss and significantly reduce tinnitus [116].

Cyclandelate, a vasodilator used in the treatment of cerebrovascular and peripheral vascular disorders, that is believed to act by blocking calcium influx [117], has been investigated for the treatment of tinnitus based on the assumption that some forms of tinnitus may arise from cerebrovascular insufficiency. In an open multicentric clinical trial of patients with tinnitus, vertigo, and visual disturbances, 90-day treatment with cyclandelate reduced the severity and frequency of these symptoms with minimal side effects [118]. However, in a subsequent placebo-controlled double-blind study, cyclandelate did not significantly change audiometric measures of tinnitus loudness and pitch and caused side effects in many patients [119].

Furosemide is a loop inhibiting diuretic used to treat congestive heart failure and edema [120]. Furosemide has been proposed as a treatment for tinnitus of “cochlear” origin because it strongly suppresses the endolymphatic potential and other cochlear responses [121]. One initial study has shown that approximately 50% of patients exhibited a reduction of tinnitus symptoms following intravenous furosemide treatment [122]. Moreover, furosemide has also been found to suppress tinnitus in approximately 40% of patients with Ménière’s disease [123]. However, high doses of furosemide can also induce temporary hearing loss and tinnitus [124].

*Ginkgo biloba* has been proposed for the treatment of a wide range of disorders including tinnitus [125]. In western countries, *Ginkgo biloba* is commonly available in form of leaf extracts, which in Europe and in the United States are among the most widely used and appreciated herbal medications. *Ginkgo* extract contains two main pharmacologically active substances such as flavonoid glycosides and terpene lactones, responsible for many biological effects. Even if some studies have suggested beneficial effects of *Ginkgo* on
tinnitus, particularly in patients with short duration symptoms [126, 127], there is a growing body of evidence from large, well-controlled double-blind placebo-controlled clinical studies clearly indicating that Ginkgo is no more effective in alleviating tinnitus symptoms than placebo [128, 129]. EGb-761 is a concentrated extract of Ginkgo biloba (enriched in flavonoids and terpenes) which has a broad spectrum of pharmacologic actions, including a free-radical scavenger effect, and which has shown efficacy for tinnitus in a phase I trial. Several other herbs have been proposed for tinnitus therapy, such as Cimicifuga racemosa, Cornus officinalis, Verbascum densiflorum, and Yoku-kan-san, but none of them have been tested in well-controlled trials [130].

Melatonin is a neurohormone that is primarily produced by the pineal gland. Since it can influence sleep and circadian rhythms, melatonin is nowadays widely used for treating sleep disturbances [131]. This effect of melatonin may have been the rationale for using this drug in the treatment of tinnitus. An open label study found statistically significant improvements on ratings of tinnitus severity and sleep quality scores [132], whereas a double-blind placebo-controlled cross-over study did not demonstrate superiority of melatonin over placebo [133]. A more recent randomized double-blind placebo-controlled study found that melatonin in combination with sulpiride reduced subjective rating of tinnitus and tinnitus loudness more than placebo [112].

Misoprostol is a synthetic prostaglandin E1 analogue which is primarily used to prevent gastric ulcers induced by non-steroidal anti-inflammatory drugs [134]. In a small, placebo-controlled cross-over study, tinnitus severity improved in 33% of subjects during misoprostol treatment (escalating to 800 mg/day), while none improved with placebo [135]. A subsequent double-blind placebo-controlled study has shown a significant reduction of tinnitus loudness with misoprostol treatment, but no differences in subjective measures of tinnitus severity [136]. A further study has shown efficacy of misoprostol in the treatment for chronic tinnitus in hypertensive and/or diabetic patients [137].

Nimodipine is a calcium antagonist, which crosses the blood–brain barrier and blocks L-type calcium channels [138]. Although a first open clinical trial suggested positive effects of nimodipine on tinnitus in some patients [139], these could not be confirmed in a second open clinical trial [140].

Vardenafil represents a potent and highly selective phosphodiesterase type 5 (PDE5) inhibitor that induces an increase of nitric oxide-mediated vasodilatation and which is marketed for treatment of erectile dysfunction and pulmonary hypertension. A prospective randomized double-blind placebo-controlled trial did not show any benefit of vardenafil over placebo [141].

Zinc is an essential catalytic or structural element of many proteins and a signaling messenger that is released by neural activity at many central excitatory synapses. Growing evidence suggests that zinc may also be a key mediator and modulator of the neuronal death associated with transient global ischemia and sustained seizures, as well as perhaps other neurological disease states [142]. While positive results have been reported in some tinnitus patients with hypozincemia, zinc therapy did not result in tinnitus improvement in patients with normal zinc levels in several double-blind placebo-controlled studies [143–148].

Conclusions

Despite the significant unmet clinical need for a safe and effective drug targeting tinnitus relief, there is currently not a single FDA-approved drug on the market. Although the available treatments for the management of the tinnitus patient are diverse, most patients and clinicians are waiting for a drug than can suppress or significantly reduce tinnitus. Thus, there is a pressing need to develop a drug targeting tinnitus relief. A wide variety of drugs with different therapeutic uses have been used off-label with some effect in a limited subset of patients. Tinnitus-related comorbidities such as depression or anxiety can especially be addressed successfully with pharmacological treatment. Since pharmaceutical companies are slowly entering the tinnitus field, this scenario most likely will change in the near future.

References


Chapter 79
The Endocannabinoid System in the Cochlear Nucleus and Its Implications for Tinnitus Treatment

Paul F. Smith

Keypoints

1. One of the main theories of tinnitus is that it is a form of sensory epilepsy, sometimes arising from neuronal hyperactivity in the brainstem cochlear nucleus.
2. Antiepileptic drugs have therefore been explored as one potential treatment option.
3. Increasing evidence suggests that cannabinoid drugs can also have antiepileptic effects.
4. Recently, it has been reported that cannabinoid CB1 and CB2 receptors and the endogenous cannabino
d 2-arachidonoylglycerol (2-AG), are expressed in the cochlear nucleus.
5. CB1 receptors appear to negatively regulate the release of glutamate, and it is possible that their down-regulation during the development of tinnitus is responsible for the neuronal hyperactivity associated with the condition.
6. This chapter explores the possibility that cannabinoid drugs might be useful in the treatment of tinnitus.

Keywords  Tinnitus • Cochlear nucleus • Cannabinoid receptors • Endocannabinoids

Abbreviations

2-AG  2-Arachidonoylglycerol
ACPA  CB1 receptor agonist
CB  Cannabinoid
CN  Cochlear nucleus
Δ9-THC  Δ9-tetrahydrocannabinol
DAG  Diacylglycerol lipase
DCN  Dorsal cochlear nucleus
FAAH  Fatty acid amide hydrolase
GABA  Gamma-amino butyric acid
IPSC  Inhibitory post-synaptic currents
LTD  Long-term depression
LTP  Long-term potentiation
MAGL  Monoacylglycerol lipase
NAPE-PLD  N-arachidonoylphosphatidylethanolamine-phospholipase-D
PEA  N-palmitoylethanolamide
PSP  Post-synaptic potentials
SCE  Standard cannabis extracts
VCN  Ventral cochlear nucleus

Introduction

Subjective tinnitus is often caused by exposure to loud noise, and sometimes by head/neck injuries or exposure to ototoxic drugs (e.g., salicylate) [1]. Many theorists believe that the mechanisms for initiating tinnitus (‘ignition’ mechanisms) may be somewhat separate from those that maintain it [2, 3]. While cochlear hair cell dysfunction may trigger noise-induced tinnitus, there is evidence that the maintenance of tinnitus is associated with neuronal hyperactivity in the central auditory nervous system. Acoustic trauma has been correlated with increased spontaneous activity in the dorsal cochlear nucleus, (e.g., [4–9], but see [10, 11] for contradictory data), the inferior colliculus [12–14], and the primary (but not the secondary) auditory cortex [15, 16]. On the basis of such studies, it has been proposed that tinnitus is a form of sensory epilepsy...
that might therefore be responsive to antiepileptic drugs [17, 18].

However, the evidence supporting the efficacy of antiepileptic drugs in the treatment of tinnitus is inconsistent [see 19 for a review]. Gananca et al. [20] performed a retrospective survey of 25 years of the use of clonazepam and concluded that it was at least partially effective in 32% of cases of tinnitus. Shulman et al. [21] suggested that for tinnitus of central origin, benzodiazepines provided long-term relief in 90% of cases. However, there are no systematic well-controlled clinical trials (i.e., double-blind, placebo-controlled) of the effects of benzodiazepines on tinnitus. Menkes and Larson [22] published a single case study reporting that sodium valproate was effective in suppressing tinnitus; however, there has never been a properly controlled clinical trial conducted to evaluate its efficacy. Carbamazepine has also been used, but other than case studies, e.g., [22, 23], only 3 clinical trials have been published. Melding and Goodey [24] reported that 56% of patients who had responded positively to lidocaine experienced relief from tinnitus following carbamazepine treatment. Sanchez et al. [25] also reported that carbamazepine was effective in reducing tinnitus in 58% of patients and abolished tinnitus in 18% of patients. However, Hulshof and Vermeij [26] reported that carbamazepine was less effective than a placebo in relieving tinnitus.

By far, the best studied antiepileptic drug in the context of tinnitus is gabapentin [27]. Following a positive case study [28], Bauer and Brozoski [29] conducted a prospective, placebo-controlled, single-blind trial of the effects of gabapentin on 39 patients with tinnitus. They found that the drug was effective in reducing tinnitus in some patients, especially those in whom the condition was related to acoustic trauma. However, Witsell et al. [30], in a more recent study using a randomized, placebo-controlled, double-blind trial reported that gabapentin had no significant effect on the severity of tinnitus. Piccirillo et al. [31] reported similar results from an 8-week double-blind, randomized trial.

Cannabinoids as Antiepileptic Drugs

In the late 1980s, Δ⁹-tetrahydrocannabinol (Δ⁹-THC), the principal psychoactive constituent of the Cannabis sativa plant, was demonstrated to act on a specific G-protein–coupled cannabinoid receptor (the “CB1 receptor”) [see 32, 33 for reviews]. By 1993, an endogenous cannabinoid named ‘anandamide’ (arachidonylethanolamide) had been discovered in the porcine brain. A second cannabinoid receptor subtype, the “CB2 receptor”, was identified in the peripheral nervous system and immune system [see 32, 33 for reviews]. During the 1990s, these findings were replicated and extended, and it became clear that the endogenous cannabinoid (“endocannabinoid”) signaling system was central to many aspects of brain function. A second endocannabinoid, 2-arachidonoylglycerol (2-AG), was discovered. It was shown that these arachidonic acid derivatives were synthesized by enzymes such as N-arachidonoylphosphatidylethanolamine-phospholipase-D (NAPE-PLD for anandamide) and diacylglycerol lipase (DAG for 2-AG) and metabolized by enzymes such as fatty acid amide hydrolase (FAAH for anandamide and 2-AG) and monoacylglycerol lipase (MAGL for 2-AG) [see 32, 34 for reviews]. As a consequence of these revolutionary developments in cannabinoid pharmacology, there has been intense interest in the development of both synthetic and natural cannabinoid compounds for the treatment of a wide range of disorders, including spasticity, pain, urinary dysfunction, and epilepsy [35].

Cannabinoids have been reported to have pro- or anticonvulsant effects under different circumstances [36, 37]. Epidemiological studies suggest that cannabis’ use is common amongst people with epilepsy and that the drug is believed by users to have anticonvulsant actions [38]. Gordon and Devinsky [36] reviewed the literature on the effects of marijuana on epileptic symptoms and concluded that although cannabis’ use can reduce seizure frequency in many cases and provoke seizure activity in some cases, it probably has no effect in most cases. Unfortunately, there have been a very few clinical studies of the effects of cannabinoids on seizure activity in humans, and no large, well-controlled, double-blind studies [36, 37]. Nonetheless, some recent reports of the anti-epileptic effects of the synthetic Δ⁹-THC, dronabinol, have been published, e.g., [39].

CB1 receptors are thought to be localized mainly presynaptically and in many cases, through the inhibition of calcium influx at presynaptic terminals, inhibit the release of classical neurotransmitters, including glutamate [40, 41]. Wallace et al. [42] used the rat pilocarpine model of epilepsy to investigate the effects of cannabinoids on seizure activity. Δ⁹-THC, as well as the synthetic cannabinoid receptor agonist R(+)

9-tetrahydrocannabinol (∆9-THC), as well as

D9-THC,

Cannabis

9-arachidonoylphosph

N-arachidonoylphosphatidylethanolamine-phospholipase-D (NAPE-PLD for anandamide) and diacylglycerol lipase (DAG for 2-AG) and metabolized by enzymes such as fatty acid amide hydrolase (FAAH for anandamide and 2-AG) and monoacylglycerol lipase (MAGL for 2-AG) [see 32, 34 for reviews]. As a consequence of these revolutionary developments in cannabinoid pharmacology, there has been intense interest in the development of both synthetic and natural cannabinoid compounds for the treatment of a wide range of disorders, including spasticity, pain, urinary dysfunction, and epilepsy [35].

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WIN55,212, completely blocked spontaneous seizure activity. The CB1 receptor antagonist SR141716A potentiated seizure duration and frequency, suggesting that endocannabinoids might have been suppressing seizure activity. Biochemical analyses of the hippocampus indicated that 2-AG was present in elevated concentrations during the seizure activity, which the authors hypothesized might play a role in termination of the seizures. Immunohistochemical and western blot analyses also showed that CB1 receptor expression was increased throughout the hippocampus during the seizure activity, again suggesting the possibility that the endocannabinoid system might serve some form of anti-epileptic function. Using the pentylentetrazole model of epilepsy in mice, Shafaroodi et al. [43] demonstrated that the CB1 receptor agonist ACPA increased the seizure threshold, whereas the CB1 receptor antagonist AM251 blocked this anticonvulsant effect.

Not all studies have demonstrated an anti-epileptic action for CB1 receptor agonists, suggesting that where CB1 receptors are localized to GABAergic terminals, cannabinoids could potentiate epileptiform activity [44]. Nakatsuka et al. [45] used patch clamp recordings in granule cells of the human dentate gyrus to show that activation of CB1 receptors could suppress inhibitory synaptic activity. Bath application of WIN55212-2 suppressed the frequency of spontaneous inhibitory postsynaptic currents (IPSCs) as well as reducing their amplitude. The CB1 receptor antagonist AM251 completely blocked these effects, suggesting that they were mediated by CB1 receptors. It is likely that the activation of CB1 receptors on presynaptic GABAergic terminals results in decreased GABA release, which was responsible for the reduction in IPSC frequency and amplitude. These and similar previous results demonstrate that CB1 receptor agonists have the potential to induce as well as decrease epileptiform activity, depending on which presynaptic terminals the drugs affect.

Wallalley et al. [46] have provided evidence that emphasizes the potential complexity of the effects of cannabis itself on epileptiform activity. Comparing standard cannabis extracts (SCEs) with and without Δ⁹-THC, they found that while Δ⁹-THC depressed depolarizing post-synaptic potentials (PSPs) in rat olfactory cortex neurons, SCEs with and without Δ⁹-THC could potentiate PSPs. This effect could be blocked by the CB1 receptor antagonist SR141716A. One of the most surprising findings from this study was that the potentiation of PSPs was actually greater when a Δ⁹-THC–free SCE was used. The authors suggested that a novel, unknown component in cannabis may override the decrease in excitatory synaptic transmission caused by Δ⁹-THC and that this constituent may be partly responsible for the pro-convulsant effects of cannabis that have sometimes been reported.

In a similar study, Wilkinson et al. [47], using seizure activity induced in rat piriform cortical brain slices by oxotremorine-M, found that an SCE had a more potent anticonvulsant action than Δ⁹-THC alone, but that the Δ⁹-THC–free extract also had anticonvulsant activity.

Marsicano et al. [48] bred mutant mice lacking CB1 receptors on principal forebrain neurons, but not on inhibitory interneurons, and found that kainic acid caused extreme seizure activity in vivo compared to wild-type littermates. In vitro, the threshold for neuronal excitation caused by kainic acid was reduced in the hippocampi of CB1 receptor–deficient mice. In wild-type mice, kainic acid administration resulted in increased concentrations of anandamide in the hippocampus. However, 2-AG or palmitoylethanolamide (PEA) levels were not affected and principal hippocampal neurons appeared to be protected. The anandamide uptake inhibitor UCM707 was also shown to protect against kainic acid–induced seizures. However, no protective effects occurred in the mutant mice, suggesting that CB1 receptors are necessary for the brain to protect against kainic acid–induced seizures. Mechoulam and Lichtman [49] have suggested that the endocannabinoid system may serve as a natural, endogenous anticonvulsant network. The fact that endocannabinoids such as anandamide and 2-AG are synthesized on demand and are rapidly metabolized adds validity to the concept that they may function as an “on-demand” defense system [37]. Recently, Ludanyi et al. [50] have reported that in hippocampi from humans with chronic epilepsy, CB1 receptor gene and protein expression, as well as DAG expression, were down-regulated in a way that correlated with the degree of sclerosis.

Wallace et al. [51] have also demonstrated that anandamide and its analogue, O-1812, have potent anticonvulsant effects in the maximal electroshock seizure model in mice. These effects could be blocked by the CB1 receptor antagonist SR141716A. Similar to the results of Wallace et al. [42], SR141716A significantly reduced the seizure threshold, consistent with the idea that activation of cannabinoid receptors by naturally occurring anandamide serves an anticonvulsant role. Anandamide and the
synthetic CB1 receptor agonist WIN-55212-2 (but not the inactive isomer WIN 55212-3) have both been shown to reduce epileptiform activity in hippocampal slices. This effect can be blocked by SR141716A [52]. It is likely that these effects were mediated by anandamide and WIN-55212-2 acting on cannabinoid receptors on presynaptic glutamatergic terminals.

N-palmitoylethanolamide (PEA) is another member of the family of endogenous lipid amides and is a putative endocannabinoid. PEA has been reported to demonstrate anticonvulsant activity in the mouse maximal electroshock and chemically induced convulsion models (e.g., pentylentetrazol, bicuculline, strychnine) [53]. In the maximal electroshock model, the anticonvulsant effects of PEA were comparable to phenytoin [53]. PEA has also been shown to have an anticonvulsant action in the kindling model of epilepsy in rats [54].

All together, this evidence suggests that the endocannabinoid system serves a critical function in the control of hyperexcitability and that endocannabinoids and cannabinoid drugs may have antiepileptic effects that might be useful in the treatment of tinnitus.

Endocannabinoids and Cannabinoid Receptors in the Cochlear Nucleus

Until recently, there have been a few studies of cannabinoid receptors in the cochlear nucleus (CN). CB1 receptors were identified in the CN in early autoradiographic studies; however, Herkenham et al. [55] concluded that the CN had the lowest density of CB1 receptors of any brain region. This finding may have discouraged researchers from investigating CB1 receptors in the CN. However, the density of receptors is not the only indication of their likely significance. Receptor affinity and efficacy (i.e., the intracellular effect of receptor activation) are also important. In fact, Brievogel et al. [56] have reported that CB1 receptors in many brainstem regions have greater coupling to their G proteins (i.e., greater efficacy) than those in limbic and neocortical areas.

The first studies of CB1 receptors in the CN were published by Zheng et al. [57] and Tzounopoulos et al. [58]. Zheng et al. [57] used immunohistochemistry and stereological methods to quantify CB1 receptor expression in the dorsal and ventral cochlear nuclei (DCN and VCN, respectively). They found substantial CB1 receptor labeling on many different cell types, such as stellate cells, giant cells, fusiform cells, and corn cells in the DCN, as well as globular bushy cells, elongate cells, and octopus cells in the VNC (Figs. 79.1 and 79.2). Some of the labeling was cytoplasmic, which seemed inconsistent with the accepted presynaptic localization of CB1 receptors; however, it has been reported that the CB1 receptor undergoes extensive trafficking between the cytoplasm and the presynaptic terminals, especially in brain regions where it is very active. An earlier western blot study by Ashton et al. [59] had reported CB1 receptor levels in the CN that were similar to the cerebellar granule cell layer and cerebellar nuclei.

These results were confirmed and extended by Tzounopoulos et al. [58], who found CB1 receptors at
In a further study, Zhao et al. [60] showed that glutamate terminals in the DCN expressed more CB1 receptors than glycergic terminals, and that both fusiform and cartwheel cells expressed DAG α and β, the two enzymes necessary for the production of 2-AG. Both forms of DAG were found in the dendritic spines of cartwheel cells but not fusiform cells, suggesting that the production of 2-AG is closer to parallel fiber synapses in cartwheel cells compared to fusiform cells. This was

![Fig. 79.2](image) High magnification view of CB1 receptor immunoreactivity in different cell types of the cochlear nucleus. (a): Granule cells in the molecular layer of the dorsal cochlear nucleus. (b): A stellate cell. (c): A cartwheel cell. (d): A giant cell. (e): A fusiform cell. (f): A corn cell. (g): A globular bushy cell. (h): An elongate cell. (i): An octopus cell. Reproduced from [57] with permission.
the first evidence for a complete endocannabinoid system in the DCN, involving, at the very least, 2-AG acting on CB1 receptors. Zhao et al. [60] concluded that the endocannabinoid system exerts a greater control over excitatory than inhibitory inputs in the DCN and that endocannabinoid signaling is a major factor affecting the balance of excitation and inhibition in this part of the central auditory system.

The expression of the second subtype of cannabinoid receptor in the brain, the CB2 receptor, is controversial. However, Baek et al. [61] found CB2 receptor labeling in the CN, which suggests the possibility that both cannabinoid receptor subtypes could be involved regulating the function of the CN (Fig. 79.3).

Cannabinoids, Cannabinoid Receptors, and Tinnitus

Only one study to date has investigated the relationship between CB1 receptors in the CN and the development of tinnitus. Zheng et al. [57] investigated the expression of CB1 receptors in the DCN and VCN in rats in which tinnitus was induced with salicylate injections. They used a modification of the conditioned behavioral paradigm developed by Jastreboff et al. [62] in order to confirm that the animals were experiencing tinnitus (Fig. 79.4). In animals with tinnitus, they found a significant decrease in the number of neurons expressing CB1 receptors in the VCN compared to control animals. On the other hand, there was no significant difference in the DCN (Fig. 79.5). Zhao et al. [60] have suggested that if CB1 receptors were down-regulated on glutamatergic terminals synapsing on fusiform cells, this would increase the excitation of fusiform cells, possibly leading to hyperexcitability.
Unfortunately, there have been no systematic studies to date of the effects of cannabinoids on tinnitus, in either animals or humans. One case report has been published in which tinnitus was eliminated by administration of the synthetic Δ⁹-THC, dronabinol [63]. However, the patient had intracranial hypertension with many other symptoms and had been previously using cannabis.

Given that agonists for the CB1 receptor have been shown to exert antiepileptic effects, that antiepileptic drugs do appear to alleviate tinnitus at least in some circumstances and that the endocannabinoid system is emerging as an important influence in the function of the CN; it seems worthwhile to pursue the possibility that cannabinoid drugs could be useful in the treatment of tinnitus. Since some of these drugs, including natural extracts (e.g., Sativex®), are already available in many countries for the treatment of nausea, wasting, chronic pain, and spasticity [see 64 for a review], it would not be difficult to test their effects in patients with tinnitus. Despite concerns about intoxication, most studies suggest that provided the levels of Δ⁹-THC are low, few adverse side effects are experienced [65, 66].

Conclusions

This chapter has explored the possible significance of the endocannabinoid system for the cochlear nucleus and the treatment of tinnitus. There is substantial evidence to suggest that tinnitus is, in many cases at least, a form of sensory epilepsy [18]. Although the results of clinical trials of the effects of antiepileptic drugs on tinnitus are not consistent, these drugs have been shown to alleviate tinnitus in some cases. Therefore, since cannabinoids have been shown to exert antiepileptic effects in many parts of the brain, it is possible that they will exert similar effects in the central auditory system. In this respect, it is important to note that both an endocannabinoid (2-AG) and two cannabinoid receptor subtypes (CB1 and CB2) have been found in the CN [57, 58, 60, 61]; the CB1 receptors are functional, are preferentially localized to glutamatergic terminals, and indeed mediate synaptic plasticity in the CN [58, 61]; and CB1 receptors have been shown to down-regulate in the VCN in an animal model of tinnitus [57]. The functional significance of the endocannabinoid system for tinnitus, therefore, deserves urgent investigation.

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References

3. Baguley, DM. What progress have we made with tinnitus? Acta Otolaryngol (Stock) 2006, 126:4–8
44. Hoffman, AF and Lupica, CR. Mechanisms of cannabinoid inhibition of GABA(A) synaptic transmission in the hippocampus. J Neurosci 2000, 20:2470–2479
Chapter 80
Treatment of Somatosensory Tinnitus

Tanit Ganz Sanchez and Carina Bezerra Rocha

Keypoints

1. Treatment of somatosensory tinnitus often needs a multidisciplinary approach.
2. Treatment of patients who have signs of bone problems, muscular tension in the temporomandibular joint area or neck, should be directed to correct these problems as the first option.
3. If correction of bone or muscular disorders of temporomandibular joint and neck fails in relieving tinnitus, “symptomatic” treatment should be initiated.

Keywords Tinnitus • Somatic • Myofascial trigger point • Cervical spine • Temporomandibular joint • Treatment

Abbreviations

Bot-A Botulinum toxin type A
CNS Central nervous system
DCN Dorsal cochlear nuclei
GET Gaze-evoked tinnitus
MPS Myofascial pain syndrome
MTP Myofascial trigger point
TENS Transdermelectrical nerve stimulation
THI Tinnitus handicap inventory
TMD Temporomandibular disorder
TMJ Temporomandibular joint
VAS Visual analogue scale

Introduction

As discussed in Chap. 43, tinnitus may have a somatosensory cause (primary origin in head and neck trauma, dental or cervical manipulation, or even in unknown chronic pain) or somatosensory modulation (auditory origin with temporary somatosensory influence in loudness, pitch or localization). Thus, it is logical to consider that these patients need evaluation of their temporomandibular joint (TMJ) and neck before deciding the best treatment option. However, otolaryngologists and neurologists are usually the first physicians to be sought by patients with tinnitus, and unfortunately little attention is often given to somatosensory influence on tinnitus [1]. Many patients with tinnitus would benefit from the opinion of a dentist or physiotherapist (according to customs and availability in each country). Whenever it is agreed that a patient with tinnitus has problems such as muscular tension in TMJ related to the TMJ or the neck, these problems should be addressed (see also Chaps. 95 and 96).

In this chapter, we will discuss different approaches for treating somatosensory tinnitus including certain methods that are generally used for patients with tinnitus but have been modified for somatosensory tinnitus.

Treatment Options to Somatosensory Tinnitus

Treatments for somatosensory tinnitus are now under development by several researchers, including a group associated with the Tinnitus Research Initiative, working to define the best options for patients with somatosensory tinnitus. This chapter will describe results from several such studies, some of which are under development.
Relaxing Muscle Tension in Jaw and Neck

Considering that patients with temporomandibular disorders (TMD) often have muscular tension in both jaw and neck – as well as tinnitus, vertigo/dizziness, and aural fullness – the first aim of the treatment of somatosensory tinnitus should be to reduce such muscle tension [2].

Methods used for treatment of TMD by the dentist and the physiotherapist are described in Chaps. 95 and 96. Many patients with such problems also benefit from performing regular stretching exercises of their suboccipital muscles at home, as well as rotation movements in the atlanto-occipital joint – especially to the restricted side – and relaxing exercises involving breathing with the diaphragm.

Such treatment of muscle tension in the jaw and neck can reduce tinnitus, as well as decrease tension-related symptoms such as vertigo, aural fullness, and pain in the jaw, neck, or headache [2]. Björne showed that the intensity of all such symptoms was significantly reduced (\(p<0.001\)) at a 3-year follow-up examination for patients who used this type of treatment.

Focal administration of lidocaine in jaw muscles (mainly the lateral pterygoid or masseter) or neck muscles (mainly trapezius and sternocleidomastoid) may temporarily reduce tension [3], and sometimes tinnitus decreases when the local anesthetic is active [2]. This means that relieving TMJ disorders and other forms of muscular tension may also relieve tinnitus. Felício et al. [4] observed that patients with TMD and auditory symptoms improved after using bite splints for 8 weeks. Wright and Bifano [5] reported 82.5% improvement of tinnitus in patients whose TMD improved with the association of cognitive therapy, bite splints, and home exercises.

Deactivating Myofascial Trigger Points (MTP)

The relationship between myofascial trigger point (MTP) deactivation and tinnitus relief was initially demonstrated by injecting a local anesthetic in these painful spots. In 1960, Travell [6] related a patient’s tinnitus to a MTP located in the deep portion of masseter, because local injection of lidocaine on the ipsilateral masseter’s myalgic spots relieved the patient’s tinnitus. In this study, the injections were repeated eight times and the relief lasted from several days up to 4 weeks. Wyant [7] described two patients whose tinnitus was relieved after injections with steroid and lidocaine in MTPs in the cervical region. In this study, both tinnitus and pain relief occurred for 4 months.

In an unpublished study of 178 patients compared with a control group [3], tinnitus completely disappeared in 15% of the cases after cervical MTP injection. There were 178 people included in the treatment group. The control group consisted of 39 participants with tinnitus, who were not treated. In a 6-month follow-up after the last injection, tinnitus improved in more than 30%, as opposed to 15% of a control group.

Relief or reduction of tinnitus from injection of a local anesthetic (without corticosteroid or adrenaline) or dry needling depends on mechanical disruption of the myofascial trigger point and inactivation of the MTP. Inactivation of MTP can be done by injection of Botulinum toxin A because of its destruction of motor endplates. It is important for optimal effectiveness that the patients perform an active range of motion following the injection [8].

A 2005 study by Eriksson and co-workers [9] showed that some tinnitus patients benefited from stretching and massage. In a double-blind placebo-controlled randomized clinical trial, we have shown a significant decrease of tinnitus loudness (\(p<0.001\)), decrease of pain intensity (\(p<0.001\)), decrease of THI scores (\(p=0.01\)), and decrease of the number of MTP (\(p<0.001\)) from inactivation of MTPs.

The treatment was based on ten sessions of real and ten sessions of sham MTP deactivation in tinnitus patients. The real treatment used digital pressure in each MTP of eight possible muscles and some home orientations (muscle stretching, postural guidance, and hot pack). Seventeen patients from the experimental group and nine patients from the control group were analyzed by a blind researcher before treatment began and again after 10 weeks of the treatment. Moreover, two patients who had bilateral tinnitus now have complete relief, one who had bilateral tinnitus no longer has tinnitus in one ear, and another who had seven different kinds of tinnitus now only has three. Three patients with normal pure-tone audiograms, (i.e., tonal thresholds from 250 to 8,000 Hz better than 25 dBHL) were the ones with the best results after treatment, regardless of tinnitus being constant or intermittent, unilateral or bilateral.

Many techniques to relieve MTP have been published, but a few have been supported by scientific evidence.
The most commonly used treatment procedures for the myofascial pain syndrome (MPS) was reviewed by Vernon and co-workers [10] showing that these methods were supported by evidence from treatment. Laser therapy, transcutaneous electrical nerve stimulation, acupuncture, and magnet therapy for MTP and MPS are treatments, which are supported, in varying degrees, scientifically. The duration of relief fluctuates among therapies. Evidence is weak, however, for ultrasound therapy and is limited for electrical muscle stimulation, high-voltage galvanic stimulation, interferential current, and frequency-modulated neural stimulation.

We describe a patient below who was cured of her tinnitus through deactivation of MTPs in our institute:

Case report: A 51-year-old female complained of bilateral tinnitus for the last 4 years (worse on the right), scoring 10 on the visual analog scale. She also had pain in her shoulders, cervical spine, and the left side of her face for 4 years (score of 9) together with other complaints such as dizziness, daily bruxism, subjective hearing loss, and depression. A physical ENT exam showed no pathologies, and she was referred to a physiotherapist because of her pain. During palpation of MTP of the left sternocleidomastoid, she noticed a complete disappearance of her tinnitus. Tinnitus returned when pressure was no longer applied. Palpation of MTP of the right trapezius caused a change in pitch of her tinnitus. Palpation of her left trapezius muscle caused temporary dizziness. Nine muscles of her head, neck, and shoulder girdle had myofascial trigger points. She was treated for her MPS by digital deactivation (pressure release, Fig. 80.1) of MTPs, together with exercise at home including muscle stretching, self-massage in the MTP, isometric strengthening, and postural guidance. During her weekly sessions in the clinic, she showed gradual and lasting improvement of her pain, tinnitus, and dizziness. After 6 months, she had total remission of these problems.

**Manual Therapies: Cervical Manipulation**

Chiropractic care is a popular and successful management option for reversible functional disorders of the cervical spine and other body structures. Some studies have demonstrated that such manipulation can relieve tinnitus [11–15].

Thus, Alcantara and co-authors [13] described how chiropractic treatment of a patient with cervical subluxation and TMD could reduce the patient’s tinnitus, vertigo, and hearing loss. The symptoms eventually ceased after 9 sessions. Kessinger and Boneva [11] documented clinical changes after some chiropractic sessions in a geriatric patient with tinnitus, vertigo, hearing loss, and cervical alterations from C3 to C7. Throughout the sessions, the patient’s symptoms were alleviated, and structural/functional improvements were also evident through radiographic examination.

Contrary to classical chiropractic treatment, Arlen’s atlas therapy is performed without traction, rotation, or extension of the cervical spine. Kaute [16] considers that irritation and tension of posterior cervical muscle may precipitate a great afferent input to the vestibular nuclei in the brainstem, and this response seems to be one of the origins of idiopathic tinnitus. Diminishing the tension via atlas therapy seems to lower the proprioception and nociception output, leading to normalization of the flow of information to the brainstem and, as a consequence, the lessening of tinnitus.

It seems that some somatosensory tinnitus could be alleviated by correcting the misalignment of the cervical spine through manual therapy (chiropractic or osteopathy), especially in the upper cervical. This readjustment may allow the entire spine to reposition itself and possibly re-adjust the input of the region through the somatosensory pathway on the auditory system. Nevertheless, as much as this topic has been receiving more attention in the current literature, it still needs further clarification.

**Transelectrical nerve stimulation (TENS):** The use of electrical stimulation is used routinely in treatment of both pain and tinnitus. Its use in treatment of tinnitus is discussed in a separate chapter (Chap. 91).

**Botulinum Toxin Type A**

Botulinum toxin type A (Bot-A) is a neurotoxin that can inhibit the release of acetylcholine at the neuro-
muscular junction [17]. Due to its known paralytic effect, it is administered locally to control muscle hyperactivity in many different disorders, as well as in cosmetics.

Besides the paralytic effect, Bot-A might have a direct antinociceptive action through the blockage of the autonomic nervous system, in addition to the neuromuscular action [18–22]. Moreover, through a peripheral mechanism, it can inhibit the sensitization of central trigemino-vascular neurons [23]. Thus, Bot-A has more than one effect to control headache, migraine, and chronic neuropathic pain. Recently, Bot-A has been used for treatment of tinnitus [24, 25].

A crossover double-blind study of the effect of Bot-A on tinnitus showed that it had little effect on tinnitus [26]. One group was first injected with Bot-A and, 4 months later, with a saline injection into three sites around the ear: 1 cm above pinna, 1 cm behind – at 2 o’clock position – and 1 cm behind auricle – 5 o’clock position. The second group was first injected with placebo and, 4 months later, with BoNT-A injection. Seven participants reported a decrease in their tinnitus after the Bot-A injection and two improved after placebo.

When tinnitus was analyzed through global clinical impression by the patient (“better”, “worse,” or “the same”) and through THI, the effects of Bot-A were significantly better than placebo (p < 0.05), when comparing pre-treatment and 4 months after injection (p = 0.04). Such results suggest that the Bot-A can play a role in tinnitus management by reducing the peripheral inputs from cervical, temporal, frontal, and periauricular muscles.

A similar study [24] showed that of 26 participants, 7 improved, 13 worsened, and 16 were unchanged. It should be considered when discussing treatment with botulinum toxin that it has potentially serious side effects, such as changing cardiac reflexes [26]. Eighty percent of people treated with botulinum toxin had abnormalities in their electrocardiogram. The fact that the treatment probably has to be repeated for long periods is a disadvantage that when used for treating other disorders such as hemifacial spasm has made many people discontinue treatment [27].

**Training Exercises Repeating the Movements That Evoke Tinnitus Modulation**

Exercise in general may be beneficial because it increases brain-derived neurotrophic factor (BDNF) [28]. Training by repetitive movements generates specific neurophysiological changes by activating neural plasticity. It has been demonstrated that activation of neural plasticity has a therapeutic effect on many disorders, such as vestibular diseases, where the repetition of specific maneuvers can decrease vestibular disturbances. Sanchez et al. [29] showed that muscle contractions may change the pattern of tinnitus, from temporary worsening to temporary improvement, with repetition of the maneuvers that modulate tinnitus. A similar strategy was able to cure one of our patients who had gaze-evoked tinnitus (GET) at the Tinnitus Research Group of the University of São Paulo School of Medicine by repeating all the eye movements that evoked her tinnitus [1].

Case study: A 39-year-old woman with profound hearing loss in the right ear of unknown etiology since her teenage days developed a left vestibular schwannoma, which was subtotally removed. She received a right Nucleus 22 cochlear implant and 1 month after the surgery, at about the time the implant was activated, she developed a GET whenever her implant was on (without the implant, she had no tinnitus even with eye movements). A hissing occurred in the right ear with right- or down-gaze and in the left ear with left- or up-gaze; these symptoms had persisted for the last 4 years before the person decided to seek help. Motivated by the known benefit of vestibular rehabilitation for vestibular disorders, we recommended a habituation program to be done twice a day. The program consisted of gaze to the extreme right ten times and holding each position of gaze for 1 s before returning to the primary position of gaze. This exercise was then repeated to the left, up, and down directions of gaze. After 2 weeks, the tinnitus caused by vertical change of gaze from down stopped and the up-gaze tinnitus decreased after the 2-week period of exercise. The loudness of the tinnitus associated with vertical eye gaze decreased from 10 to 1 on the visual analog scale. However, her horizontal GET persisted and began to respond to treatment only when her gaze exercise program was modified by increasing the duration of each extreme of right and left gaze from 1 to 5 s. After 4 weeks, her right and left gaze-evoked tinnitus improved, and the loudness decreased from 10 to 6 and 10 to 2 on the visual analog scale, respectively. The increase of the duration of each gaze position from 5 to 30 s, for 3 more weeks, caused all GET to cease and not occur again out to an 18-month follow-up and at the end of all exercises. Her entire treatment program lasted 14 weeks.

The central element of vestibular rehabilitation is the repetition of a set of exercises, which compensates for the central nervous system abnormalities using eye movements, as well as cervical and body maneuvers. In the particular case discussed above, the habituation of GET occurred with the repetition of eye movements that used to trigger it. That the vertical component of patient’s GET responded sooner to treatment than the
horizontal component indicates that more than one neural network or process is involved in the habituation therapy.

Due to this surprising and long-term cure of tinnitus, we decided to use the modulation phenomenon as a basis for treating other patients through the repetition of the movements that modulate tinnitus. This treatment should be individually targeted to each patient, considering the particular movements that evoke tinnitus modulation, such as this example from work at the Tinnitus Research Group of the University of São Paulo School of Medicine.

Case Report: A 65-year-old woman with normal pure-tone thresholds complained about her constant bilateral (mostly left ear) “refrigerator’s engine” tinnitus. The loudness of her tinnitus increased from bilateral compression of her temporal area. She was asked to perform a therapy involving repetitive pressure on temporal regions from where the tinnitus could be modulated making ten repetitions, three times a day. Her tinnitus began to decrease after 7 days of such training. After 2 months, her tinnitus could no longer be modulated by pressure on her temporal area for several days on the right side; after 4 months, her tinnitus could no longer be modulated, and the baseline tinnitus on the right side disappeared for several days. Tinnitus modulation and the perception of baseline tinnitus also decreased in the left ear, although this process took a longer time than for the right ear.

We are presently studying more details about the treatment of tinnitus through the repetition of maneuvers that modulate it.

In conclusion, individuals with somatosensory tinnitus or tinnitus that can be modulated may improve by coordinated exercise of the muscles that can modulate the tinnitus.

Acknowledgments The authors thank the Tinnitus Research Initiative for the creation and support of a workgroup of Somatosensory Tinnitus and Modulating Factors.

References
3. Estola-Partanen, M (2000) Muscular tension and tinnitus: an experimental trial of trigger point injections on tinnitus [dissertation]. Tampere: Faculty of Medicine, University of Tampere


Chapter 81
Tinnitus Treatment: Botulinum Toxin

Miguel J.A. Láinez, Alejandro Ponz, and Anna Piera

Keypoints

1. Somatosensory tinnitus (objective or subjective) is tinnitus that can be modulated by stimulation of the somatosensory system.
2. Abnormal interactions between the auditory and the somatosensory nervous system that may occur at several levels of the central nervous system cause somatosensory tinnitus.
3. This chapter discusses how administration of a botulinum toxin can alleviate tinnitus and the mechanism of its action, and how that relates to its effects on chronic pain.
4. A proven benefit of botulinum toxin in patients with objective tinnitus is also discussed.

Keywords  Somatosensory tinnitus • Botulinum toxin • Autonomic pathway • Headache • Dorsal cochlear nucleus

Abbreviations

BoNT-A Botulinum toxin type A
CGRP Calcitonin gene related peptide
CNS Central nervous system
DCN Dorsal cochlear nucleus
HFS Hemifacial spasm
MSN Medullary-somatosensory nucleus
PAM Posterior auricular muscle
PTA-2 Pure tone average 2
SDS Speech discrimination scores
THI Tinnitus handicap inventory
TMJ Temporomandibular joint disorders

Introduction

While the pathophysiology of the different forms of tinnitus remains poorly understood, there is increasing evidence from electrophysiologic and functional neuroimaging studies that severe chronic tinnitus is caused by abnormal functioning of the central nervous system (CNS) [1] (see Chap. 10) brought about by activation of neural plasticity (see Chap. 12).

Somatosensory Tinnitus

The finding that people can develop tinnitus from forceful head and neck contractions [2] is an example of somatosensory tinnitus. Temporomandibular joint disorders (TMJ) are also often associated with tinnitus [3] (see Chap. 53), thus another example of somatosensory tinnitus. Effective treatment of the underlying disorder may resolve somatosensory tinnitus in some cases, but not in others [4]. The neural mechanisms of somatosensory tinnitus have been discussed elsewhere in this volume (see Chap. 9).

Reports have shown that somatic stimulation of the head or upper neck can suppress tinnitus through somatosensory pathways (see Chap. 80), supporting some forms of somatosensory tinnitus treatment such as administration of central muscle relaxants (benzodiazepines, etc.), acupuncture, biofeedback, and electrical stimulation for relaxing the muscles.
Botulinum Toxin Type A

Botulinum toxin type A (BoNT-A) is a neurotoxin. Administered locally, it can inhibit the release of acetylcholine at the neuromuscular junction [5]. It is used therapeutically in disorders of muscle hyperactivity, including movement disorders, dystonia, spasticity, cerebral palsy, gastrointestinal disorders, and urological disorders. BoNT-A was first used to treat strabismus [6], and it is now widely used in cosmetics to diminish wrinkles and frown lines because of its paralytic effect [7]. It should be noted that botulinum toxin has system effects that include cardiovascular reflexes [8].

In vitro and in vivo studies [9] have demonstrated that BoNT-A inhibits the release of nociceptive mediators such as glutamate, substance P, and calcitonin gene-related peptide (CGRP) from nociceptive fibers [10], suggesting that BoNT-A may have a direct antinociceptive action through its effects on the autonomic nervous system in addition to its neuromuscular action [11–13]. Moreover, BoNT-A, through a peripheral mechanism, has also been shown to inhibit central sensitization of central trigeminovascular neurons [14], which takes an integral part in the development, progression, and maintenance of migraine headaches [14, 15]. Central sensitization is also considered to be a potential mechanism underlying the development of chronic daily headaches in patients with migraine (see Chap. 61) [16].

Clinical trials have suggested that BoNT-A may be an effective and safe prophylactic headache medication in the treatment of migraine and other headache disorders [17]. Thus, evidence has been presented that the blockage of the autonomic pathways, and not just its paralytic effect, contributes to the ability of BoNT-A to control headaches, chronic neuropathic pain, and migraines [10, 15, 18]. These similarities between tinnitus and pain were the reason that we investigated the effect of BoNT-A on tinnitus.

Botulinum Toxin in Tinnitus Treatment

In a prospective double-blind study of the effect of BoNT-A with 30 participants with tinnitus, 26 participants completed the two parts of the study, and the results were included in the analysis. Seven of the participants’ tinnitus improved, in 3 it worsened, and in 16 it was unchanged; following placebo, the tinnitus of 2 participants improved, in 7 it worsened, and in 17 it was unchanged.

In this study, 30 patients with tinnitus were randomly placed into one of two treatment arms. Patients received either BoNT-A (20–50 units) or saline injection at the first treatment, and the opposite treatment 4 months later. Tinnitus and hearing were evaluated using questionnaires similar to the tinnitus handicap inventory (THI). Audiograms, pure-tone average-2 (PTA-2), and speech discrimination scores (SDS) were obtained prior to the first and second injection for all participants. BoNT-A or placebo were injected into three sites around the ear; 1 cm above the superior aspect of the auricle, 1 cm behind the auricle at the 2 o’clock position, and 1 cm behind the auricle at the 5 o’clock position [18].

When tinnitus was classified as “better”, “worse,” or “same” (global clinical impression estimated by patients), the treatment and placebo groups were statistically and significantly ($p<0.005$) different. Also, THI scores decreased significantly between pretreatment and 4 months after BoNT-A injection ($p=0.04$). The results of this study suggest that administration of BoNT-A may be useful in the management of tinnitus.

BoNT-A administered to the middle-ear cavity has been used to treat patients with tinnitus due to myoclonic tensor tympani contractions [19–21].

Animal studies of the possible adverse effects of administration of BoNT-A into the middle-ear cavity showed no negative effects [22].

In a study of patients with tinnitus from hemifacial spasm (HFS) in whom posterior auricular muscle is affected [23], BoNT-A was applied on the side to which the tinnitus was referred, obtaining a symptomatic improvement in 9 of 14 patients. Thus, patients who have spasm in their posterior auricular muscles (PAMs) may be candidates for treatment with botulinum toxin.

Conclusions

The effect of a local application of BoNT-A on tinnitus is assumed to be achieved through a reduction of inputs to the CNS from receptors in cervical, temporal, frontal, and periauricular muscles. This is assumed to produce a reduction of the activity in the medullary-somatosensory nucleus (MSN) (Nucleus Z), thereby
reducing the input to the dorsal cochlear nucleus (DCN). This is assumed to be the basis for the use of BoNT-A in management of chronic headaches, and similar action may explain its effect on subjective tinnitus in patients with somatosensory tinnitus.

BoNT-A is proven to be effective for treatment of patients with objective tinnitus from palatal and middle-ear myoclonus when injected into the middle-ear cavity or injected in the palatal muscles with laryngoscopic guidance.

References

Part VII
Surgical Treatments
Surgery has a definite role in the management of tinnitus associated with certain conditions as follows:

1. Surgery should be considered if hearing can be improved by surgery. Therefore, surgery plays a role in the management of tinnitus associated with conductive hearing loss. In patients with otosclerosis, tinnitus is most likely to disappear after stapes surgery (Chap. 83). Other options are tympanoplasty procedures in patients with tinnitus and chronic otitis media. Individuals with objective tinnitus due to middle-ear myoclonus will benefit from a surgical section of the tensor tympani or stapedial tendon.

Conductive and cochlear hearing loss can also be improved by conventional hearing aids. In cases of failure, surgically implanted devices can be considered (Chap. 76). If tinnitus is associated with profound bilateral hearing loss or deafness, tinnitus suppression has been reported as a secondary benefit of cochlear implantation. Recently, cochlear implantation was discussed as a new treatment option irrespective of hearing restoration for patients with severe tinnitus due to unilateral deafness (see Chap. 77).

2. Some forms of tinnitus are associated with a clear structural cause that can be improved with surgery. Pulsatile tinnitus can occur due to vascular loops in the vicinity of the eighth nerve. Microvascular decompression procedures of these loops have shown some benefit in tinnitus suppression (Chap. 84). Surgical treatment options for patients with pulsatile tinnitus of venous origin are ligation of the internal jugular vein, occlusion of the sigmoid sinus, or closure of a dural fistula. Tumors of the eighth nerve, like vestibular schwannoma, can cause tinnitus (Chap. 39). There are different surgical and non-surgical treatment options available to treat individuals with vestibular schwannoma. The surgical approaches differ in the conservation or destruction of the auditory part of the eighth nerve. The impact of surgical tumor removal on tinnitus suppression is discussed in Chap. 85. Sectioning of the eighth nerve (cochlear neurectomy) has also been tried for tinnitus suppression in individuals with tinnitus who do not have vestibular schwannoma. Improvement rates of less than 50% have to be considered in this destructive procedure as well as a chance of the condition worsening and development of complete hearing loss [1]. Therefore, candidates for this type of surgery should have no useful hearing on the affected side and should understand that effects of surgery are unpredictable.

3. Some individuals with Ménière’s disease may have tinnitus reduction from surgical treatment options. The different techniques include transtympanic application of gentamycin, endolymphatic sac surgery, and destructive procedures like labyrinthectomy or vestibular neurectomy (Chap. 83).

4. Some people with temporomandibular joint (TMJ) disorder have severe tinnitus. In most cases, nonsurgical treatment options like physical therapy or local injections of anesthetics and corticoids can restore the joint function sufficiently. Some individuals with severe conditions may benefit from surgery. Surgical intervention can range from arthroscopy to a partial or total TMJ implant (Chap. 95).

5. Only a few incidences of tinnitus are direct consequences of pathologies of the ear or the auditory
nerve. In most individuals with tinnitus, it is assumed that the tinnitus occurs as a series of reactions in the central nervous system due to deprivation of input to the central auditory pathway. Neuroimaging and neurophysiologic data suggest that chronic tinnitus is associated with focal brain activation of the auditory cortex or other parts of the auditory nervous system, such as the inferior colliculus (see Chap. 12). Therefore, targeted modulation of tinnitus-related neural hyperactivity has been considered as a new promising treatment strategy (Chap. 86). Besides the non-invasive neuromodulation techniques like repetitive transcranial magnetic stimulation (rTMS) and transcranial direct current stimulation (tDCS), an invasive direct electrical stimulation was suggested as a novel treatment option. If rTMS is capable of suppressing tinnitus transiently, the effect might be maintained by the surgical implantation of electrodes over the area of electrophysiological signal abnormality on the auditory cortex for direct electrical stimulation (Chap. 90).

Reference

Chapter 83
Surgical Treatment: The Ear

Tobias Kleinjung

Keypoints

1. Surgical restoration of hearing can improve tinnitus complaints in patients with tinnitus associated to conductive hearing loss.
2. Tinnitus is most likely to disappear after stapes surgery.
3. New-onset tinnitus or worsening of a pre-existing tinnitus can occur as an unwanted side effect of middle ear surgery.
4. Some patients with advanced Ménière’s disease might benefit from a surgical approach to their tinnitus.

Keywords Tinnitus • Ear surgery • Tympanoplasty • Stapedotomy • Stapedectomy • Ménière’s disease

Abbreviations

PORP Partial ossicular replacement
TORP Total ossicular replacement

Introduction

Any kind of conductive hearing loss may be accompanied by tinnitus, as outlined in detail in Chap. 34. Surgical efforts to improve hearing loss can, in some cases, also bring about the partial or complete remission of tinnitus. This chapter will discuss the possibility of reducing tinnitus through surgical operations in order to treat conductive hearing loss. The text will describe surgical procedures involving the middle ear that are indicated for treatment of different forms of objective tinnitus. Finally, it will also discuss otological surgery techniques used in the management of Ménière’s disease. The topics of cochlear implant surgery (see Chap. 77) and surgery of the internal auditory canal (see Chap. 85) are dealt with in separate chapters. If the accompanying tinnitus is due to hearing loss, irrespective of its duration, then restoration of hearing can be beneficial to management of tinnitus, in addition to improving hearing (see Chap. 10).

Surgery of the External Auditory Canal

Space-occupying lesions that completely or partially obliterate the external auditory canal and lead to conductive hearing loss must be removed. This applies both to benign changes, such as auditory canal exostoses, and to malignant tumors. If normal hearing is restored after uncomplicated healing, any tinnitus that may have been present preoperatively can also be expected to resolve completely.

Middle Ear Surgery

Myringotomy with Tube Insertion

Myringotomy, followed by aspiration of fluid build-up in the middle ear and insertion of a small tube in the opening of the tympanic membrane, brings immediate relief of symptoms in cases of otitis media with
effusion. This procedure may also result in remission of tinnitus along with a reduction of aural fullness and conductive hearing loss. Myringotomy, with or without tube insertion, can also positively influence the course of acute otitis media that does not respond favorably to pharmacological therapy.

**Tympanoplasty**

“Tympanoplasty” is the term used to describe the surgical repair of the tympanic membrane after a perforation. This process includes inspection of the ossicular chain and, if necessary, its reconstruction by ossiculoplasty.

According to Wullstein, depending on the extent of reconstruction involved, there are five different types of tympanoplasty [1]. Tympanoplasty Type I merely involves the restoration of the perforated tympanic membrane by grafting. The ossicular chain is intact. In Type II and III procedures, ossiculoplasty is an integral part of tympanoplasty. Tympanoplasty Type II is a procedure in which the patient’s own auditory ossicles (parts of the incus or the head of the malleus), i.e., organic material, are used for the reconstruction. In tympanoplasty Type III, alloplastic materials are used.

The defective ossicles are repaired using synthetic prostheses that replace the incus and are placed on the intact stapedial head (partial ossicular replacement prosthesis, PORP) or by prostheses that replace the incus and stapedial suprastructure and are placed directly on the intact stapes footplate (total ossicular replacement prosthesis, TORP). Tympanoplasty Types IV and V no longer play a role in middle ear surgery today.

The techniques of tympanoplasty have an application in the treatment of chronic otitis media. In chronic mesotympanic otitis media (chronic suppurative otitis media), reconstruction of the sound conduction mechanism is necessary in 20–25% of cases. In cholesteatoma, 80% of patients require tympanoplasty Type III [2]. Depending on the underlying pathology, the technique of tympanoplasty may be combined with procedures involving the external auditory canal (e.g., canalooplasty) and mastoid (e.g., mastoidectomy). The technique of tympanoplasty is used to correct malformations of the middle ear and following persistent traumatic eardrum perforation. Since the advent of microscopic middle ear surgery in the 1950s, many tympanoplasty techniques have been described. The techniques differ in terms of the approach, such as transcanal, endaural, retroauricular, graft material, used for tympanic membrane replacement (e.g., temporalis fascia, cartilage), and the design of the prosthesis and materials used (e.g., homologous incus, hydroxyapatite, gold, titanium) [2]. All methods aim at achieving complete eradication of infection, repairing the defective tympanic membrane, and improving hearing. These are the topics primarily addressed in the literature. Publication of results regarding relief of preoperative tinnitus has been few (summarized in Table 83.1). Nevertheless, results currently available show approximately 30% of patients who had tympanoplasty are no longer aware of tinnitus. In two of the three published studies, complete remission of tinnitus was achieved [3, 4] in about one-third of patients and more than 40% had partial remission and 4–8% became worse (Table 83.1). The assessment offered by Helms in an older study from 1981 [5] showed that one-third became worse. The improvement in tinnitus symptoms after surgery may be attributed primarily to closure of the airbone gap. Accordingly, Lima Ada et al. [4] found a good correlation between postoperative hearing improvement and the reduction in tinnitus. In those patients who continue to suffer from tinnitus despite adequate hearing improvement, there must have been other causes for the reduced sound stimulation of tinnitus [4].

**Stapes Surgery**

Conductive hearing loss and tinnitus are the main symptoms of otosclerosis. With the development of microscopic middle ear surgery in the 1950s, surgical mobilization of the stapes in otosclerosis became the

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focal point of interest among middle ear surgeons. In 1958, Shea performed the first stapedectomy, where the stapes was replaced by prosthesis [6, 7]. In the following years, the technique of stapedotomy has evolved into a standard procedure, where only the suprastructure of the stapes is removed, a perforation of the footplate is made, and a piston prosthesis is attached to the long process of the incus extending into the perforation of the stapes footplate (for review, see [8]). The introduction of laser surgery permitted “no-touch” perforation of the stapes footplate [9]. The main objective of stapes surgery is to improve hearing. The most published studies regard hearing improvement achieved from different techniques. Hearing is improved in about 90% of patients [10] and approximately 60% of patients have an air-bone gap of between 0 and 10 dB [11]. No improvement in hearing occurs in approximately 8% of patients and a deterioration of hearing loss, including deafness, occurs in 2% of patients [10]. Few studies regarding the effect on tinnitus from stapes surgery have been published. Table 83.2 provides a summary of comparable studies conducted since 1990. On the average, complete remission of tinnitus was achieved in approximately half of patients from stapes surgery. Partial remission was achieved in 30% and approximately 80% of those who had stapes surgery benefited from the operation. Most of the remainder had no change, and fewer than 5% of patients reported worsening of their tinnitus.

Many studies showed improvement regarding tinnitus that was independent of the hearing improvement [12, 13], but one older study by Glasgold et al. [14] showed a correlation between hearing improvement and tinnitus improvement. Ayache et al. [15] found no difference in reduction of tinnitus between stapedotomy and stapedectomy. Sakai et al. [16] and Gersdorff et al. [12], on the other hand, noted better results after stapedotomy than with stapedectomy. No significant correlation between gender, tinnitus frequency, tinnitus duration, or extent of hearing loss and the effect on the tinnitus from stapes surgery was reported [15]. These factors, therefore, do not have prognostic value for stapes operation. It is unclear whether the positive effect of the stapes surgery is due to the improvement of hearing or some other factors related to mobilization of the fixed footplate. The fact that many patients already experience an improvement in tinnitus immediately after surgery – i.e., in a state when the auditory canal is packed – favor the latter hypothesis.

### Middle Ear Surgery for Objective Tinnitus

Objective tinnitus may be either vascular or muscular in nature. Objective tinnitus often accompanies disorders such as glomus tumors. When patients with glomus tumors are treated surgically, complete eradication of the pathological process is the main aim in the resection of such tumors. The patient’s pulsatile tinnitus most often disappears, which is an additional benefit of the surgery. After embolizing the vessels feeding the tumor, resection of a glomus tumor located in the middle ear cavity is done using the same approaches as those for tympanoplasty. Reconstruction of the tympanic membrane and ossicular chain may be necessary. Surgery to excise glomus jugulare tumors is different and requires a wide approach via the lateral skull base [17].

Treatment of objective tinnitus, caused by contraction (repetitive myoclonus) of the middle ear muscles that results in rhythmic tinnitus, is to section the tendons of the stapedius or tensor tympani muscles [18, 19].

When the Eustachian tube fails to close normally, disabling breath-synchronous tinnitus may result.

### Table 83.2 Effects of stapes surgery on tinnitus

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<td>Da Silva Lima et al. [47]</td>
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<td>Ramsay et al. [49]</td>
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<td>Szymanski et al. [13]</td>
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The first remedy may be inserting a tympanostomy tube [20]. Other surgical procedures aim at narrowing or occluding the Eustachian tube from the middle ear or the nasopharynx. These methods are in a preliminary trial phase. Irreversible measures, such as the paratubular implantation of a Teflon graft [21], may lead to chronic otitis media due to Eustachian tube dysfunction if there is overcorrection. Endoscopic application of absorbable substances (hyaluronic acid, collagen) into the tubal elevation may bring transient symptom relief and is easy to regulate [22, 23].

**Tinnitus as a Risk in Middle Ear Surgery**

While middle ear surgery offers benefit regarding management of tinnitus, it can also, as a side effect, cause tinnitus or worsen existing tinnitus. Postoperative deafness is a serious complication of any middle ear surgery that can occur from intra-operative damage to cochlear structures. Together with deafness and vertigo, tinnitus may occur or pre-existing tinnitus may become worse [24]. The risk of serious postoperative inner ear damage in stapes surgery has been reported to be between 0.5 and 1% [2].

Operations for chronic otitis media extensive cholesteatoma resections, in particular, may have higher risks of postoperative tinnitus from damage to the inner ear from stapes luxation or development of a semicircular canal fistula [25]. In operations of large cholesteatoma involving the stapes, it is therefore recommended that some cholesteatoma be left behind. The remaining cholesteatoma then can be resected in a second operation after 6–9 months [26].

Surgical procedures directed at the middle ear or the external auditory canal involve extensive drilling which carries a risk of noise-induced hearing loss and increased risk of tinnitus. In addition, the effect of surgery and anesthesia on the central nervous system may explain postoperative increases in tinnitus.

**Ear Surgery and Ménière’s Disease**

Two kinds of surgical treatment for advanced Ménière’s disease are in use (see Chaps. 38 and 60). One is conservative and spares hearing while the other is a destructive procedure. Conservative procedures compromising endolymphatic sac decompression, gentamycin infusion, are indicated when the symptoms are dominated by a frequent occurrence of severe attacks of vertigo with some residual hearing preserved. A third conservative procedure is an application of air puffs to the inner ear via a ventilation tube in the eardrum [27]. Surgery on the indication of tinnitus is rarely performed, and only in connection with deafness and reduced vertigo, when the tinnitus is highly distressing and resistive to other treatments.

Endolymphatic sac surgery is the most common surgical procedure for Ménière’s disease. Its purpose is to treat the endolymphatic hydrops by inserting a permanent drain from the endolymphatic sac to the middle ear space. This is achieved by wide exposure of the endolymphatic sac following decompression of the sigmoid sinus in a mastoidectomy [28]. After saccotomy, silicone sheeting is inserted into the sac lumen to allow permanent drainage. The risk of suffering additional sensorineural hearing loss with this technique has been estimated at less than 2% [29]. Some debate surrounds the success rates obtained with this procedure in terms of the control of vertigo and tinnitus. Most of the reports published in the literature relate to vertigo, with successful vertigo control claimed to be achieved in 70–90% of cases [30, 31]. The success rate for tinnitus control with endolymphatic sac surgery is lower than that for vertigo control, with improvement or complete remission of tinnitus being reported in the 30–40% range [32, 33].

These reported success rates of endolymphatic sac decompression are being questioned after studies by Thomsen et al. [34] and Bretlau et al. [35], which showed similar effect of a placebo (sham) operation involving a classic mastoidectomy without decompression as was obtained in real endolymphatic sac decompression. Both real and placebo operations had success rates of 75%.

A similar placebo effect has also been attributed to insertion of a ventilating tube into the tympanic membrane with no additional measures [36], a technique that continues to find use in surgical practice [37]. Success rates for destructive procedures, such as labyrinthectomy, in improving tinnitus are not higher as compared with endolymphatic decompression. Labyrinthectomy is reported to improve tinnitus in 40% of the patients [38]. This rarely performed procedure may be considered for tinnitus and vertigo control.
in patients with Ménière’s disease who have very poor hearing or are deaf. Higher success rates have been achieved when labyrinthectomy is combined with cochleovestibular neurectomy [39]. According to Jones et al. [40], good postoperative control of tinnitus can be expected with this technique in slightly less than 70% of cases. Effective control of vertigo symptoms can be achieved with this combination in nearly 100% of the patients [39].

Intratympanic gentamicin perfusion is designed to achieve chemical partial ablation of the vestibular system while still preserving cochlear function. For that, gentamicin is instilled into the middle ear cavity via a tube inserted in the ear drum or by direct puncture of the tympanic membrane. It diffuses across the round window membrane to reach the inner ear [41]. Several published studies have shown that good vertigo control can be achieved in 70–90% of the patients [28, 41, 42]. It is an advantage of this method that morbidity is low and the incidence of sensorineural hearing loss has been reduced to about 20% of all those treated [43]. This technique is currently regarded as the standard therapy for controlling vertigo attacks [42]. Little has been reported regarding the effect on tinnitus from gentamicin treatment. In one study, Lange et al. [44] reported improvement in tinnitus in 26 out of 56 patients treated (46%). Two small studies reported improvements in tinnitus in only 5 and 27% of patients, respectively [45, 46].

References

2. Hildmann, H, Sudhoff, H, Middle ear surgery 2006, Heidelberg: Springer
5. Helms, J, Tympanoplasty and Tinnitus (author’s transl) Laryngol Rhinol Otol (Stuttg), 1981 60(3):99–100
17. Fisch, U and Mattox, D, Microsurgery of the skull base 1998, Stuttgart: Georg Thieme
22. Mees, K and Beimert, U, [Correction of the gaping eustachian tube osteum with injectable collagen] Laryngol Rhinol Otol (Stuttg), 1988 67(2):87
45. Smith, WK, Sandooram, D, and Prinsley, PR, Intratympanic gentamicin treatment in Meniere’s disease: patients’ experiences and outcomes J Laryngol Otol, 2006 120(9):730–5
Chapter 84
Long-Term Follow-Up of Microvascular Decompression for Tinnitus

Jacques Magnan, Benoit Lafont, and Charbel Rameh

Keypoints

1. The concept of cochlear nerve compression by an offending vessel as a cause for a small percentage of tinnitus cases is a controversial topic.

2. The diagnosis of tinnitus secondary to a neurovascular conflict requires a combination of the following:
   (a) Unilateral tinnitus
   (b) Radiographic presence of a vascular compression of the cochlear nerve
   (c) Prolonged latencies of waves I–III in ABR

3. Following these criteria, 43 patients between 1993 and 2006 underwent an endoscope-assisted microvascular decompression of the cochlear nerve via a minimally invasive retrosigmoid approach.

4. Our results were studied 1 week, 2 months, and 2 years postoperatively. On the long term, there was resolution of tinnitus in 9 cases (21%), a marked decrease of tinnitus in 13 cases (31%), and no change of tinnitus in 19 cases (44%).

5. A significant statistical correlation was found between the resolution of tinnitus and the improvement of hearing postoperatively.

6. The best results were in cases where the subarcuate artery was responsible for the conflict. The worse results were when the course of the offending vessel (AICA) was inside the internal auditory canal or between the facial and vestibulocochlear nerves.

7. Although the results of vascular decompression of the cochlear nerve involving select cases with incapacitating tinnitus are less rewarding than those reached in hemifacial spasm or trigeminal neuralgia, they confirm the hypothesis of vascular compression syndrome of the auditory nerve and the need for better selection criteria.

Keywords Tinnitus • Endoscope assisted • Microvascular decompression • Results

Abbreviations

ABR Auditory brainstem responses
AICA Anterior inferior cerebellar artery
CCAP Cochlear compound action potential
CISS Constructive interference in steady state
CPA Cerebella-pontine angle
CSF Cerebrospinal fluid
FT Fourier transform
IAC Internal auditory meatus
MRI Magnetic resonance imaging
MVD Microvascular decompression
PIA Posterior inferior cerebella artery
VNG Vestibulonystagmography

Introduction

There are several etiologies for tinnitus. Among these, cochlear nerve compression by an arterial loop or a vein in the cerebellopontine angle (CPA) represents a very small percentage, which is difficult to quantify (see also Chaps. 10 and 40). Therefore, the vascular compression theory of the auditory cranial nerve is still questionable and has not yet gained the wide acceptance that trigeminal and facial nerve decompression for trigeminal neuralgia and hemifacial spasm has received.
By presenting our results on endoscope-assisted microvascular decompression (MVD) of the auditory nerve, we will try to shed the light on this concept which was first suggested in 1934 by Dandy [1], popularized in 1975 by Jannetta [2], and utilized in 1993 by Møller and Jannetta [3] for the management of severe tinnitus.

**Microsurgical Anatomy of the Cochlear Nerve**

A prolonged contact or “conflict” between the cochlear nerve and an adjacent vascular structure will lead to an alteration in the myelin sheath. Studies have shown a better resistance of the peripheral myelin compared to the central myelin to such insults. Thus, the configuration of central myelin in the cochlear nerve has significant implications. The area of central myelin forming the root entry zone of the nerve is truly a cone shape and extends as far as the internal auditory canal. Therefore, a blood vessel can cause symptoms by compression not only at the junctional area close to the brainstem, as in the case of the motor facial nerve, but anywhere along the course of the nerve, whether cisternal or intracanalicular. The area of the nerve surrounded by central myelin, called the transitional glial zone or “Obersteiner-Redlich” zone, is fragile and very sensitive to external compressions. Its length is variable between the different cranial nerves. It is 1.2 mm for the glossopharyngeal nerve, 1.7 mm for the facial nerve, 2.6 mm for the trigeminal nerve, and 8.3 mm for the cochlear nerve [4].

The permanent and pulsatile contact of an artery with the transitional zone of the nerve will lead to histological and physiopathological changes in the root of the nerve [5].

1. Local demyelination (Fig. 84.1) will result in aberrant connections between the axons and ultimately to the phenomenon of localized hyper-excitability (Fig. 84.2).
2. Stimulation of neighboring axons and the orthodromic recruitment (ephapse phenomenon) [6].
3. Antidromic stimulation of neighboring neurons and the permanent nuclear hyperactivity (Kindling effect) [7, 8].
4. Endoneural fibrosis of the compressed zone: this leads to an alteration in nerve conduction and a change in the auditory brainstem-evoked response (ABR) [5].

**Patient Selection Criteria for Operative Treatment**

The specification of tinnitus due to a neurovascular conflict is difficult. The decision to perform a microvascular decompression is not made unless a full workup has been done. Patients with tinnitus are quite
demanding of a radical solution, and thus decision making should be very careful. The patients must have specific clinical evaluations and a battery of tests before being advised of surgery.

Most important is the unilaterality of tinnitus. In addition to the audiogram, we order a magnetic resonance imaging (MRI) to search for any associated pathology along the course of the auditory pathways, in particular at the level of the CPA.

**Clinical Criteria**

Tinnitus has to be unilateral, severe (following Reed’s classification, significantly affecting the quality of life and considered as a handicap for the patient) [9], present over several months, and persistent despite therapy with the available medical treatments.

The pulsatile nature and the tonality of the tinnitus are not decision-influencing criteria. In addition, tinnitus associated with a low-frequency fluctuating hearing loss as in Ménière’s disease or with progressive hearing loss was for us an exclusion criterion for MVD.

**Radiological Criteria**

MRI is able to reproduce the anatomy of the CPA and demonstrate the presence of a neurovascular conflict. It also eliminates other potential causes of unilateral tinnitus such as an acoustic neuroma. MRI in both T1 and T2 three-dimensional Fourier transfer (FT) is the most effective method of delineating both the acoustic facial nerve bundle and the surrounding vascular structures in the CPA and the IAC. T2 is carried out using constructive interference in steady-state (CISS) sequence (Fig. 84.3). Also, postcontrast reformatted turbo flash in the axial plane helps in delineating the conflict vessels [10]. The image assessment includes all the CPA with serial thin slices of 0.4 mm thickness.

To confirm the diagnosis, the single presence of a neurovascular contact is not sufficient. Other radiological criteria are required (Figs. 84.3, 84.4, 84.5):

1. Perpendicular contact between the vascular loop and the nerve along two different perpendicular planes.
2. Displacement of the cochlear nerve, with a certain distance between the facial and cochlear nerves.
3. Imprint on the cochlear nerve and reduction in its diameter.
4. Brainstem distortion caused by the vascular structure at the level of the root entry zone of the cochlear nerve.

The presence of an arterial loop in the internal auditory canal (IAC) may be responsible for tinnitus but is not a sufficient criterion by itself.

**Electrophysiological Criteria**

1. Pure-tone audiometry: Pure-tone audiometry can identify an unilateral sensorineural hearing loss, on the same side as the tinnitus. In general, the hearing loss is of moderate severity over all the frequencies tested (250–8 KHz) but occurs predominantly at the higher frequencies. This hearing loss is of statistical significance when we compare it to the contralateral ear ($p<0.05$) and is proof that the nerve is in a pathological state.

2. Auditory brainstem response (ABR): ABR, in the case of a retrocochlear lesion, will reveal a prolongation of the latencies or a desynchronization. We use Møller criteria to identify pathological ABPrs with a prolonged interwave latency of peaks I–III of more than 0.2 ms (Fig. 84.6) [3].

3. Vestibulonystagmography (VNG): Vestibular tests allow a search for an ipsilateral vestibular deficit on the same side as the tinnitus, though it may be a subclinical deficit. The vascular compression might be responsible for a vestibular deficit following the same mechanism as that of tinnitus.

In summary, the diagnosis of a neurovascular conflict with the cochlear nerve requires the combination of a

![Fig. 84.6](image) A typical ABR confirming a vascular compression of the right cochlear nerve with prolonged interwave latencies I–III and I–V, and de-synchronisation of the wave.
clinically incapacitating unilateral tinnitus, radiological presence of a vascular compression, and electrophysiological alteration of ABR with increased latencies or de-synchronization of waves.

These criteria should convince the surgeon of the involvement of the CPA vessel in the etiology of tinnitus and the possible benefit of the decompression. It was not uncommon that simply making the diagnosis of a vascular compression on the cochlear nerve as a cause of tinnitus reassured the patient, rendered the tinnitus more tolerable, and consequently did not require operative treatment.

**Minimally Invasive Surgery of the Cerebellopontine Angle Using a Keyhole Retrosigmoid Approach**

For this procedure, the patient is in a dorsal decubitus position, with the head flexed and turned to the contralateral side. General endotracheal anesthesia uses analgesics, hypnotics, or neuroleptics (Diprivan, Sulfentanyl). The patient is hyperventilated to obtain a hypocapnea (Pressure of CO$_2$ around 25 mmHg at the time of dural opening) to diminish the intracranial pressure and help spontaneous cerebellar retraction. Due to this process, lumbar puncture and mannitol solution are no longer used.

The electrodes monitoring the facial or cochlear nerve are put in place after the induction of anesthesia. With intraoperative ABR monitoring, we obtain the cochlear compound action potentials (CCAPs) from the entire surface of the cochlear nerve with a surface multipolar electrode. The return of the ABR to normal after the decompression is a sign of good prognosis, but unfortunately it is not constant.

A curvilinear retro-auricular skin incision is made, two finger widths behind the pinna. It is 6–8 cm long and passes over the posterior part of the anticipated craniotomy. The cutaneous flap is anteriorly based, while the underlying musculoperiosteal flap is fashioned to be posteriorly based. The mastoid emissary vein is identified. Drilling for the craniotomy is centered on the emissary vein and is done using a cutting then a diamond burr. Bone dust is collected to make a bone pâté that will be used in closure. The craniotomy is usually elliptical in shape, 20 mm×10 mm in dimensions. It must reach the posterior border of the sigmoid sinus without skeletonizing it. Any mastoid cells should be obliterated with bone wax to prevent cerebrospinal fluid (CSF) rhinorrhea.

Opening of the dura is done under the operating microscope. The dural flap is based and suspended anteriorly. The cerebellum is protected using a synthetic dura mater. The posterior cistern is opened inferiorly at the level of the lower cranial nerves. The arachnoid wrapping surrounding the acoustico-facial nerve bundle is dissected to expose the neurovascular conflict. The offending vessels characteristically induce pressure and distort the cochlear nerve at any place along its course. The compressive effect is most commonly due to sharp-angled loops at areas of vessel bifurcations. Two other typical aspects are when the loop pinches the nerve or when the subarcuate artery constricts the nerve.

Using the 4 mm 30° endoscope, a panoramic view of all the CPA structures is obtained. The tip of the endoscope is passed above and below the acoustico-facial bundle to identify the precise location and course of the offending vessels.

Whatever be the location of the neurovascular conflict, the purpose of the microvascular decompression procedure is to change the axis of the offending vascular loop and keep it away from the offended nerve. This surgical maneuver is done under the operating microscope. First, the offending vessel is carefully mobilized using microelevators and microhooks, respecting the labyrinthine and perforating arteries. Then, the MVD is performed using a small Teflon pad which is adjusted with a microhook to isolate the nerve from the artery. Teflon is an inert material very well tolerated in the CPA. The most common offending vessel, the AICA, has a course between the seventh and eighth nerve in 50% of cases, which prevents a fully efficient and complete microvascular decompression from being completed. An intracanalicular AICA loop requires a complementary drilling of the internal auditory canal. CCAPs are monitored during decompression (Figs. 84.7 and 84.8) by placing the recording electrode beneath the flocculus or above the cochlear nerve depending on the course of the offending vessels.

The operation ends by another endoscopic control of the quality of the surgical act to confirm the good positioning of the Teflon pad.

At the end of the procedure, the dura mater is sutured meticulously using 5/0 silk sutures. Then, the craniotomy is filled with a mixture of bone pâté and fibrin glue.
The musculo-aponeurotic flap is sutured in place. The subcutaneous layer is sutured followed by skin closure.

The duration of the intervention is usually between 1.5 and 2 h.

The patient stays in the recovery room for 2 h and is then transferred to his/her room without the need for an intensive care unit (ICU), no matter what the patient’s age is [11].

Results

Literature

The literature has sufficient data to prove that MVD can cure severe incapacitating tinnitus in carefully selected patients [3, 12–15]. However, the success rate of this procedure is inferior to that of MVD for hemifacial spasm or trigeminal neuralgia. This is mainly due to the anatomic difficulty in isolating the offending vessels as well as the multitude of other confounding etiologies for tinnitus. Following microvascular decompression for tinnitus, a result considered as “good” is a patient who had a total resolution or a significant decrease in tinnitus [3, 12, 13]. The frequency of “good” results in the literature has ranged from 33 to 77% with more success in women than in men [3]. Some authors have performed revision surgeries before obtaining their final “good” results of 77% [12]. The only report that exceeded this range was by Okamura et al [16] who had 94% good results following the surgery. However, their selection criteria were much more flexible than the ones we used for our patients, and they included cases of low-frequency fluctuating hearing loss and intermittent tinnitus [16].

As for the offending vessels and their relation to the results of MVD, it seems that when the vertebral artery was the offender, decompression gave the best results, compared to AICA or PICA decompression [13].

Some authors have found a relation between the duration and character of the tinnitus and the postoperative results [3], while others found no such relation [13]. Several authors also report that ABR returns to normal in patients who had good results for the tinnitus [13], and that hearing loss is also improved [16].

Our Results

Between 1993 and 2006, 60 patients with tinnitus underwent endoscope-assisted microvascular decompression. Among them, 43 patients (22 women and 21 men) had a long-term follow-up. Their ages ranged from 37 to 71 years, and the average patient age was 57 years. The mean duration of tinnitus before surgery was 3.2 years (the range was between 1 and 8 years).

Tinnitus

In the immediate postoperative period (7 days following surgery), we have found the following results (Table 84.1):

1. Total relief in 16 patients (37%)
2. Marked improvement in 11 patients (26%)
3. No improvement in 16 patients (37%)
4. Worsening in 1 patient (2%)

At 2 months postoperatively, the results are the following:

1. Total relief in 9 patients (21%)
2. Marked improvement in 14 patients (33%)
3. No improvement in 19 patients (44%)
4. Worsening in 1 patient (2%)
Fig. 84.8 Modification of CCAPS during the vascular decompression surgery.

Table 84.1 Changes in tinnitus following microvascular decompression

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<th>Post-operative Follow up Time</th>
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<td>Total relief</td>
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<td>1 week</td>
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<tr>
<td>2 months</td>
<td>16</td>
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<td>2 years</td>
<td>14</td>
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At 2 years postoperatively, the results are the following:

1. Total relief in 9 patients (21%)
2. Marked improvement in 13 patients (31%)
3. No improvement in 19 patients (44%)
4. Worsening in 2 patients (4%)

The characteristics of the tinnitus (pulsatility, pitch, persistence, or intermittency) are very labile parameters that were not quantified.

In our series, the results did not change significantly with time after 2 months postoperatively. This is not consistent with the data of other authors who mention some improvement until up to 1 year.

**Postoperative Hearing**

It is reasonable to believe that when pulsatile compression on the cochlear nerve is released, hearing loss due to such compression might be improved. We considered that there is an improvement in hearing if there is a gain superior or equal to 5 dB over a minimum of three standard frequencies of the audiogram. As such, we found that 12 patients had an improvement of hearing over three or more frequencies. On the preoperative audiograms, we notice that the high frequencies were mostly altered (4,000 and 8,000 Hz). We found an improvement by more than 5 dB over these two frequencies in 17 patients, but this improvement did not reach statistical significance ($p > 0.05$). Hearing remained unchanged in 20 subjects. On the other hand, eight patients presented a 5–10 dB worsening of their hearing, and one patient had a more significant loss of more than 30 dB. Two causes might explain this complication: the manipulation of the arteries, or the thermal effect from the light of the tip of the endoscope. In the year following the surgery, two other patients had a loss of hearing: one patient had a sudden hearing loss 11 weeks after the surgery, which never recovered, and another patient had a fluctuating hearing loss.

**Correlation Between Tinnitus and Hearing**

There exists a significant statistical correlation between the resolution of the tinnitus and the improvement in hearing. In fact, a 5 dB improvement over at least three frequencies of the audiogram is statistically associated with a resolution of the tinnitus (Fisher exact test) ($p = 0.034$ for an improvement over three frequencies, $p = 0.009$ over four frequencies, and $p = 0.050$ over five frequencies).

In addition, the sum of the auditory gains over the seven tested frequencies (250, 500, 1,000, 2,000, 4,000, 6,000, and 8,000 Hz) is statistically better in patients without any postoperative tinnitus compared to those who still have tinnitus. These former patients present an average sum auditory gain of 20 dB ($p = 0.039$).

**Correlation Between Tinnitus and ABR**

The ABR returned to normal (I–III < 0.2 ms relative to the non-operated side) in all patients who had successful results (total relief and marked improvement patients).

**Correlation Between Tinnitus and VNG**

Out of our 43 patients, only one had vestibular test normalization with relief of tinnitus.

**Vascular Structures Responsible for the Conflict**

We have grouped the causes of cochlear nerve compression into the following groups:

1. AICA only: 18/43 (39%) (Fig. 8.4.9)
2. AICA and PICA (pinching the nerve): 4/43 (9%) (Fig. 8.4.10)
3. PICA with or without vertebral artery: 12/43 (28%) (Fig. 8.4.11)
4. Subarcuate artery (“hugging” the cochlear nerve): 7/43 (19%) (Fig. 8.4.12)
5. Vein: 2/43 (5%) (when present, the cause effect is questionable).

Though it might be difficult to predict the clinical results in relation to the incriminated vessel, it seems that the seven patients whose conflicts were related to the subarcuate artery had the best results. Among the
Fig. 84.9 AICA at the level of the porus compressing the cochlear nerve (tinnitus was reduced in this patient). (a): AICA and acoustico-facial nerve bundle within the arachnoid wrapping. (b): Closer view showing the offending contact between the vessel and the nerve at the entrance of the internal auditory canal. (c): Decompression of the vascular loop which is mobilised medially and inferiorly. (d): Vascular decompression done, not the labyrinthine artery emerging from the loop.

Fig. 84.10 Different loops of the AICA pinching the cochlear nerve at its root entry zone (tinnitus was relieved in this patient). (a): Surgical aspect of the same vascular structure AICA looping around the cochlear nerve and compressing both sides of the nerve. (b): Vascular decompression using several small Teflon pads and isolating the cochlear nerve.
seven operated cases, four had a total resolution of tinnitus, two had a decrease in tinnitus, and one had no change. On the other hand, in cases of an aberrant vein or a vascular loop inside the internal auditory canal, there was no improvement at all. For the other cases, no statistical correlation was found between the incriminated vessel and the postoperative result.

**Morbidity**

In our neuro-otological experience (1,117 retrosigmoid approach surgeries for functional indications), we have very little morbidity. This should be the rule in the functional surgical indications. In the surgeries of cochlear nerve decompression, no neurological sequelae were present postoperatively. There was no facial paralysis. There was one case of a profound hearing loss in the immediate postoperative period, probably related to a thermal heating effect from the tip of the endoscope. This happened in the earlier stages of our development of this technique. Two other hearing loss cases occurred in the first year after surgery. In one case, a sudden hearing loss appeared and did not recover despite adequate medical treatment. In the other case, a fluctuating hearing loss occurred, with vertigo, establishing the diagnosis of Ménière’s disease that manifested itself 6 months postoperatively.

In total, based on our experience with the MVD surgery using the minimally invasive endoscope-assisted...
retrosigmoid approach, we can reduce the patient morbidity whatever the patient age is, but we did not succeed in increasing the rate of successful results above 50%. MVD of the auditory cranial nerve is less rewarding when compared to MVD of the trigeminal or the facial cranial nerves. This can be explained first by the difficulty in making a certain diagnosis, and second, by the complexity of the offending vascular loop anatomy which could limit the surgical maneuver.

**Conclusion**

When the neurovascular conflict is proven, decompression represents the only real treatment for these cases of tinnitus. This hypothesis is supported by the comparable management of hemifacial spasm and trigeminal neuralgia.

Our patient selection criteria are: unilateral tinnitus, positive offending vascular loop on MRI, and retrocochlear ABR abnormalities.

Out of 43 patients operated on by MVD of the cochlear nerve, the result was relief of tinnitus in 21% and decrease in its intensity in 31% of patients. There was a significant statistical correlation between the resolution of tinnitus and the postoperative hearing improvement confirming the concept of compression of the cochlear nerve. The failures of the eighth cranial nerve MVD must be analyzed with respect to the complex process of patient selection that is not yet definitively defined, the surgical findings, which can limit the efficacy of the surgical procedure, and the duration of the tinnitus with a central component.

**References**

6. Arvanitaki A (1942) Effect evoked in an axon by the activity of a contiguous one, J Neurophysiol 5, 89–108
9. Reed GF (1960) An audiometric study of two hundred cases of subjective tinnitus, AMA Arch otolaryngol 71, 84–94
Chapter 85
Vestibular Schwannoma

Dirk De Ridder

Keypoints

1. Tinnitus is a common symptom in individuals with vestibular schwannoma (VS).
2. In 70–80% of cases, VS is ipsilateral to the tumor. It is the principal symptom in 10% and is moderate to severe in 14% of individuals.
3. Tinnitus in VS is typically associated with hearing loss (95%) and disorders of balance (50%).
4. Tinnitus in VS is mostly of high pitch.
5. Tinnitus is the predominant symptom in very small and very large tumors.
6. On the average tinnitus is not altered by gamma knife treatment, translabyrinth surgery, or retrosigmoid microneurosurgery, but the tinnitus of individual patients may improve or worsen after such treatments.
7. After retrosigmoid microneurosurgery tinnitus is often less troublesome than before the surgery.
8. Tinnitus might worsen after retrosigmoid tumor removal in operations in which attempts are made to spare hearing.

Keywords  Tinnitus • Vestibular schwannoma • Acoustic neuroma • Gamma knife • Microsurgery • Natural history

Abbreviations

Gy  Gray
SRS  Stereotactic radiosurgery

Introduction

The name “acoustic neuroma” (correctly known as vestibular schwannoma) is a misnomer because histologically the benign lesions derive from the neurilemma sheath, most commonly of the superior vestibular nerve and not the auditory nerve [1]. These tumors occur in two forms: 95% are unilateral, mostly presenting after the age of 50; 5% occur bilaterally as a part of the signs of neurofibromatosis type 2, presenting at younger age (often before the age of 30) [2]. In neurofibromatosis type 1, unilateral VS are rare, with an incidence of about 2%. Bilateral VS are virtually non-existent in neurofibromatosis type 1 [2].

The incidence of VS is 1/100,000, 6% of intracranial tumors [2]. Postmortem studies indicate that the actual incidence is much higher (0.8%) [3] because many schwannoma remain asymptomatic.

Some epidemiological studies have found an association between the use of mobile phones and the incidence of VS [4–6], whereas other studies have not found such association [7]. Similarly, some epidemiological studies suggest an association between loud noise and VS [8].

The symptoms and signs of VS depend on the size of the lesion. When still intracanalicular, VS typically presents with a triad of symptoms: unilateral tinnitus (73%) [9], hearing loss (90–98%) [9–11], and balance problems (50%) [2]. The most common symptom is unilateral hearing loss [11], predominantly in high frequencies [2]. The hearing loss is progressive in most individuals with VS, but sudden deafness occurs in
Low-frequency hearing loss occurs in individuals with large tumors [13]. Disequilibrium is more common than vertigo, and when disequilibrium is the first symptom, it seems to be correlated with rapidly growing tumors [14]. VS-related vertigo often occurs without nausea [10]. Tinnitus usually develops when the tumor is still intracanalicular [11] and is reported as the principal symptom in 1 of 10 patients who seek medical help [9]. Tinnitus is more common in small and large tumors than in medium-sized tumors [11]. When tumors extend into the cisternal space, hearing loss worsens and associated symptoms from compression of nearby cranial nerves may develop, such as facial numbness (in the midface area), facial palsy, otalgia, changes in taste, and, rarely, hoarseness or dysphagia. Compression of the brainstem and cerebellum can induce ataxia, diplopia, and other cerebellar signs. When a VS becomes very large, it obstructs the aqueduct causing collapse of the fourth ventricle leading to hydrocephalus and ultimately results in death if left untreated.

The rate of growth of a VS is unpredictable, but the average growth is 0.7 [15] to 1 mm/year [16], with 92% growing less than 2 mm/year [16]. In about 60% of patients, the annual tumor growth rate is <1 mm/year; in about 30%, 1–3 mm/year; and in about 10%, >3 mm/year [17]. Growth is manifest in 90% of tumors in the first 3 years after presentation [18], and if a VS growth is demonstrated by serial imaging, it usually continues to grow (63.9%). Only 30.6% arrest without treatment and 5.6% regress in size [19]. Intracanalicular tumors might grow slower than cisternal tumors [16], but this is not supported by all studies [17]. This might be related to the fact that growth rate is variable. In intracanalicular lesions, between 21.3 [20] and 70.6% [21] of patients demonstrate no tumor progression, but this variability signifies that potentially in as much as 76.6% of cases the tumor grows with resultant hearing loss [20].

Treatment for VS is still controversial. Treatment options consist of a wait and scan approach, stereotactic radiosurgery (SRS) (Gamma knife) and microsurgery [22]. A conservative approach can be elected in intracanalicular, asymptomatic, or elderly patients (>65 years) [14, 16, 17, 20, 21, 23, 24]. Furthermore, patients who fail conservative management have clinical outcomes that are not different from those who undergo primary treatment without a period of conservative management [16].

Microsurgery is elected for large and giant tumors, as large VS are more difficult to control with SRS and liable to produce ataxia due to transient expansion post-irradiation [25].

For small- and medium-sized VS, both microsurgery and SRS are valuable options. SRS, compared to surgical resection, seems to show superior outcomes for VS patients. A long-term tumor control rate of 94% after 5 years and 92.8% after 8 years [25] (transient facial palsy lower than 1%, and a probability of functional hearing preservation between 50 and 95%) can be achieved in experienced gamma knife centers treating large volumes of VS patients with state-of-the-art SRS [26]. Therefore, it has been suggested that unless a long-term follow-up examination indicates tumor progression at currently used radiation doses, SRS should be considered the best management strategy for the majority of VS patients [27, 28]. However, SRS does not arrest tumor growth in all patients: in 10% of patients who had SRS, tumor progression continues [29]. A second SRS treatment can often arrest further growth [29, 30], but if it does not, then radical surgery results in facial nerve worsening in half of the patients [31] due to severe adhesions or changes of the facial nerve [32]. Partial or subtotal resection can be proposed when SRS fails since the residual tumor does not seem to grow after subtotal resection [32]. A large study comparing 5,005 operations and 1,485 patients treated by SRS (gamma knife surgery) demonstrated that 96% of patients had total removal rates after microsurgical treatment, with a 1.8% recurrence rate, which compares favorably to recurrence rates after gamma knife surgery [33].

Based on these data, the following approach has been proposed: for lesions smaller than 2 cm, a conservative wait and scan approach; for growing lesions or lesions between 2 and 2.5 cm, either gamma knife or microsurgery; and for lesions bigger than 2.5 cm, microsurgery [22]. A combination of subtotal resection with adjunct SRS has been suggested as well to improve functional outcome in patients with VS that are larger than 4 cm [34, 35].

**Tinnitus and Vestibular Schwannoma**

Tinnitus is a very common symptom in VS: between 45 and 80% of VS patients have tinnitus [2, 9–11, 36–42]. It usually develops when the tumor is still
intracanalicular [11] and is reported as the principal presenting symptom in 1 of 10 patients who seek medical help [9].

When tinnitus is the main presenting symptom, it seems to be perceived as severe [37]. The presence of tinnitus in VS is unrelated to patient age, gender, audiometric thresholds between 2,000 and 4,000 Hz, ipsilateral auditory brainstem response abnormalities, or caloric test abnormalities [37]. VS are most commonly associated with high-frequency hearing loss; the tinnitus is usually high pitched and localized to the tumor ear [2]. There is a correlation between the presence of tinnitus and the type of hearing loss with a tendency for patients without hearing loss to be less likely to experience tinnitus [37]. Greater age at the time of diagnosis seems to be associated with greater severity of the tinnitus. Abnormal caloric responses are also associated with a greater severity of tinnitus [37].

The presence of tinnitus is related to the tumor size [37]: it is more common when the tumor is small or when it is larger than 4.5 cm. This suggests that the tinnitus might be related to two different pathophysiological mechanisms, similarly to what has been described for microvascular compressions (see Chap. 40). The rate of tinnitus in VS is higher in individuals with functional hearing than in deaf individuals [43]. Deafness does not mean relief from tinnitus, and tinnitus persists in 46% of individuals who were deaf before the operation for VS [43]. This may suggest that the tinnitus is initially caused by aberrant signal transmission in the auditory nerve [37], whereas at a later stage, the tinnitus results from deafferentation, similarly to the tinnitus that occurs in connection with microvascular compressions (Fig. 85.1).

Treatment Options for Tinnitus in Patients with Vestibular Schwannoma

Conservative Treatment

It is uncertain if treating a small tumor leaves the patient with a better chance of obtaining relief from future tinnitus by observing it without treatment [22]. Between one third [33] and one half [44] of patients who are followed conservatively develop progressive hearing loss, which could result in more tinnitus. However, it is not known whether or not this is the case.

Microsurgery

After VS microneurosurgery, tinnitus disappears in 0–45%, becomes better in 16–17% of patients, while 30–60% do not experience any change, and 8–29% become worse [45, 46]. Neither tumor size nor age at the time of the operation has an impact on the tinnitus that occurs after surgery [45]. Therefore, these factors cannot be used as predictors for who will improve, remain unchanged, or worsen postoperatively. Furthermore, there is no association between changes in the tinnitus and changes in the quality of life following surgical treatment of VS, suggesting that tinnitus may be of relatively minor importance in the overall quality of life of patients following microneurosurgery for VS [45]. If no tinnitus is perceived before surgery, almost 40–50% develop it afterwards [39, 47].

Tinnitus and VS size

Fig. 85.1 Tinnitus percentage related to tumor size, based on [37]
Retrosigmoid (Suboccipital) Resection

Tinnitus disappears in 25.2%, improves in 33.3%, remains unchanged in 31.6%, and worsens in 9.9% after tumor removal using a retrosigmoid approach [48]. Although the proportion of patients complaining of frequent tinnitus increases postoperatively, the number of patients who find the tinnitus troublesome decreases markedly [49]. There is no difference in tinnitus incidence whether the auditory nerve is resected or not during surgery [48]. If no tinnitus is present before surgery, it develops postoperatively in 8.5%, again, whether the auditory nerve was resected or not [48]. Risk factors for developing tinnitus after microsurgery are sudden drops in perioperative blood pressure and hearing preservation surgery. A sudden drop in blood pressure during or after surgery is a negative prognostic factor not only for hearing preservation but also for ipsilateral tinnitus in patients undergoing vestibular schwannoma surgery [50]. When attempting to achieve hearing preservation during removal of a VS in patients without preoperatively tinnitus, 85% develop tinnitus postoperatively compared to 31% of patients in operations where hearing preservation is not attempted in the translabyrinth resection group [36]. However, a number of smaller studies seem to have opposite results: 50% of patients who had tinnitus preoperatively complained of it postoperatively, and only 8% developed tinnitus as a result of VS surgery which saved hearing [40].

Translabyrinthine Resection

The tinnitus in individuals with VS is usually not bothersome [2, 37, 41, 42]. Fourteen percent suffer from moderate to severe tinnitus according to a tinnitus handicap inventory (THI), and postoperatively the tinnitus handicap is neither alleviated nor exacerbated by translabyrinthine surgery as a group. On an individual basis, the tinnitus handicap was worse in 6.5%, unchanged in 87%, and better in 6.5% [42]. There is a 35% risk for developing tinnitus when no preoperative tinnitus is present and a 15% chance that tinnitus disappears when present preoperatively with translabyrinthine surgery [41]. Patients who had a probable or definite nerve section had significantly lower postoperative tinnitus severity [51]. Based on these results, it may be suggested that patients with no tinnitus preoperatively undergoing this form of surgery are unlikely to develop tinnitus after the operation, and if they do it will not be severe enough to significantly affect their quality of life [52]. When asked what affects the postoperative quality of life, less than 4% of the patients mentioned tinnitus [53].

Gamma Knife Surgery (SRS)

Tinnitus only changes or develops in a few patients (4%) after SRS for VS [28, 54, 55]. Preoperative tinnitus does not seem to exacerbate on a group level, thus similar to microsurgery [56]. Also in patients with small VS and no hearing, results from SRS and microsurgery are not significantly different with regard to tinnitus [57]. Some studies, however, do show less tinnitus in about 45% of patients who were treated with SRS [55, 58]. Whether hearing preservation in SRS is related to this is unknown. The functional hearing preservation at 3 years is 80% in patients with tinnitus as a first symptom [59]. Based on a systematic review of hearing results, on average 57% hearing preservation rate can be expected [60, 61]. The probability of preserving functional hearing is not only dependent of the presence of tinnitus as a first symptom. Hearing preservation is higher in patients who have an initial symptom other than hearing decrease (91.1%), in patients younger than 50 years (83.7%), and in those treated with a dose to the cochlea of less than 4 Gy (90.9%) [62]. Three years after GKS for VS, 71% of patients have unchanged hearing levels [63].

Conclusion

Tinnitus is a common symptom in VS, but rarely distressing. In only 10% of patients, harboring a VS tinnitus is the principal symptom and in only 14% of these, the tinnitus is perceived as moderately to severely distressing. Using gamma knife (SRS) or translabyrinthine surgical removal does not alter the tinnitus at a group level, even though some individual patients may experience less tinnitus after the treatment. Retrosigmoid surgery seems to have an impact on the tinnitus, decreasing its severity, except when associated with surgery which saves hearing. This should be remembered when treating patients with VS presenting with tinnitus.
References

Chapter 86
Neuromodulation: Introduction

Berthold Langguth and Dirk De Ridder

Keypoints

1. Tinnitus is related to altered activity in the central nervous system.
2. Tinnitus can be treated by interfering with this abnormal activity in the central nervous system.
3. Potential therapeutic approaches include sensory modulation, sensory stimulation, brain modulation, and brain stimulation.
4. Pilot studies have shown first promising results for somatosensory stimulation, neurobiofeedback, transcranial direct current stimulation, repetitive transcranial magnetic stimulation, and epidural electrical stimulation.
5. All therapeutic strategies presented are still at an early stage of development.
6. More research is needed to further improve the efficacy of the presented techniques.

Keywords

Tinnitus • Brain stimulation • Nerve stimulation • Neural activity • Neural excitability • Neurobiofeedback • Transcranial magnetic stimulation • Electrical stimulation • Transcranial direct current stimulation • Nerve stimulation • Neuromodulation

Abbreviations

EEG Electroencephalogram
fMRI Functional magnetic resonance imaging
rTMS Repetitive transcranial magnetic stimulation
tDCS Transcranial direct current stimulation

As described in detail in Part I of this book, there is now compelling evidence that the phantom perception of sound is a consequence of altered activity in the central nervous system. An increasing number of animal studies and experimental neuroimaging studies in individuals with tinnitus together with experience from similar studies in patients who seek treatment for tinnitus have contributed to an increasingly detailed identification of the neural correlate of tinnitus. With the neural correlate of tinnitus, we mean the minimal neuronal mechanisms, which are jointly sufficient for the conscious percept [1] of tinnitus. In short, alterations of neural firing and oscillatory activity [2–4], alterations of neural synchrony and temporal coherence [5], and changes in the tonotopic maps of the auditory cortices [6] have been observed in connection with tinnitus and described in many studies (Chap. 10). Importantly, these changes are not restricted to one specific brain area. Rather, they can be conceived as alterations of a network involving auditory and non-auditory brain areas [7–9] (for more details see Chaps. 17, 20, and 21). These changes of neural activity seem to arise from dysfunctional activation of neural plasticity (Chap. 12) induced by altered sensory input, which is auditory deprivation in most cases (Chap. 11) [10, 11], but can also be altered somatosensory input (Chap. 9) [12]. Frontal and parietal brain areas seem to have an important modulatory role [7, 9, 13, 14].

Any causally oriented therapy should aim to normalize this disturbed neuronal network activity. In principle, there are two possibilities. The first approach consists of normalizing the disturbed auditory input to the auditory cortex, which can be done indirectly by hearing aids [15] or cochlear [16, 17], auditory
nerve [18], and brainstem implants [19], which all were shown to improve tinnitus in selected patients. If this approach is not possible, there is still the option to use other sensory pathways for influencing the disturbed networks (e.g., somatosensory stimulation [20, 21]) or a combination of different sensory modalities (e.g., virtual reality) [22]. Whereas the efficacy of somatosensory stimulation has been demonstrated in several studies for a subgroup of patients (see Chap. 91), virtual reality treatment is currently under investigation [22].

Another option is to supply the missing information directly to the auditory cortex [23] or interfere with the distributed “tinnitus network” directly. This can be done at the auditory cortex by implanted electrodes [24–26] (Chap. 90), by transcranial direct current stimulation (tDCS) (Chap. 89), and by repetitive transcranial magnetic stimulation (rTMS) (Chap. 88) [27–30]. Another option is to target other hubs [8] of the distributed “tinnitus network” [7, 9], e.g. both auditory and non-auditory areas [31]. Neurobiofeedback can also be used to modulate the abnormal tinnitus-related activity or metabolism by operant conditioning of the reward system [32]. Neurobiofeedback can be performed to normalize pathological oscillatory activity directly [EEG based (Chap. 87)] [33] or indirectly (fMRI based) [34]. Whereas stimulation by implanted electrodes or pharmacotherapy can be performed permanently, rTMS, tDCS, or neurofeedback can only be applied for a limited amount of time. Nevertheless, these methods hold therapeutic potential since all of them can induce plastic changes, which outlast the treatment period. These long-lasting effects with limited treatment time can be explained by either learning-like effects (activation of neural plasticity) or the disruption of dysfunctional networks, which then allow the re-establishment of a more physiological state.

All techniques presented in Chaps. 87, 88, 89, 90 and 91 have been only recently introduced as treatment options for tinnitus. All these approaches have been based on new insights in brain-based pathophysiology of tinnitus and have shown promising results in pilot studies. However, none of them have supplied enough evidence for a general application in routine treatment. Being at an early stage of development, an increase in efficacy can be expected for all these methods in the next years, especially with increasing knowledge of both the pathophysiology of the different forms of tinnitus and the neurobiological mechanisms involved in the modulatory effects of the different interventions being used already.

References

Chapter 87
Neurobiofeedback

Thomas Hartmann, Isabel Lorenz, and Nathan Weisz

Keypoints

1. While neurofeedback has been used for the treatment of various diseases for about 40 years, research on using it as a treatment against tinnitus has begun only recently.
2. This is mainly due to the fact that the first studies concerning electrophysiological abnormalities in tinnitus patients were done in the early 2000s.
3. This chapter first outlines the history of neurofeedback as well as the theory behind it.
4. This is followed by a short description of the electrophysiological abnormalities in tinnitus patients applied in the studies provided at the end of the chapter.
5. These studies not only show effects on electrophysiological measurements but also demonstrate a great impact on tinnitus sensation and distress.

Keywords
Chronic tinnitus • Neurofeedback • EEG

Abbreviations

ADHD Attention deficit and hyperactivity disorder
dB Decibel
EEG Electroencephalography
fMRI Functional magnetic resonance imaging
HL Hearing level
MEG Magnetoencephalography
QEEG Quantitative electroencephalography
TQ Tinnitus questionnaire

Theory and History

Bioneurofeedback (also known as neurofeedback, electroencephalography (EEG), biofeedback, or EEG operant conditioning) exploits a simple learning rule: the operant modification of signals acquired from the brain of a participant or patient. Although, advances in technology allow for more sophisticated forms of neurofeedback than was possible earlier, the basic principle has not changed over the past 40 years: A signal is acquired from the participant’s brain in the form of a recorded EEG, relevant aspects of this signal are extracted (e.g., power in a distinct frequency band), and fed back to the participant in real time. As soon as the signal reaches a predefined target, the participant is rewarded. It is important to note that this principle is agnostic to both the signal and the reward used. Furthermore, there are no assumptions about the direct behavioral relevance of the signal for the patient, as, for instance, there is no direct link between a certain group of cortical oscillations and a particular disorder the patient might suffer from. Moreover, changes of the respective signals normally do not have an immediate relevance. Thus, it is vital for every successful neurofeedback training approach to increase the behavioral relevance of the signal for the patient (e.g., by choosing an appropriate reward). A further aspect to be emphasized is that the participant cannot be aware of the acquired signal without the help of a feedback, which leads to the ultimate goal of any neurofeedback approach: learning via operant modification to control a signal, putatively reflecting a distinct brain state, which is normally beyond the individual’s awareness and thereby uncontrollable.

Following the seminal work of Miller [1], demonstrating that autonomic functions can be modified...
through operant conditioning, Sterman and Friar showed that not only it is possible to use operant conditioning to increase sensorimotor rhythms but also this modification leads to a decrease in the amount of seizures experienced by an epileptic patient [2]. Similar encouraging results were found 4 years later for attention-deficit hyperactive disorder (ADHD) [3], training enhancement in the alpha and reduction in the theta band. These patients successfully learned to control their EEG oscillations and modify them into the desired direction, and the ADHD symptoms improved on 13 behavioral categories like “Out-Of-Seat-Behavior” and “sustained attention”. A worsening of the symptoms was reported when the contingency of the training was reversed, resulting in a reward for decreasing alpha and increasing theta oscillations.

These studies present important results, but they were based on single cases. Controlled studies involving groups of patients as well as control groups and/or treatments were needed to confirm these results. One of the first controlled studies concerning epilepsy and neurofeedback was conducted in 1993 [4]. Twenty-five patients suffering from epilepsy learned to control their “Slow Cortical Potentials” (SCP) an event-related component indexing neuronal excitability. One year after the training, 13 of 18 patients reported a significant decrease in seizure incidence. In this study, patients not only learned to reduce the SCPs, leading to lower excitability and thereby preventing seizures. The protocol involved training both directions, thus teaching the patients to actually control this aspect of their brain waves in a more complete manner. Trying to achieve a transfer from laboratory experience to real-life situations, transfer trials and distraction were introduced. Transfer trials are trials in the same neurofeedback setting as used during the ‘real’ training, but without any feedback provided for the patient. Hence, the participants should be enabled to incorporate the strategy learned during the training and transfer it into their everyday routine. Distraction is used to further enhance the transfer to everyday life, as the patient is required to apply the strategy in situations outside the laboratory with some kind of distraction, such as background noise. Transfer trials and distraction ought to be considered as an important aspect in modern neurofeedback therapy.

Several controlled studies have now demonstrated promising effects of neurofeedback on epilepsy, ADHD, and other disorders such as depression (for a review see [5]).

Independent of the type of disorder, neurofeedback training always involves prior identification of an abnormal recordable signal pattern differentiating patients from healthy controls (e.g., a significant increase or decrease of power in distinct frequency bands). A further challenge is the demand for almost instantaneous feedback, excluding several signal-processing algorithms such as averaging data from numerous trials. Identification of abnormal signals can be achieved by either controlled studies or using QEEG (Quantitative EEG). QEEG is a method comparing EEG signals acquired and processed with a standardized setting to a normative database of either healthy individuals or patients exhibiting abnormal oscillatory activity due to a certain defined condition. Thus, significant deviations from the standard EEG-recordings can be found and used for neurofeedback trainings [5].

The design of neurofeedback training may be regarded to be independent from the disease or the signal to be trained. It always involves the acquisition of the signal using appropriate devices which is then further processed using either proprietary software bound to the specific equipment or freely available software like ConSole [6]. Although EEG signals are usually utilized for neurofeedback, today other signal sources, such as fMRI, are used as well. The software then reduces the information of the signal to an essential minimum which is then made visible and/or audible to the patient. A common example of such a neurofeedback cue is an object moving from the left to the right side on a computer screen, whereas the information of the signal is represented by the height of the symbol on the screen. If the participant in the study is able to reach a pre-defined target (e.g., to “move the symbol” above a certain height), he/she receives a reward, which can be positive visual feedback (e.g., smiley face) appearing on the screen or, in some cases, monetary compensation as well.

Treatment of subjective tinnitus by means of neurofeedback is a relatively new application. In the next section, we will give a short review of the identified abnormal spontaneous EEG patterns, followed by an overview of current neurofeedback approaches pursued in our laboratory.
Electrophysiological Correlates of Tinnitus

Using neurofeedback as a treatment for tinnitus, it is important to identify abnormal aspects of brain activity, which are correlated with measures of subjective tinnitus, such as its intensity and/or distress. This is fundamental, both for the conceptualization of a neurofeedback strategy and for the assessment of the training success. If behavioral measurements such as questionnaires assessing core symptoms of tinnitus are associated with the amount of modification gained throughout the training process, the argument that the abnormal EEG pattern is a critical clinical marker is strengthened.

During the last few years, several studies were published on electrophysiological correlates of tinnitus [7–9], stimulating the emergence of innovative theories and models as well as treatment approaches [10–14].

Although not directly related to neurofeedback, it is of value to consider current electrophysiological findings regarding possible and actual applications of neurofeedback. (For a more extensive review see Chap. 20.)

Results of Central Mechanisms of Tinnitus

In their 2004 review, Eggermont and Roberts provide an extensive overview on central mechanisms underlying tinnitus derived largely from animal studies [7]. Unlike studies involving humans, animal studies can directly explore the instantaneous effects of certain tinnitus-inducing treatments, like noise trauma and high doses of salicylate, in central and peripheral structures using single- and multi-unit recordings. Besides proving that subjective tinnitus is a central phenomenon, the review concludes that changes in neural activation related to peripheral changes cannot be isolated from the rest of the brain, but are likely to lead to changes in the balance of intracortical inhibition/excitation. This can lead to drastic changes of spike rate and temporal aspects of spiking activity (synchrony) in several areas of the brain, most notably in the auditory cortex. Particularly, changes in synchrony can be, if sufficiently large, captured using non-invasive techniques, such as EEG or MEG, and are reflected in alterations in ongoing oscillatory activity. In tinnitus, ongoing (spontaneous) synchronized activity probably engages the higher order brain regions that are responsible for conscious perception of tinnitus. Recently, this view has been further elaborated upon in a model framework by Weisz et al. [8] and extended by notions on inter-areal coupling of distant brain regions (see Chap. 20).

Basis for Neurofeedback Therapy of Tinnitus

Empirically, our neurofeedback tinnitus therapy was based on the identification of electrophysiological signals that differ markedly from people not experiencing chronic tinnitus. Our first paper on abnormal spontaneous brain activity was published in 2005 [9] not only demonstrating that tinnitus patients exhibit higher energy in the delta band and lower energy in the alpha band compared to healthy controls but also showing that a correlation exists between tinnitus distress and abnormal oscillatory activity patterns in right temporal and left frontal areas. These results were later supported and extended in a study revealing, furthermore, a marked increase of gamma band power in tinnitus patients [8]. Another paper shows a decrease in delta band power during residual inhibition [15].

The above-mentioned studies all point in the same direction: the resting state of brain oscillations is different in individuals with tinnitus and in individuals who do not have tinnitus in the delta, alpha, and gamma band (see Weisz et al. [8]). Thus, it seems reasonable to suggest that these EEG anomalies in individuals with tinnitus can be important for the therapy of tinnitus aimed at normalizing these oscillatory patterns. Cortical oscillations like respiratory rate or blood pressure are autonomous functions or a reflection of these. Therefore, operant modification of cortical oscillations should be possible by means of neurofeedback, as it has been demonstrated before regarding other aspects of electrophysiological signals.
Treating Tinnitus with Neurofeedback: An Overview Over Recent Studies

Studies exploring the effect of neurofeedback on subjective tinnitus are few. Two studies have supported the assumption that distress in general is associated with a reduction of power in the alpha band of EEG recorded from posterior sites and enhancement of power in the beta band [16, 17]. On the basis of these findings, it has been hypothesized that the vicious circle between strain, anxiety, and depression initiated in tinnitus can be interrupted through relaxation and by up-regulating the alpha activity (sign of increased relaxation) as well as down-regulating the beta activity (sign of decreased stress).

The approaches described in this chapter differ essentially from other studies in that the activity being modified is different in terms of assumed anatomical localization and generator types. While posterior recording sites have been the regions of interest in many studies, we focus on recordings from temporal and frontal regions, which we believe are mainly involved in the psychoacoustic and distress aspects of chronic tinnitus.

It is important that alpha oscillations in our approach are interpreted as an indicator of the excitatory–inhibitory balance in cortical neurons [8].

Here, we present the results of two recent studies by our workgroup in detail (Study 1 and 2) and pilot data from a new and innovative study (Study 3). Although all three studies differ in methodological details, the basic principles remain unchanged insofar as the objective of all training is to reestablish the excitatory–inhibitory imbalance putatively underlying tinnitus via a normalization of the ongoing spontaneous activity, particularly in the alpha band. The differences in the presented approaches lie mainly in which frequency bands are trained and how the feedback is presented to the patients.

Study 1

In the first study, 21 patients with chronic subjective tinnitus participated in a training aimed at controlling alpha power (5 patients), delta power (5 patients) or a ratio of alpha and delta power (11 patients). EEG was recorded at four fronto-central positions and the average power (in case of training a single frequency band) or ratio (in case of training alpha and delta simultaneously) of the respective frequency bands was displayed as the height of a fish “swimming” across the screen. No instructions on how to solve the task were given, except the notice that the position of the fish represented the cortical oscillations which had to be modulated by mental activity. Additionally, the participants were asked not to engage in muscular activities and to avoid eye blinks throughout the training session. Training success was monitored by matching the participant’s perception of their tinnitus to the intensity of their tinnitus to a 1-kHz test-tone using an audiometer and by measuring the power of the trained frequency bands during a 5-min resting condition before and after the training. The distress related to the tinnitus was surveyed once a week using a German adaptation of the Tinnitus Questionnaire [18]. Results showed a significant enhancement of the alpha–delta ratio within sessions and a significant linear trend between sessions. Thus, patients did not only learn to control their cortical EEG oscillations within a single session but also experienced an effect between sessions over the entire length of the training. Furthermore, a significant reduction of tinnitus intensity and tinnitus distress was revealed. The average tinnitus intensity was significantly reduced from 25 to 16 dB HL, and the average tinnitus distress measure decreased from 27 to 19 points at the end of the training. It is important to note that the amount of reduction of tinnitus intensity was strongly correlated with enhancements in the alpha/delta ratio, disregarding the exact training protocol. No significant differences were found between the different training groups (Alpha alone, Delta alone, Alpha/Delta ratio) or regarding tinnitus-related measures or ongoing oscillatory activity. This supports our notion that normalization of ongoing oscillatory activity might contribute to a reversal of the abnormal excitatory/inhibitory imbalance.

Although the study yielded promising results, it was not free of methodological problems. Thus, it is not clear and cannot be deducted post hoc what the patients actually trained as only the ratio of alpha/delta or one of the frequency bands was fed back. An increase of this ratio may have been an increase of alpha, a decrease of delta, or both, while a static ratio could have also been an increase in both bands or no change in these frequency bands at all. As the other two groups only trained one of the two frequency bands, no evidence about the effect of training both frequency bands could be concluded from the study. We thus developed a new training, providing two-dimensional feedback to the patients.
Study 2

Sixteen patients participated in the second study. EEG was recorded from 31 electrodes covering the whole scalp. The data were projected online on a source montage with eight sources covering major areas of the brain. Alpha and Delta power were computed for both temporal sources. During the training, patients saw a football (serving as the feedback cue) moving in the middle of the screen, which was supposed to be moved upwards, indicating increased alpha power, and sidewards (to the right-hand side), indicating decreasing delta power. A coordinate system was superimposed on the screen, dividing it into four quadrants, wherein the right upper quadrant was the patients’ target to reach (i.e., increased alpha power and decreased delta power).

Training success was, again, monitored using electrophysiological measurements as well as the tinnitus intensity matched to a 1-kHz test-tone using an audiometer and by measuring the power of the trained frequency bands during a 5-min resting condition before and after the training [18].

Patients were able to normalize their alpha and delta power significantly, which means there was a significant enhancement of alpha power and a significant reduction of delta power after the training. Behavioral measures also demonstrated a certain relief from the tinnitus. Thus, there was a significant decrease of TQ values from an average of 22 points before the start of the training to an average of 17 points after the last training session. Tinnitus intensity was also significantly reduced from an average of 26 dB to an average of 23 dB HL.

Although this study exhibits an alleviation of tinnitus symptoms in some patients, it is also clear that many patients were not able to learn the task, mainly due to the abstract nature of the task and insufficient instructions.

Study 3

We offered the participants one possible strategy (out of numerous others) to be successful in the neurofeedback task as described below. In contrast to the previous studies, an amplitude-modulated sound with a frequency spectrum close to the individual’s tinnitus was presented to both ears. Sound stimulation normally leads to desynchronization (decrease) of alpha oscillations recorded from auditory areas and is also modulated by top–down influence such as from attention [19]. By training a suppression of alpha desynchronization and thereby reducing cortical excitation, the aim was to aid patients in finding strategies of drawing away attention from their own internally generated sound.

Preliminary analyses of the results obtained from nine patients demonstrate highly significant effects regarding alpha normalization: Alpha power was increased by about 80% from the first to the last session. Behavioral measures point to an alleviation of tinnitus distress inasmuch as TQ values were significantly decreased from an average of 28 points to an average of 20 points.

Summary

Although neurofeedback has been available in clinical practice and research for 40 years, only recent advances in computer technology, amplifiers, and signal-processing routines made it possible to develop sophisticated techniques for biofeedback trainings. It is now possible to use knowledge about abnormal oscillatory patterns in the EEG that occurs in individuals with a disease to design a neurofeedback training program that is aimed at normalizing these patterns and thereby alleviating the disease condition.

Here, we have briefly reviewed the literature on abnormal cortical oscillations in individuals with tinnitus. Although such studies have been few and the results have not always been consistent, the central origin of tinnitus is now undisputed. Findings from our workgroup showed a decrease in alpha components of the EEG and an increase of delta and gamma activity in individuals who have tinnitus. On the basis of that, we designed and tested three kinds of neurofeedback trainings, which differed in methodological issues but shared the goal of normalizing these cortical oscillations.

In accordance with early neurofeedback studies, we showed that the participants were able to learn how to control the oscillations in their EEG. We could also show that this normalization had a positive impact on the perceived loudness of their tinnitus and/or the distress caused by their tinnitus.

Research on the cortical processes involved in the generation of tinnitus is new, and limited understanding
of the phenomenon involved is an obstacle in achieving success in therapy using biofeedback. Models and theories incorporating recent knowledge of the brain’s internal processes are evolving and will provide a better understanding of tinnitus as a central phenomenon. Together with new developments in techniques of signal processing and in neurofeedback, we may expect that innovative neurofeedback designs against tinnitus will be devised in the future. The recent findings suggest that training of coherences or connectivity between brain regions involved in the processing or generation of tinnitus will be promising areas in the future.

References

6. Hartmann T. ConSole [Internet]. Available from: http://console-kn.sf.net
**Chapter 88**

**Transcranial Magnetic Stimulation**

Tobias Kleinjung, Berthold Langguth, and Eman Khedr

**Keypoints**

1. Repetitive transcranial magnetic stimulation (rTMS) is a noninvasive method for applying electromagnetic fields to the brain.
2. rTMS can induce alterations of neuronal activity that outlast the stimulation period.
3. By modulating the excitability of the auditory cortex, rTMS can influence tinnitus perception.
4. Single sessions of rTMS over the temporal or temporoparietal cortex have been successful in transiently reducing tinnitus perception.
5. Repeated sessions of rTMS have resulted in tinnitus relief in a subgroup of patients lasting from several days to several months.
6. However, effect sizes of rTMS in the treatment of tinnitus are only moderate, and interindividual variability is high.
7. Further research is needed before this technique can be recommended for routine clinical use.

**Keywords** Tinnitus • Transcranial magnetic stimulation • Functional imaging • Cortical excitability • Neuromodulation

**Abbreviations**

DLFP Dorsolateral prefrontal cortex
EEG Electroencephalography
FDG [18F]deoxyglucose
MRI Magnetic resonance imaging
MT Motor threshold
PET Positron emission tomography
rTMS Repetitive transcranial magnetic stimulation
TMS Transcranial magnetic stimulation

**Transcranial Magnetic Stimulation**

Transcranial magnetic stimulation (TMS) is an experimental tool for stimulating neurons via brief magnetic pulses delivered by a coil placed on the scalp [1]. In brief, the stimulator delivers a short-lasting, high-intensity current pulse through an insulated stimulating coil. This induces a magnetic field perpendicular to the coil which penetrates the scalp and brain with little attenuation (see Fig. 88.1). The magnetic field reaches a maximum of approximately 1.5–2 T (same size as that of an MRI scanner) in about 100 μs and then decays back to zero [2]. The magnetic coils that are used have different shapes. Round coils are relatively powerful. Figure eight-shaped coils are more focal with a maximal current delivered at the intersection of the two round components [3] (see Fig. 88.2).

Because the field changes rapidly with time, it induces an electrical current in the brain under the coil which is similar in duration and amplitude to a conventional electrical stimulator used to activate peripheral nerves. Due to the strong decline of the magnetic field with increasing distance from the coil, the direct stimulation is limited to superficial cortical areas, but the stimulation effects can be propagated transsynaptically to functionally connected remote areas and thus indirectly affect large areas of the brain.
Whereas single magnetic pulses do not seem to have longer lasting effects on the brain, the application of multiple pulses, called repetitive TMS (rTMS), leads to effects on the brain that outlast the duration of the stimulation. These effects resemble those seen in animal experiments where repeated stimulation of many pathways has been shown to produce changes in the effectiveness of synapses in the same circuits [4]. Low-frequency (≤1 Hz) rTMS has been repeatedly shown to result in a decrease in cortical excitability [5], whereas high-frequency (5–20 Hz) rTMS results in an increase in excitability [6]. These changes include the phenomena of long-term potentiation (LTP) and long-term depression (LTD), which have been shown to be important in learning and memory [7]. Repetitive TMS can also be used to transiently disturb ongoing neural activity in the stimulated cortical area, thus creating a transient functional lesion. Such an approach can help to identify whether a given brain area is critically involved in a specific behavioral task.

Because of these unique and powerful features, rTMS has been widely used in various fields, including cognitive neuroscience and several clinical applications (for review see [8, 9]). However, despite its practical usefulness, the mechanisms of how rTMS stimulates neurons and interferes with neural functions are still incompletely understood.

**Rationale for the Application of TMS in Tinnitus**

Tinnitus is often associated with a lesion in the peripheral auditory system. It often occurs together with presbycusis, Ménière’s disease, noise trauma, sudden deafness, or drug-related ototoxicity [10, 11]. However, these pathologies are not directly causing tinnitus. Rather, the neuroplastic changes which occur in the brain as reaction to sensory deafferentation represent the neural correlate of most forms of tinnitus [12] (see Chaps. 10 and 12). Thus, the mechanisms involved in tinnitus generation share similarities with those responsible for phantom pain after limb amputation [11]. Support for these hypotheses comes from functional imaging studies demonstrating that tinnitus is associated with neuroplastic alterations in the central auditory system and associated areas (see Chap. 18). In detail, positron emission tomography (PET) investigations revealed abnormal asymmetry in the auditory cortices of tinnitus patients with higher levels of spontaneous neuronal activity on the left side, irrespective of tinnitus laterality [13–15]. However, changes of neural activity are not limited to the central auditory pathways. Temporoparietal regions, as well as frontal and limbic areas, are also involved [16–18].
Since rTMS has the ability to focally modulate cortical activity, it has been assumed that it can interfere with the abnormal neural activity in the auditory cortex associated with tinnitus and thereby influence the perception of tinnitus. If this is the case, repeated applications of rTMS might represent a potential treatment for some forms of tinnitus by producing longer lasting modulation of cortical activity. Additional support for this approach comes from clinical trials in which rTMS was used in an attempt to treat other pathological conditions with potential cortical hyperactivity, such as auditory hallucinations [19], writers’ cramp [20], and obsessive compulsive disorders [21].

Studies Using Single Sessions of rTMS in Tinnitus

Within the last few years, results of several studies using single sessions of rTMS have been published (for references see Table 88.1). The goal of these studies was to transiently reduce tinnitus perception (see Table 88.1). In these kinds of studies, trains of high-frequency rTMS (10–20 Hz) were mainly administered. In a pilot study, stimulation of the left temporoparietal cortex with high-frequency rTMS (10 Hz) resulted in a transient reduction of tinnitus in 57% of the participants [22]. This result has been confirmed in a large series of 114 patients with unilateral tinnitus [23]. In this study, repetitive TMS at frequencies between 1 and 20 Hz was applied over the auditory cortex contralateral to the site of tinnitus perception. The best tinnitus suppression was achieved using higher stimulation frequencies, and patients who had their tinnitus for a shorter duration had the best results. These studies indicate that rTMS can be a valuable diagnostic tool for differentiating different forms of chronic tinnitus. This approach has been used for screening purposes to select patients for surgical implantation of cortical electrodes [24, 25] (see Chap. 90).

Two studies [26, 27] confirmed the result of transient tinnitus reduction after high-frequency stimulation of the left temporoparietal cortex, whereas one study [28] demonstrated reliable tinnitus suppression in only 1 out of 13 subjects after a single session of high-frequency rTMS. Additionally, one small study has shown [26] that the participants with significant tinnitus reduction after rTMS also had good response to anodal transcranial direct current stimulation (tDCS).

Different methods have been used to identify the target for stimulation. In one study, changes of cerebral blood flow were determined before and after lidocaine injection [17]. Single sessions of low-frequency (1 Hz) rTMS with the coil navigated to individually determined areas in the temporoparietal cortex resulted in tinnitus reduction in 6 out of 8 participants lasting up to 30 min.

Studies Using Repeated Sessions of rTMS in Tinnitus

The application of low-frequency rTMS in repeated sessions followed the hypothesis that longer lasting
Table 88.1  Effects of single sessions of rTMS on tinnitus

<table>
<thead>
<tr>
<th>Authors</th>
<th>N</th>
<th>Stimulation site</th>
<th>Coil positioning</th>
<th>Frequency</th>
<th>Intensity</th>
<th>Pulses/session</th>
<th>Control condition</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plewnia et al. [22]</td>
<td>14</td>
<td>Various scalp positions</td>
<td>10–20 EEG system</td>
<td>10 Hz</td>
<td>120% MT</td>
<td>30</td>
<td>Stimulation of nonauditory cortical areas</td>
<td>In 8 patients (58%) tinnitus suppression after left temporal/temporoparietal stimulation</td>
</tr>
<tr>
<td>De Ridder et al. [23]</td>
<td>114</td>
<td>Auditory cortex contralateral to tinnitus site</td>
<td>Anatomical landmarks</td>
<td>1, 5, 10, 20 Hz</td>
<td>90% MT</td>
<td>200</td>
<td>Coil angulation</td>
<td>In 60 patients (53%) good or partial tinnitus suppression after active rTMS, in 33% suppression after sham rTMS</td>
</tr>
<tr>
<td>Fregni et al. [26]</td>
<td>7</td>
<td>Left temporoparietal areas</td>
<td>10–20 EEG system</td>
<td>10 Hz</td>
<td>120% MT</td>
<td>30</td>
<td>Sham coil and active stimulation of mesial parietal cortex</td>
<td>In 3 patients (42%) tinnitus suppression after left temporoparietal stimulation, no effect for both control rTMS conditions</td>
</tr>
<tr>
<td>Folmer et al. [27]</td>
<td>15</td>
<td>Left and right temporal cortex</td>
<td>10–20 EEG system</td>
<td>10 Hz</td>
<td>100% MT</td>
<td>150</td>
<td>Sham coil</td>
<td>In 6 patients (40%) tinnitus suppression after active rTMS, in four of the patients after contralateral rTMS in two patient after ipsilat. TMS; in 2 patients suppression after sham rTMS</td>
</tr>
<tr>
<td>Londero et al. [28]</td>
<td>13</td>
<td>Contralateral auditory cortex</td>
<td>fMRI guided neuronavigation</td>
<td>1, 10 Hz</td>
<td>120% MT</td>
<td>30</td>
<td>Stimulation over nonauditory cortical areas</td>
<td>Eight patients were stimulated over the auditory cortex with 1 Hz; in 5 of them (62.5%) tinnitus suppression; no suppression after 1 Hz rTMS of nonauditory targets; no suppression after 10 Hz, in 2 patients suppression after stimulation of a control position</td>
</tr>
<tr>
<td>Plewnia et al. [17]</td>
<td>8</td>
<td>Area of maximum tinnitus related PET activation (temporoparietal cortex)</td>
<td>Neuronavigational system, based on H2O PET with and without Lidocaine</td>
<td>1 Hz</td>
<td>120% MT</td>
<td>300, 900, 1,800</td>
<td>Control position (occipital cortex)</td>
<td>In 6 patients (75%) tinnitus reduction after active rTMS, better suppression with more pulses</td>
</tr>
<tr>
<td>Author(s)</td>
<td>N</td>
<td>Stimulation site</td>
<td>Coil positioning</td>
<td>Frequency</td>
<td>Intensity</td>
<td>Pulses/session</td>
<td>Control condition</td>
<td>Results</td>
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<tr>
<td>De Ridder et al. [47, 48]</td>
<td>46</td>
<td>Auditory cortex contralateral to tinnitus site</td>
<td>Anatomical landmarks</td>
<td>5, 10, 20 Hz tonic; 5, 10, 20 Hz burst</td>
<td>90%MT</td>
<td>200</td>
<td>No placebo condition</td>
<td>14 placebo-negative patients were analyzed: In those with narrow band/white noise tinnitus burst TMS was more effective in tinnitus suppression as compared to tonic TMS, whereas for pure tone tinnitus no difference was found between burst and tonic stimulation</td>
</tr>
<tr>
<td>Poreisz et al. [46]</td>
<td>20</td>
<td>Inferior temporal cortex</td>
<td>10–20 EEG electrode system, T3</td>
<td>Continuous theta burst, intermittent theta burst, immediate theta burst</td>
<td>80%MT</td>
<td>600</td>
<td>No placebo condition</td>
<td>Significant tinnitus reduction only for continuous theta burst immediately after stimulation</td>
</tr>
<tr>
<td>Meeus et al. [51]</td>
<td>50</td>
<td>Auditory cortex contralateral to tinnitus site</td>
<td>Anatomical landmarks</td>
<td>1,5, 10, 20 Hz tonic; 5, 10, 20 Hz burst</td>
<td>50% maximal stimulator output (independently of individual MT)</td>
<td>200</td>
<td>No difference between tonic and burst rTMS in pure tone tinnitus (about 50% average suppression in unilateral and 30% in bilateral tinnitus). For bilateral narrow band tinnitus superiority of burst stimulation compared to tonic stimulation; better effects in patients with lower MT</td>
<td></td>
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</table>
improvement of tinnitus complaints can be achieved by reducing auditory cortex hyperactivity. An increasing number of studies using this approach as a treatment for tinnitus have been published recently (Table 88.2). Most rTMS treatment studies applied low-frequency rTMS in long trains of 1,200–2,000 pulses repeatedly over 5–10 days. In all controlled studies, a statistically significant improvement of tinnitus complaints has been documented. However, the degree of improvement and its duration varied across studies, probably due to differences in study design, stimulation parameters, and selection criteria of the participants.

Repetitive TMS has been applied over temporal or temporoparietal cortical areas. One placebo-controlled study with 14 participants used [18F]deoxyglucose (FDG) PET and a neuronavigational system for the exact positioning of the TMS coil over the site of maximum activation in the auditory cortex [14] (see Fig. 88.3). After active treatment, the participants experienced a significant decrease in their tinnitus, as reflected by the score of the tinnitus questionnaire, whereas sham treatment showed no effect. Treatment effects were still detectable 6 months after treatment. Another study concerned the effects of 2 weeks of rTMS applied over the cortical area where lidocaine-induced activity change was largest as determined by [15O]H2O PET [29]. This approach also resulted in moderate, but significant effects after active stimulation. Placing the coil over the left temporal area according to the 10–20 EEG coordinate systems [30] resulted in a significant reduction of tinnitus severity after 10 sessions of 1 Hz rTMS. Beneficial effects of low-frequency rTMS have been confirmed by several further controlled studies [31–33].

While some studies demonstrated effects that outlasted the stimulation period for as much as 12 months [14, 34, 35], others were not able to achieve long-lasting effects [29, 32]. The number of daily sessions may be an important factor regarding long-term effects in tinnitus patients [36], thus similar to the experience from TMS applications for other disorders such as depression [37] and auditory hallucinations [38].

A recent case report showed that rTMS may be used as a maintenance treatment to manage chronic tinnitus [39]. In this patient, tinnitus could be reduced by rTMS each time it reoccurred using one to three sessions of rTMS; it finally remained stable on a low level after the third stimulation series. The positive effect of this maintenance stimulation could also be confirmed by reduced cerebral metabolism in PET imaging after treatment. The approach to use rTMS for maintenance treatment of tinnitus is further supported by the observation that those patients, who respond once to rTMS treatment, also experience further positive effects from a second series of rTMS [40].

Enhancement Strategies

In previous studies, when repeated sessions of rTMS were introduced as a therapeutic approach, stimulation has been performed at a frequency of 1 Hz [41]. This was motivated by the finding that 1 Hz rTMS reduces neuronal excitability over the motor cortex [5] and by the successful use of low-frequency rTMS in treatment of neuropsychiatric disorders, which are associated with focal hyperexcitability [42]. This concept has been challenged by a recent study with a relatively large sample size which compared effects of 1, 10, and 25 Hz rTMS [34]. Whereas sham rTMS treatment had no effect, active stimulation over the left temporoparietal cortex resulted in a reduction of tinnitus irrespective of the stimulation frequency. A follow-up assessment 1 year after treatment suggested a trend for higher efficiency of stimulation at 10 and 25 Hz, as compared to 1 Hz [35].

Experimental data from motor cortex stimulation in healthy subjects indicate that the effect of low-frequency rTMS can be enhanced by high-frequency priming stimulation [43]. However, in a clinical study, high-frequency priming stimulation failed to enhance the therapeutic efficacy of low-frequency rTMS for the treatment of tinnitus [44].

Repetitive TMS can be applied in a tonic and a burst mode. The burst stimulation technique has been proposed for enhancing rTMS effects. In detail, bursts of three stimuli at a frequency of 50 Hz (interval of 20 ms between each stimulus), applied every 200 ms (5 Hz, Theta burst), have been shown to induce more pronounced and longer lasting effects on human motor cortex than tonic stimulation [45]. Single sessions of continuous theta burst stimulation (3 pulses at 50 Hz, repeated at 200 ms intervals for up to 600 pulses for 40 s continuously) over the temporal cortex in tinnitus patients did only result in short-lasting reduction of tinnitus loudness, comparable to effects achieved with single sessions of tonic stimulation, whereas other
| Authors                | N  | Stimulation site                                                                 | Coil positioning                                                                 | Frequency (Hz) | Intensity | sessions | Pulses/session | Design                  | Control condition | Results                                                                                     |
|------------------------|----|----------------------------------------------------------------------------------|----------------------------------------------------------------------------------|----------------|-----------|-----------|-----------------|-----------------|--------------------------------------------------------------------------------------------|
| Kleinjung et al. [14]  | 14 | Area of maximum PET activation in the temporal cortex, (12 left, 2 right)       | Neuronavigational system, based on FDG-PET                                        | 1              | 110% MT  | 5         | 2,000           | Sham-controlled, cross-over | Sham coil        | Significant reduction of tinnitus after active rTMS as compared to sham rTMS; lasting tinnitus reduction (6 months) |
| Langguth et al. [30]   | 28 | Left auditory cortex                                                             | 10–20 EEG system                                                                  | 1              | 110% MT  | 10        | 2,000           | Open            | No control condition                                                         | Significant reduction of tinnitus until end of follow-up (3 months) |
| Plewnia et al. [29]    | 6  | Area of maximum tinnitus related PET activation (temporoparietal cortex; 3 left, 3 right) | Neuronavigational system, based on H$_2$O PET with and without Lidocaine          | 1              | 120% MT  | 10        | 1,800           | Sham-controlled, cross-over | Occipital cortex | Significant reduction of tinnitus after active rTMS, as compared to the control condition; no lasting effects |
| Kleinjung et al. [56]  | 45 | Left auditory cortex                                                             | Neuronavigational system, based on structural MRI                                 | 1              | 110% MT  | 10        | 2,000           | Open            | No control condition                                                         | Significant tinnitus reduction after rTMS, lasting up during follow-up period (3 months); responders were characterized by shorter tinnitus duration and less hearing impairment |

(continued)
<table>
<thead>
<tr>
<th>Authors</th>
<th>N</th>
<th>Stimulation site</th>
<th>Coil positioning</th>
<th>Frequency (Hz)</th>
<th>Intensity</th>
<th>sessions</th>
<th>Pulses/session</th>
<th>Design</th>
<th>Control condition</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rossi et al.</td>
<td>16</td>
<td>Left secondary auditory cortex</td>
<td>Eight patients: neuronavigational systemEight patients: according to 10–20 EEG system, halfway between T3 and C3/T5</td>
<td>1</td>
<td>120% MT</td>
<td>5</td>
<td>1,200</td>
<td>Sham-controlled, cross-over</td>
<td>Coil angulation + electrical stimulation of facial nerve</td>
<td>Significant reduction of tinnitus after active rTMS, as compared to the control condition, no lasting effects</td>
</tr>
<tr>
<td>Smith et al.</td>
<td>4</td>
<td>Area of maximal PET activation in the temporal cortex, neuronavigational system</td>
<td>Neuronavigational system, based on FDG-PET</td>
<td>1</td>
<td>110% MT</td>
<td>5</td>
<td>1,800</td>
<td>Sham-controlled, cross-over</td>
<td>Coil angulation</td>
<td>Modest response to active treatment in 3 patients (75%)</td>
</tr>
<tr>
<td>Khedr et al.</td>
<td>66</td>
<td>Left temporoparietal cortex</td>
<td>10–20 EEG system</td>
<td>1, 10, 25</td>
<td>100% MT</td>
<td>10</td>
<td>1,500</td>
<td>Sham-controlled, parallel group design</td>
<td>Occipital cortex</td>
<td>Significant reduction of tinnitus after all three active rTMS conditions, as compared to the control condition; tinnitus reduction lasting during follow-up period (4 months and 12 months)</td>
</tr>
<tr>
<td>Langguth et al.</td>
<td>32</td>
<td>Left auditory cortex, Neuronavigational system, based on structural MRI</td>
<td>1; 6 + 6</td>
<td>110% MT (90% MT for 6 Hz rTMS)</td>
<td>10</td>
<td></td>
<td>2,000</td>
<td>Randomization between two active treatment conditions, parallel group design</td>
<td>No sham control condition</td>
<td>Significant improvement for both stimulation conditions, no difference between conditions, no lasting effects</td>
</tr>
<tr>
<td>Authors</td>
<td>Stimulation site</td>
<td>coil positioning</td>
<td>Frequency (Hz)</td>
<td>Intensity (%) MT</td>
<td>Sessions</td>
<td>Design</td>
<td>Control condition</td>
<td>Results</td>
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<tr>
<td>Lee et al. [65]</td>
<td>Left temporoparietal cortex</td>
<td>??</td>
<td>0.5</td>
<td>100% MT</td>
<td>5</td>
<td>Open study</td>
<td>No control condition</td>
<td>No significant reduction of tinnitus</td>
<td></td>
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<tr>
<td>Kleinjung et al. [53]</td>
<td>Left auditory cortex; left dorsolateral prefrontal cortex</td>
<td>neuronavigational system, based on structural MRI</td>
<td>1, 20 (DLPFC) + 1</td>
<td>110% MT</td>
<td>10</td>
<td>Two active treatment conditions, parallel group design</td>
<td>No sham control condition</td>
<td>Directly after stimulation significant improvement for both stimulation conditions, at 3 month follow-up significantly better results for the combined frontal and temporal stimulation</td>
<td></td>
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<tr>
<td>Kleinjung et al. [55]</td>
<td>Left auditory cortex</td>
<td>Neuronavigational system, based on structural MRI</td>
<td>1; 1 + Levodopa</td>
<td>110% MT</td>
<td>10</td>
<td>Randomization between two active treatment conditions, parallel group design</td>
<td>No sham control condition</td>
<td>Significant improvement for both stimulation conditions, no difference between conditions, no lasting effects</td>
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<tr>
<td>Marcondes et al. [33]</td>
<td>Left temporoparietal cortex</td>
<td>10–20 EEG system</td>
<td>1</td>
<td>110% MT</td>
<td>5</td>
<td>Sham controlled, parallel group design</td>
<td>Sham coil</td>
<td>Significant improvement after active rTMS but not after sham rTMS, beneficial treatment effects still detectable at 6 months follow-up</td>
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</table>

MT = motor threshold
theta burst protocols had no effect at all [46]. In two other studies, single sessions of burst stimulation were compared with tonic stimulation [47, 48]. Burst stimulation had similar effects as tonic stimulation in patients with pure-tone tinnitus but was superior in patients with noise-like tinnitus. A possible explanation for this finding is that pure-tone tinnitus may be due to increased neuronal activity in the classical (lemniscal) auditory pathways, which mainly fire tonically, whereas noise-like tinnitus may be the result of increased activity in the nonclassical (extralemniscal) pathways, characterized by burst firing [47, 49, 50]. A follow-up study of the same group could replicate this result for bilateral tinnitus, but not for unilateral tinnitus [51]. Furthermore, this study suggests that higher stimulation intensity may result in slightly better tinnitus suppression.

The neurobiology of chronic tinnitus suggests that neuronal changes are not limited to the auditory pathways [16]. Recent progress in neuroscientific research demonstrated that hyperactivity within primary sensory areas alone is not sufficient for conscious tinnitus perception. Rather, synchronized co-activation of frontal and parietal areas seems to be necessary [52]. In one pilot study, 32 patients received either low-frequency temporal rTMS or a combination of high-frequency prefrontal and low-frequency temporal rTMS [53]. Directly after therapy, there was an improvement of the tinnitus questionnaire score for both groups, but there were no differences between groups. Evaluation after 3 months revealed a remarkable advantage for combined prefrontal and temporal rTMS treatment. These data indicate that modulation of both frontal and temporal cortex activity might represent a promising enhancement strategy for improving TMS effects in tinnitus patients.

Combination of rTMS with pharmacologic intervention has been suggested for potentiating of rTMS effects. It is known from animal experiments that neuronal plasticity can be enhanced by dopaminergic receptor activation [54]. However, in a clinical pilot study, the administration of 100 mg of levodopa before rTMS was not successful in enhancing rTMS effects in tinnitus patients [55].

There is some evidence from several studies that the histories of patients who are treated may affect the therapeutic outcome of rTMS in tinnitus patients. Several studies reported that patients who have had their tinnitus for a short duration had better treatment outcomes [14, 23, 34, 56]. Normal hearing was also identified as a positive clinical predictor for good treatment response [33, 56]. Interestingly, short tinnitus duration and normal hearing have been demonstrated to be positive predictors in other treatment options for tinnitus as well [57, 58].
Methodological Considerations

Tinnitus is a phantom perception of sound and is susceptible for placebo effects. Evaluation of treatment efficacy requires adequate methodology for control of unspecific effects related or unrelated to the treatment (see Chap. 22). The majority of controlled studies, published so far, have compared the effect of the active treatment with placebo treatment in cross-over designs. Potential shortcomings of this approach include carry-over effects and missed long-term effects due to short observation periods. Also, different kinds of sham treatments have been reported so far. Besides the sham coil system [14, 59], which mimics the sound of the active coil without generating a magnetic field, an angulation of an active coil tilted 45° [32] or 90° [31] to the skull surface or a stimulation of nonauditory brain areas [29] have been described. Finding an optimal control condition for treatment studies is also difficult because of limitations in blinding of patients and operators to different stimulus conditions and due to the fact that TMS itself results in auditory and somatosensory stimulation in addition to the anticipated specific effect. One possible solution is a control condition which involves electrical stimulation of the facial nerve [31].

In most studies, validated tinnitus questionnaires and visual analog scales serve as primary outcome measurements due to the lack of objective signs of tinnitus. One pilot study demonstrated that an improvement in tinnitus rating after stimulation was also reflected by a reduction of activity in the PET scan after rTMS therapy, as compared to pretreatment values [32]. Therefore, functional imaging might be suggested as an important objective marker of treatment effects.

Safety Aspects

An extensive body of data has confirmed that rTMS is a safe and well-tolerated technique [60] when performed within a range of parameters defined according to a consensus on safety guidelines [60, 61]. Most available data regarding safety originate from rTMS studies in depressed subjects. A study showed that 2–4 weeks of daily prefrontal rTMS resulted in no sign of structural MRI changes [62], no significant changes in auditory thresholds, and no significant electroencephalogram abnormalities [63]. Adverse auditory effects such as hearing loss or auditory hallucinations have not been reported after temporal rTMS. The risk of rTMS induced epileptic seizures, which had been reported in individual cases after high-intensity and high-frequency rTMS, has been largely reduced since the introduction of safety guidelines [61]. Mild adverse effects such as physical discomfort on the skull during stimulation or transient headache after stimulation are reported by about 10% of treated patients. It is essential that contraindications such as electronic implants (e.g., cardiac pace makers and cochlea implants), intracranial pieces of metal, or previous epileptic seizures are considered before treatment with rTMS.

Conclusion

In summary, the results from an increasing number of studies must be considered as preliminary due to small sample sizes, methodological heterogeneity, and high variability of results. However, the results of these studies are promising and show a similar percentage of beneficial effects. Data on the effect of the duration of treatment are still controversial. Effects outlasted the stimulation period up to 12 months in some studies; others could not demonstrate any after-effects. Replication in multicenter trials with many patients and long-term follow-up are needed before further conclusions can be drawn [64]. Further clinical research is also needed to get a clear definition of which subgroups of patients with tinnitus benefit most from rTMS and how their medical histories affect the outcome. Better understanding of the pathophysiology of the different forms of tinnitus and the neurobiological effects of rTMS will be critical for optimizing or even individualizing treatment protocols.

References


Chapter 89
Transcranial Direct Current Stimulation (tDCS): A New Tool for the Treatment of Tinnitus?

Sven Vanneste and Dirk De Ridder

Keypoints

1. Transcranial direct current stimulation (tDCS) is a non-invasive technique of cortical stimulation encompassing a relatively weak constant current flow (between 0, 5 and 2 mA) through the cerebral cortex via scalp electrodes.
2. Several studies already revealed that tDCS can influence working memory, decision making, risk-taking behavior, impulsiveness, and emotions responsive to visual material in healthy humans.
3. Major depression and tinnitus have also shown promise in few pilot studies.
4. This chapter will review tDCS and its potential as treatment for tinnitus.

Keywords Tinnitus • Transcranial direct current stimulation • Transcranial magnetic stimulation • tDCS • TMS • Neuroplasticity

Transcranial Direct Current Stimulation

Transcranial Direct Current stimulation (tDCS) is a non-invasive procedure of cortical stimulation that was introduced in the 1960s. In these early studies, it was shown that subthreshold direct current stimulation increases spontaneous neuronal activity in the brain [1–3]. Apart from changes of spontaneous discharge rates, direct current stimulation of levels below behavioral threshold was shown to modulate the cortical response to thalamic stimulation in animals [3, 4]. It was further demonstrated that in addition to an acute effect of direct current stimulation, this technique can also induce a long-lasting after-effect on neuronal excitability and activity [1, 3, 5, 6]. Although these findings on direct current stimulation were obtained in animal studies, most of the results can probably be applied to humans. Nevertheless, it took almost 40 years before direct current stimulation gained attention as a possible tool for patient treatment and research in humans.

When tDCS is applied in humans, a relatively weak constant current (between 0, 5 and 2 mA) is passed through the cerebral cortex via scalp electrodes. Depending on the polarity of the stimulation, tDCS can increase or decrease cortical excitability in the brain regions to which it is applied [7]. Currently, tDCS is usually applied through two surface electrodes, one serving as the anode and the other as the cathode. Some of the applied current is shunted through scalp tissue and only a part of the applied current passes through the brain. Anodal tDCS typically has an excitatory effect on the local cerebral cortex by depolarizing neurons, while the opposite is the case under the cathode where hyperpolarization occurs. This effect of tDCS typically outlasts the stimulation by an hour or longer after a single treatment session of sufficiently long stimulation duration [8–11].

Blocking voltage-dependent ion channels pharmacologically abolishes any effect of depolarizing anodal tDCS on cortical excitability and does not influence the impact of hyperpolarizing cathodal tDCS. This means that the effect of tDCS on the cerebral cortex might be a subthreshold modulation of neuronal resting membrane potential [12], but this cannot explain the after-effects of tDCS. The NMDA receptor antagonist dextrometorphan blocks the after-effect of tDCS, whereas the NMDA receptor agonist D-cycloserine
partially extends this effect [13–15]. This means that the after-effects of tDCS might depend on a modification of NMDA receptors efficacy. This tDCS polarity-dependent alteration of NMDA receptor function seems to be initiated by the respective membrane potential shift and probably by the accompanying cortical activity modification, because sodium channel blocker carbamazepine eliminates both the immediate and after-effect. Intraneuronal calcium concentration also contributes, since calcium channel antagonists eliminate the excitability-enhancing after-effect of anodal tDCS [12]. The mechanism of action of tDCS thus appears to depend on NMDA receptor activity [16] and involve a combination of hyper- and de-polarizing effect on neuronal axons.

Increasing the size of the reference electrode and reducing the size of the stimulation electrode allow for more focal treatment effects [17]. Moving the electrodes a few centimeters shifts the efficacy of tDCS dramatically [18]. Moreover, the electrical field strength is relatively homogeneous under the electrodes, but diminishes exponentially away from the electrode [7, 19]. In contrast to the relatively focal electrophysiological effect under the electrodes, widespread remote effects of tDCS on different cortical and subcortical areas were revealed in a PET study, suggesting that these effects might be caused by neural connection and not by direct electrical stimulation [20].

Safety of tDCS in Humans

To date, more than 2,000–3,000 individuals have participated in tDCS studies with no significant adverse effects reported using standard protocols consisting of 1–2 mA intensity, electrode size between 25 and 35 cm², and stimulation between 20 and 30° min per session. Slight tingling under the electrodes, headache, fatigue, and nausea might occur [21] and high current amplitudes (2 mA) can induce burns under the electrodes (Frank et al. 2010; Palm et al. 2008). tDCS does not elevate serum neuron-specific enolase levels (i.e. a sensitive marker of neuronal damage) [9]. No brain edema, no alterations of the blood–brain barrier, and no cerebral tissue damage were detectable by magnetic resonance imaging after tDCS [13]. Also, no worsening of cognitive function has been observed as a consequence of treatment [22]. However, for safety reasons, electrodes should not be positioned above cranial foraminae and fissures because these could increase effective current density in neural tissue.

TMS vs. tDCS

As a method for neuromodulation, tDCS has been compared with repetitive transcranial magnetic stimulation (rTMS) [23, 24]. As in rTMS, the effect produced by tDCS depends on the stimulation duration, the stimulation intensity, and the location of stimulating electrodes. While both techniques allow focal neuromodulation, there are fundamental differences between these methods; whereas TMS is thought to exert its effects by inducing action potentials in cortical neurons, tDCS is believed to modulate neuronal excitability without inducing neuronal firing.

Moreover, the two methods differ in several practical aspects, and tDCS has several advantages over rTMS. Since tDCS produces less artifacts, such as acoustic noise and muscle twitching, it is more suitable for double-blind, sham-controlled studies and clinical applications of tinnitus research. The equipment for tDCS is compact and portable and less expensive. Seizure incidents have not been reported in tDCS studies, and the effects of tDCS seem to last longer than that of rTMS, which makes it more suitable as a treatment tool. The use of tDCS should therefore be considered as a complementary tool to rTMS.

Rationale for Using tDCS in the Treatment of Tinnitus

Considerable evidence that neural plasticity can cause tinnitus has been recently presented [25, 26]. Neural plasticity is a property of the nervous system to change its function and its organization [27], change synaptic efficacy, generate or reduce synapses, and produce new connections by developing and eliminating axons and dendrites (see Chap. 12). For tinnitus in particular, a reorganization of the auditory cortex consisting of a shift in tonotopic maps contralateral to the tinnitus side has been demonstrated [28]. Abnormal symmetry in the auditory cortex activity in tinnitus has been shown, indicating a higher level of spontaneous activity [29–32].

Other studies revealed changes in non-auditory brain areas, namely in frontal and limbic areas [33–36]. In the
subcortical anterior cingulate area, including the nucleus accumbens, a reduction of gray matter density has been shown in tinnitus as compared to controls [37]. MEG studies have found a reduction in alpha (8–12 Hz) and an increase in delta (1.5–4 Hz) in temporal regions, left frontal, and right parietal areas [38] as well as increased functional connectivity between the right frontal lobe and anterior cingulum [39].

Since tDCS has the ability to modulate cortical neural activity, it seems likely that application of tDCS to specific regions could alter tinnitus. Several studies already revealed that tDCS can have beneficial effects on disorders such as depression and pain. Preliminary results suggest that tDCS applied to the temporal lobe [24] and the dorsal lateral prefrontal cortex can suppress tinnitus [58].

tDCS of the Temporal Lobe

Neuroimaging and electrophysiological studies have shown high spontaneous activity in the central auditory nervous system and changes in the tonotopic map of the auditory cortex in some individuals with tinnitus [33, 40–42]. Based on these findings, Fregni et al. initiated a tDCS study on a small sample (N=7) with bilateral non-pulsatile tinnitus (tinnitus duration range: 1–17 years) in an attempt to modulate neural activity in the left temporoparietal cortex [24]. A single session of anodal tDCS applied over the left temporoparietal area and cathode placed contralateral over the supraorbital area resulted in a transient reduction of tinnitus, similar to what has been shown to occur after applying TMS at 10 Hz [24]. No effect was found from a single session of cathodal tDCS applied over the left temporoparietal area with the anode placed over the contralateral supraorbital area. This was surprising, since it was cathodal, tDCS has a general inhibitory effect. One possible reason might be that cathodal tDCS was too weak to disrupt ongoing activity. Anodal tDCS, however, had a transient suppressive effect on the tinnitus of the participants in this study. As large electrodes were used, it was assumed that anodal tDCS would additionally excite surrounding areas that might, by competition or inhibitory connections, decrease the pathologically increased activity of some areas related to tinnitus pathophysiology. Yet, an alternative explanation might be that anodal tDCS affects targeted not only the brain region but also distant cortical and subcortical structures because these regions are connected to the areas that were stimulated. This would be in accordance with a recent PET study that demonstrated anodal tDCS of the motor cortex compared with cathodal tDCS induced a more widespread increase of regional cerebral blood flow [20].

The results from this study thus suggest that anodal stimulation of the temporoparietal area can produce an immediate reduction of tinnitus lasting a short time. Cathodal stimulation of the temporoparietal area may produce a similar effect, provided the duration of the stimulation is sufficiently long. This may be comparable with TMS, where short session of high-frequency TMS induces immediate change in tinnitus perception, while several sessions of low-frequency TMS induce prolonged decreases in tinnitus [43, 44].

tDCS of the Dorsolateral Prefrontal Cortex

New insights into the neurobiology of tinnitus suggest that neuronal changes are not limited to the classical auditory pathways [37, 38, 41, 45]. In particular, the dorsolateral prefrontal cortex (DLPFC) seems to play a specific role in auditory processing; the DLPFC has a bilateral facilitatory effect on auditory memory storage [46] and contains auditory memory cells [47]. The DLPFC also exerts early inhibitory modulation of input to the primary auditory cortex in humans [48] and has been found to be associated with auditory attention [46, 49, 50] resulting in top–down modulation of auditory processing [51]. This was further confirmed by electrophysiological data indicating that tinnitus might occur as the result of a dysfunction in the top–down inhibitory processes [52].

Interestingly, several tDCS studies focused on DLPFC and found successful results for treating major depression [53] and mood changes in depression [54], as well as reducing impulsiveness [55] and pain threshold [56, 57]. As the DLPFC is involved in attention-mediated top–down control of auditory processing and tinnitus, and tDCS seems to be a promising tool for modulating the DLPFC, it is possible that bifrontal application of tDCS might be a useful technique for the suppressing tinnitus. The common rational is to modify activity in the prefrontal cortex and also to re-establish the balance of left and right prefrontal cortex activation.

In a preliminary study involving 418 individuals with non-pulsatile tinnitus, it was shown that tDCS,
with the anode over the right DLPFC and the cathode over the left DLPFC, could cause tinnitus suppression in 29.9% of the participants [58]. In contrast, recent results of a study of 28 individuals with non-pulsatile tinnitus indicate that bilateral application of tDCS, with anode on the left and cathode on the right DLPFC, has no suppressive effect on tinnitus. Taken these results together, suppression of tinnitus by tDCS seems to be related by the ability to enhance excitability of the right prefrontal cortex and reducing the excitability of the left prefrontal cortex. A comparison between both groups did not show differences in tinnitus duration, tinnitus laterality, or tinnitus type and could therefore not explain the obtained results.

In conclusion, these studies indicate that anodal stimulation of the right DLPFC can produce an immediate reduction of tinnitus in some individuals. However, repeated sessions of tDCS might have better effects than single sessions as used in the present pilot study. Previous studies have already shown that anodal tDCS of the left DLPFC can affect depressive symptoms after one daily 20-min session of tDCS for a duration of 5 days [54]. Further studies are needed to explore the potential of frontal tDCS in reducing tinnitus-related distress.

Conclusion

Preliminary studies suggest that tDCS can modulate tinnitus in some individuals. However, further clinical and neurobiological research is needed before tDCS can be considered a practical treatment option for routine use. Therefore, multicenter placebo-controlled randomized trials with many patients and longer follow-up periods are required in order to estimate the efficacy of tDCS for the treatment of tinnitus. Further research is also needed to define selection criteria for patients for tDCS treatments. It may be possible to optimize and individualize stimulation protocols.

References

Chapter 90
Auditory Cortex Stimulation for Tinnitus

Dirk De Ridder and Sven Vanneste

Keypoints

1. The most frequent cause of tinnitus is hearing loss.
2. The auditory deprivation can lead to pathological theta–gamma coupling linked to a decrease of alpha oscillations also known as thalamocortical dysrhythmia.
3. Auditory deprivation also leads to auditory tract and auditory cortex tonotopic reorganization via activation of neural plasticity.
4. Presenting the missing information can reverse cortical reorganization.
5. Cortical stimulation can also reorganize tonotopic organization.
6. Auditory cortex stimulation can decrease tinnitus.
7. Auditory cortex stimulation interferes with ongoing oscillatory activity.
8. One in three patients responds to tonic stimulation and one in three to burst resulting in two out of three patients responding to auditory cortex stimulation.
9. Average improvement for auditory cortex stimulation is 50%.
10. Individuals with pure-tone tinnitus respond best to tonic stimulation of the cortex.
11. Individuals with noise-like tinnitus respond best to burst stimulation.
12. Tinnitus duration gender, or age is not predictive for successful stimulation.

Keywords Auditory cortex stimulation • Tinnitus • Deafferentation • fMRI • Gamma • Darwin • Plasticity • Reorganization

Abbreviations

BOLD Blood oxygen level dependent
EEG Electroencephalography
ERP Event related potential
fMRI Functional magnetic resonance imaging
Hz Hertz
iEEG Intracranial EEG
MEG Magnetoencephalography
MSI Magnetic source imaging
PET Positron emission tomography
rTMS Repetitive transcranial magnetic stimulation
TRI Tinnitus research initiative

Introduction

Until recently, people suffering from tinnitus were told “to learn and live with it.” This was largely due to the fact that there were no treatments available because of a lack of knowledge on how tinnitus is generated and the fact that tinnitus was considered solely an ear problem. In recent years, however, our understanding of the brain mechanisms involved in the generation of tinnitus has increased quite substantially [1, 2]. Even though for the majority of individuals with tinnitus, the original problem was located to the ear, and more specifically to hearing loss, most forms of tinnitus are caused by pathologic changes in the function of the brain.

Tinnitus Intensity

Tinnitus can be considered as a phantom phenomenon [3], similar to phantom pain [4–7]. People are well aware that when a hand or another part of the body is...
amputated, the missing part can generate phantom feelings [8]. This occurs in up to 85% of amputations, and in 15% the feeling is expressed as phantom pain [8]. Hearing loss, being considered analogous to amputation, can thus induce a phantom percept (such as phantom sound), better known as tinnitus. Much of the advancement in brain research on tinnitus is based on what is known from pain [5–7].

A heuristic model of tinnitus (see Chap. 21) is based on studies of consciousness suggesting that any conscious percept, including tinnitus, is related to gamma band activity (30–80 Hz) [9]. At rest, the auditory thalamocortical loop produces oscillations at alpha frequencies (8–12 Hz). When there is hearing loss, the cells that do not receive information from the cochlea will initially oscillate at lower frequencies (theta, 3–7 Hz) because there is less information to be processed [10]. In the brain high frequency activity (>10 Hz) in cells will suppress the activity in surrounding cells through lateral inhibition [11]. However, 10-Hz activity does not produce lateral inhibition [11], and lateral inhibition from activity at lower frequencies (<10 Hz) is decreased [12]. At low theta frequencies, lateral inhibition will decrease, inducing a halo of high-frequency gamma activity, also called the “edge effect.” So when there is hearing loss, due to slower firing rate caused by the nerve fibers transmitting the missing frequencies, a decrease in lateral inhibition (i.e., less suppression of the surrounding activity arises) will result in an associated halo of faster gamma band activity (30–80 Hz) at the lesion edge. This is called thalamocortical dysrhythmia [13]. Magnetoencephalography (MEG) studies have indeed shown that in tinnitus patients, decreased alpha is linked to increased gamma [9, 14, 15] and theta coupled to gamma [12, 13], supportive of this idea.

The brain can be considered a Darwinian structure, analogous to nature [16]. An animal can be considered analogous to the synapse that connects nerve cells. Synapse formation is analogous to animal reproduction; competition for connections is analogous to competition for resources and survival of the fittest synapse analogous to survival of the fittest animal [17]. Therefore, the cortex cells that do not receive information, also known as deafferented cells, will look for information in order to survive and open connections to neighboring cells, processing the same incoming information as the neighboring cells but generating the perception of the frequency to which these cells are programmed to process. This hypothetical mechanism is called Darwinian plasticity [18] and is basically analogous to dendritic plasticity [19, 20]. Subsequently, the decreased oscillation rates of the deafferented cells will increase to the same rate as the halo as such abolish thalamocortical dysrhythmia. Due to increased lateral inhibition of the deafferented area and lesion edge, a new halo of low-frequency activity will develop around the lesion edge. This could be called reverse thalamocortical dysrhythmia (see Chap. 21). No consensus exists yet on the exact mechanism of how tinnitus is generated in the brain, but the hypothetical mechanism could explain that the tinnitus pitch matches the frequencies of the deafferented nerve cells.

Another explanation of how certain features of tinnitus may be related to plastic changes in the nervous system suggests that “re-routing” of activity through thalamic and cortico-cortical connections could transmit information from a lesion edge toward the deafferented area. This is parallel to axonal sprouting into a deafferented region. The main difference between these two models is that in Darwinian plasticity the auditory cortex cells that do not receive information any more will attract information, whereas in classical plasticity the auditory cortex cells that are adjacent to the cells that do not receive information invade the area of deprived cells [21]. Probably both dendritic and axonal sprouting occur simultaneously.

Irrespective of whether Darwinian plasticity or classical plasticity can explain generation of the neural activity that causes tinnitus, the end result is that the deafferented cells become hyperactive and cause a phantom sound.

### Reafferenting the Auditory Cortex

The tinnitus treatment based on this model focuses on reafferenting the auditory cortex, which means that the missing information is supplied back to the auditory cortex and differs from the classical tinnitus treatments which predominantly target the ear. This does not mean that the ear is not important in tinnitus generation, as the changes in the brain are induced by a lack of information from the ear. Therefore, everybody suffering from tinnitus should undergo a complete tinnitus work-up by a specialized neuro-otologist (see Chap. 46).
Reafferenting the auditory cortex can happen in two ways: by compensating for the hearing loss to normalize the input to the cortex or by reafferenting the cerebral cortex. Reestablishing input to the cortex can be done by using hearing aids [22] (see Chap. 74) or cochlear implants in completely deaf people [23] (see Chap. 77), both of which are capable of reducing tinnitus. If this fails, the auditory cortex can be reafferented electrically by supplying the missing information directly or indirectly to the deafferented area of the auditory cortex. The electrical information can be supplied indirectly by electrical stimulation of the auditory nerve [24, 25], cochlear nuclei [26], inferior colliculus, or thalamus. The missing input to the cortex can be supplied directly to the deafferented area of the cortex by electrical stimulation [4, 27–30]. Electrical stimulation of the cortex may also result in suppressing the tinnitus by interfering with the neural network that functions abnormally instead of specifically reafferenting the cortex.

Electrical simulation through an electrode that is placed on the area of cortical hyperactivity can reestablish normal organization of the reorganized maps [31] through egocentric selection [32] of the entire tonotopic pathway all the way to the cochlea [33]. Another explanation for tinnitus relates the disorder to hypersynchronous gamma band activity [9, 13, 34]. The gamma band activity might code the tinnitus intensity [34], and the tinnitus percept per se could be the result of an emergent network property [15, 35, 36]. In a recent study using MEG during electrical stimulation of the auditory cortex, the stimulation increased spectral correlation across low and high gamma band activity; between alpha and beta activity, but delta/theta activity decreased, suggesting that auditory cortex stimulation does indeed affect thalamocortical dysrhythmia [37].

Electrical stimulation might, thus, do nothing more than disrupt the abnormal thalamocortical dysrhythmia embedded in a larger tinnitus network (see Chap. 21), and subsequently the emergent property of the network, the tinnitus, disappears.

Interfering with Tinnitus-Related Distress

Interfering with tinnitus-related distress requires an understanding of the pathophysiology of tinnitus distress. The exact mechanisms are unknown, but based on the available literature and data from research on pain [38, 39], dyspnea [39], and post-traumatic stress syndrome (PTSD) [40], it seems likely that a “general distress network” exists, consisting of the amygdala, anterior cingulate, and anterior insula. Activity in this network might generate a feeling of distress, perceived as tinnitus distress if the activity in the “general distress network” is synchronized with the dysrhythmic activity in thalamocortical loop, i.e., with the (theta and/or) gamma activity in the auditory cortex [15] (see Chap. 21). This is consistent with the hypothesis that broad band [41] and gamma synchronization [42] is a potential binding mechanism to generate a unified percept of the simultaneously presenting stimuli.

Auditory Cortex Stimulation in Clinical Practice

Auditory cortex stimulation is based on a four-step rationale (Fig. 90.1):

1. Tinnitus is related to synchronized gamma band activity.
2. Synchronized gamma band activity correlates with the blood oxygen level-dependent (BOLD) imaging signal of functional MRI (fMRI).
3. fMRI-guided neuronavigated transcranial magnetic stimulation (TMS) can suppress gamma band-related tinnitus.
4. If TMS is successful in tinnitus suppression, fMRI-guided neuronavigated electrode implant can suppress tinnitus permanently.

For successful treatment of tinnitus, two problems must be addressed. First of all, the exact localization in the brain of the auditory cortical hyperactivity must be determined. This is done by means of magnetoencephalography (MEG) [30] or fMRI [29, 43, 44]. In fMRI, two MRI scans are combined: a morphological brain scan and a scan performed during auditory exposure. In the MRI machine, the tinnitus pitch is first determined by tinnitus matching. Subsequently, the tinnitus pitch the patient perceives is presented via earphones, assuming that the generator of this frequency is the same as the tinnitus generator. Since both the activities seen on fMRI scan, i.e., the BOLD effect
and the tinnitus [9], or tinnitus intensity [34], are related to gamma band synchronization [45], the activity seen on fMRI can be correlated to the anatomical location of the tinnitus generator. In a similar way, the MEG can be fused with an MRI scan to obtain a magnetic source imaging (MSI) in order to localize the area in the brain generating the neural activity that causes the perception of a phantom sound. These tinnitus-matched frequencies behave differently than the non-tinnitus-generating frequencies in the auditory cortex (Kovacs, unpublished data), suggesting that presenting the tinnitus-matched frequencies might indeed be capable of demonstrating a part of the auditory cortex that functions abnormally.

The second problem is localizing the zone of hyperactivity exactly from results of the scan of the patient’s brain. The use of neuronavigation guided by the fMRI is helpful in this task.

Subsequently, a non-invasive stimulation can be performed targeting this area on the auditory cortex that shows abnormal activity. This is done by TMS.

If this non-invasive test shows that the abnormal neural activity assumed to cause tinnitus has been successfully suppressed, an electrode can subsequently be placed extradurally, overlying the secondary auditory cortical area of hyperactivity for permanent tinnitus suppression (on the exact same site as where the TMS was successful). The electrode is activated by an internal pulse generator, similar to a cardiac pacemaker, placed in the abdomen. The stimulation parameters (frequency, amplitude, and pulse width) can be changed postoperatively by remote control to find the best parameters for maximal tinnitus control.

In summary, the hypothesis behind auditory cortex stimulation for treatment of tinnitus is: (1) tinnitus is related to gamma band synchronization of the auditory cortical activity, (2) the anatomical location of the tinnitus generator can be determined by fMRI, (3) the activity of neurons in this location can be modulated by non-invasive TMS applied with the aid of neuronavigation, and (4) if TMS can suppress the tinnitus electrical stimulation through an electrode implanted on the same area, it can permanently provide the same tinnitus suppression by electrical stimulation as was achieved using TMS.
Implantations

Three different methods have been developed for auditory cortex stimulation via implanted electrodes:

1. Extradurally, on an area overlying the secondary auditory cortex [28, 29].

2. Intradurally, on the surface of the brain, placing the stimulating electrode in an existing groove or sulcus (intraculcal grey matter) of the primary auditory cortex. Such placements of the stimulating electrode provide predominant stimulation of cell bodies and not so much of the incoming or leaving fibers [28, 46], and

3. Intradurally, inside the brain similar to deep brain stimulation. Such placement of the stimulating electrode will activate intraparenchymatous white matter, thus, nerve tracts of fibers coming into and leaving the primary auditory cortex [30].

The electrodes that are implanted can have 4–16 electrode contacts (Figs. 90.1 and 90.2).

The surgery for an extradural electrode placement has been described in some detail [29, 47] and has minimal risk of complications. An incision is made 5–6 cm above the external ear canal, based on the fMRI. The location of the auditory cortex varies among different individuals and between the left and right side. The location of the incision in the skin is therefore guided by the fMRI. The skin incision is about 5 cm long and followed by a split of the temporal muscle. A small 1 by 5 cm hole is made in the skull and the small sensory fibers that innervate the dura are coagulated. This is necessary because the electrical stimulation used for tinnitus suppression may also cause pain by activating these fibers. After that is completed, the stimulating electrode is placed on the exact spot with 1–2 mm accuracy based on the fMRI and sutured to the dura. The small skull defect is repositioned and fastened with small titanium screws and plates.

For intradural grey matter stimulation, the electrode is inserted in the posterior part of the Silvian fissure after opening the dura (Fig. 90.2). For intraparenchymal white matter stimulation, the stimulating electrode is inserted into the auditory cortex, parallel to the Sylvian fissure. In both intradural procedures, the dura is closed after the insertion of the electrode. After that, the lead to the stimulating electrode is tunneled to the chest where it is connected to an extension lead, and further tunneled to the abdomen where it is passed through the skin to the outside of the body.

The electrode leads that exit the abdomen are connected to a stimulator, usually after 3 days, as on the first day the tinnitus is often markedly decreased from the operation. During external trial stimulation, the different electrode contacts are activated, one by one, or more than one at a time, depending on what gives the best suppression. The trial sessions are limited to one hour because it is difficult for the patient to keep concentrating for longer times. Once a good suppressive effect is obtained, which can occur after one day (sometimes, it takes a week or even a month), a programmable internal pulse generator (IPG) is implanted in the abdomen and the electrode is connected with a new extension lead to the IPG.

Stimulation is not performed continuously because of the risk for eliciting an epileptic seizure. Usually, the stimulator is active for 5 s and switched off for 5 s. During these 5 s that the IPG is silent, the tinnitus remains suppressed because of residual inhibition. As the patient does not feel the electrical impulses, he or she does not know whether the stimulator is on or off. During the first period after the implantation, the tinnitus returns very quickly when the stimulator is turned off. After a couple of seconds, the sound starts to come back, thus the residual inhibition is not very long. However, after years of stimulation when the

![Fig. 90.2 Intraoperative picture showing the opened dura and electrode, inserted in the posterior part of the Silvian fissure](image-url)
stimulator is switched off or the battery has become drained, it may take weeks before the tinnitus returns full scale. It can only be hoped that after many years of stimulation, the tinnitus might stay away for longer and longer periods of residual inhibition and finally forever, even without further stimulation.

**Results**

In total, 43 patients with intractable grade 3 and 4 tinnitus, i.e., severe tinnitus according to the tinnitus questionnaire [48], were implanted with a cortical electrode overlying the secondary auditory cortex. Before implantation, all patients underwent tests in two TMS sessions on separate dates performed by a person not involved in the surgery. If TMS resulted in suppression of the tinnitus (>20% improvement on a visual analog scale (VAS)) on two separate occasions, the patients were regarded to be eligible for implantation.

Although all patients reacted to TMS, 1 out of 3 patients did not respond to the cortical stimulation after implantation. Two out of 3 patients responded to cortical stimulation with an average decrease in the perceived tinnitus loudness of 51.3%. A significant but weak positive correlation ($p<0.05$) between suppression effect from the test TMS and cortical stimulation after implantation was exists in responders to both TMS and electrical stimulation.

Of the patients who were implanted with a cortical electrode, the use of burst stimulation (5 stimuli of 1 ms pulse width, 1ms interpulse interval, at 500 Hz delivered 40 times a second) improved the total results dramatically. When only tonic stimulation is used, only one in three patients obtained tinnitus suppression. However, using burst stimulation, half of the non-responding patients could benefit. In a similar way, half of the patients who did respond to tonic stimulation had better suppression with burst stimulation. Earlier studies have shown that the suppression effect on TMS depends on how long the patient had tinnitus prior to TMS, indicating that the suppression effect on TMS decreases over time [15, 49, 50]. Similar results were obtained for the patients who were eligible for implantation. A negative correlation was found between tinnitus suppression from TMS and tinnitus duration $r=-0.37$, $p<0.05$. However, no correlation was found between the suppression effect based on electrical cortical stimulation and tinnitus duration. The latter result is quite interesting, as it suggests cortical stimulation is different from TMS and that it affects the neural activity that causes tinnitus in a different way than TMS since it seems to help in a way independent from tinnitus duration.

However, it was revealed that tinnitus type (pure tone, narrow band noise, or both) and laterality, whether unilateral or bilateral, whether unilateral or bilateral had a significant influence on the amount of suppression. Pure-tone tinnitus was suppressed more than narrow band noise or the combination of pure tone and narrow band noise and unilateral tinnitus was suppressed more than bilateral tinnitus.

Another study of eight patients using a similar technique but different hardware [27] showed only temporary tinnitus suppression in six of the patients studied. However, tinnitus distress decreased, even without suppression of tinnitus intensity. This is more similar to the effect of 1-Hz rTMS [51] than the results of our previous study [52]. Two explanations have been proposed for these differences [53], namely, a different stimulation device with different stimulus parameters and different electrodes. The stimulation parameters may be important for the ability to induce tinnitus suppression. Using an electrode with only two contacts limits the way the electrodes can be programmed. The tinnitus always reoccurs after auditory cortex stimulation using

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**Fig. 90.3** Summary of the results obtained by electrical stimulation of auditory cortex for the treatment of tinnitus

| 43 patients | 14 (32.56%) do not respond to cortical stimulation | 8 (50%) patients respond better on burst stimulation
| 29 (67.44%) respond to cortical stimulation | suppression effect for tonic stimulation 38.10% |
| 16 (55.17%) respond to tonic stimulation | suppression effect for burst stimulation 53.01% |
| suppression effect 51.31% | |
| 13 (44.83%) do not to tonic stimulation, but do respond to burst stimulation | 8 (50%) patients do not respond better on burst stimulation
| suppression effect 52.18% | |
| suppression effect 51.30% | |

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D. De Ridder and S. Vanneste
implanted electrodes and therefore requires the use of several different stimulation programs using multiple electrode contacts to run alternately to prevent tinnitus recurrence [29, 52].

The fact that tinnitus distress does decrease without tinnitus intensity reduction could possibly be explained by disruption of the phase synchronization between the "general distress network" and the thalamocortical dysrhythmia. Intracortical microstimulation in the auditory cortex of animals disrupts not only local ongoing activity but also long-range connections in a larger network [54], similarly to what has been described in humans using TMS of the auditory cortex [55].

Side Effects

Side effects are limited and do not occur at stimulation parameters required for tinnitus suppression. Side effects may occur when high-frequency, high-intensity stimulation is used. Different kinds of side effects have occurred when testing is performed to discover the limits of the applied stimulation design, so that permanent stimulation can be programmed without these side effects. A feeling of intoxication, altered spatial localization of external sounds, difficulty in finding appropriate words, dizziness, vertigo and hearing perception changes (hearing perceived as being clearer, related to even their own voice) [28], as well as out of body experiences [56] were noted. Some patients with tinnitus have an associated feeling of aural pressure, the feeling as if there is water inside the ear. In all patients with successful tinnitus suppression, these associated pressure feelings decreased, however. The stimulation designs that best suppressed tinnitus and best suppressed pressure feelings are not always identical.

Complications

Complications are limited but can be severe. Epileptic seizures occurred in 3 of the 43 patients studied. In these three patients, the epileptic seizures most likely occurred because of prolonged stimulation without interruption and also occurred while the patients still had an external stimulator which could not be programmed by the investigator (it relies on patient cooperation). Therefore, patients with epilepsy are not candidates for this procedure. One patient developed an epileptic seizure during prolonged trial stimulation. Therefore, trial programs should probably best be limited to 1-h sessions.

The major complications occurred with intradural implants. One of the four patients implanted with the electrode directly on the surface of the primary auditory cortex developed a postoperative intracranial bleed in the superior temporal sulcus at a distance from the Sylvian fissure, where the electrode was inserted. Speech disturbances occurred as a result. However, a decrease in tinnitus also occurred as a result of the bleeding. One of the four patients developed an intracranial abscess that required surgical evacuation, with good outcome. Thus, this treatment should be preferentially performed extradurally. Extradural techniques in our last 30 patients experienced no serious complications.

Failures of Auditory Cortex Stimulation

Not all patients benefit from stimulation via implanted electrodes as detailed above.

Some of the conceivable reasons for failure are:

- Contralateral auditory cortex is not involved in all patients
- TMS is not a good predictor of subsequent implant success
- The stimulation design is not optimal for the individual patient
- The neural network that causes the tinnitus has become permanent
- The adjustments in the network that causes tinnitus also change over time

We have placed the stimulating electrode on the contralateral secondary auditory cortex for unilateral tinnitus and on the right secondary auditory cortex for bilateral tinnitus. It is possible that all electrodes should have been implanted at the left auditory cortex similarly to what has been done for rTMS [51]. This suggestion is supported by the finding that PET studies usually show increased metabolism on the left auditory cortex [57, 58] in individuals with tinnitus, irrespective of the
side on which the tinnitus is perceived, and that TMS applied to the left side can suppress this metabolic activity [59]. On the other hand, fMRI [44, 58, 60, 61] and EEG [34] and MEG [9, 31] studies suggest that the neural generator of tinnitus is located contralaterally to the tinnitus side.

It is clear that TMS, even though performed twice and placebo controlled before every implant, is not a perfect predictor of success in subsequent implants. However, a correlation does exist between the amount of tinnitus suppression obtained by TMS and by stimulation via implanted electrodes.

Another reason for failure may be that the stimulus protocol used is not optimal for the individual patient. For example, noise-like tinnitus does not seem to respond to tonic stimulation, responding to burst stimulation instead [47]. If burst stimulation had not been used, the results obtained in the series of 43 patients discussed above would have been poor, with only 33% of patients showing benefit from the stimulation by implanted electrodes. In this study, 45% of patients only experienced benefit from burst stimulation.

Regarding the difference between the effect of TMS and stimulation with implanted electrodes, it is possible that TMS reaches more fibers or penetrates deeper into the auditory cortex than the implanted electrode. It is conceivable that TMS reaches auditory cortex fibers that go to the parahippocampus directly and therefore influences the parahippocampus where the implant does not. (The parahippocampal involvement in tinnitus is detailed in Chap. 21.)

Another explanation for the failure of cortex stimulation may be that the tinnitus network might have become too hard wired for stimulation to disrupt it. Even though this is conceptually possible, it is unlikely that this is the cause of the observed failures. While it is true that the results from TMS are affected by how long the patient has had the tinnitus [49, 55, 62–64], the results of our studies of patients with implanted electrodes indicate that the tinnitus duration does not seem to affect the outcome. Therefore, this argument is most likely not valid for explaining treatment failure. It, however, points to yet another interesting difference between TMS and stimulation from intracranial electrodes.

The tinnitus network might change in time. The recent development of network science [65–68] (see Chap. 21) with its application to tinnitus by the seminal work of Schlee et al. [15, 35, 36] has altered the way researchers think about the pathophysiology of tinnitus. Since tinnitus should be considered an emergent property of a large network, it is possible that the weight of the hubs and their individual’ connectivity change in time [15]. Therefore, whether stimulation is beneficial or not could be dependent on the state of the network; the exact state cannot be derived from group data but should be analyzed on an individual level of the patients eligible for implant. Further studies exploring the differences of resting-state activity, as recorded by EEG or MEG, between responders and non-responders could help to elucidate these prognostic problems.

The Future of Neurostimulation for Tinnitus

Based on the new network science, it should be possible to retrieve good alternative targets to the auditory cortex for neuromodulation. This requires a thorough analysis of resting-state data of an individual patient, looking for the hubs in a scale-free network model (see Chap. 21) of tinnitus. Once these methods become easily accessible, results of the promising technique of neuromodulation should improve.

Conclusion

Brain stimulation is an option for patients with severe and intractable tinnitus. With proper selection of the patients, extradural stimulation is capable of suppressing tinnitus completely or partially in 67% of patients. Extradural stimulation is preferred because of less risk of complications than intradural placements of the stimulating electrodes. Using the results from TMS seems logical as a prognostic criterion. However, even with TMS as a preoperative test, 33% or more of patients will still fail to benefit from stimulation through implanted electrodes. Several reasons why not all patients benefit from auditory cortex stimulation may exist. Development of new stimulation designs as well as the application of network science might, in the near future, improve results of the techniques.
References

**Chapter 91**

**Cutaneous Stimulation**

Aage R. Møller

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**Keypoints**

1. Electrical stimulation of the skin around the ears was one of the first described methods used clinically for treating tinnitus, but the results vary widely among authors.

2. Recordings from cells in the dorsal cochlear nucleus (DCN) show that stimulation of the skin around the ears can cause both increased and decreased excitation of neurons in the dorsal cochlear nucleus.

3. The skin around the ears is innervated by the upper spinal cord (C₂) and the caudal trigeminal nucleus, the neurons of which project to the DCN and the external nucleus (ICX) of the inferior colliculus. This is believed to be the basis for the observed effect on tinnitus from electrical stimulation of the skin.

4. Electrical stimulation of the middle ear mucosa (bony capsule of the cochlea) has also been used to treat tinnitus. The mucosa is innervated mainly by the trigeminal nerve.

5. A few studies have shown that cutaneous electrical stimulation in other places of the body can modulate tinnitus, indicating that the nonclassical auditory pathways are involved in such forms of tinnitus.

**Keywords** Cutaneous stimulation • Tinnitus • Trigeminal nucleus • Dorsal column nuclei

**Abbreviations**

- DCN Dorsal cochlear nucleus
- DR Dorsal raphe nucleus
- IC Inferior colliculus
- ICX External nucleus (of IC)
- LC Locus coeruleus
- Sp5 Spinal trigeminal nucleus
- STT Spinothalamic tract
- TENS Transderm electrical nerve stimulation
- V1 Upper branch of the trigeminal nerve
- VAS Visual analog scale

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**Introduction**

Cutaneous electrical stimulation has been described as one of the earliest methods for managing tinnitus by means of electrical stimulation [1]. Most studies have used various forms of electrical stimulation through electrodes placed on the skin around the ears [2]. Results from stimulation of the skin at other places of the body or peripheral nerves, such as the median nerve at the wrist, [3] have also shown to be an influence on some individual’s tinnitus. Electrical stimulation of the mucosa in the middle ear (bony capsule of the cochlea) has also been shown to modulate tinnitus in some individuals [4].

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**Electrical Stimulation of the Skin Around the Ears**

One of the first to describe the results of cutaneous electrical stimulation around the ears was Shulman and Tonndorf [2, 5]. Different investigators have used different kinds of electrical stimulation, such as
impulses presented at different rates, high frequency electrical current, and direct current (DC). The beneficial effects reported by the different investigators vary greatly.

Some investigators [2] using a commercially available device reported beneficial effect of 82% in a study of 27 individuals. The effect lasted 3 months or more in 47% of the participants. However, when the effect of using the same device was tested in a single-blind crossover study by other investigators [6], the benefit was found to be minimal (only 7% of the participants reported improvement). More recent studies of transcutaneous electrical stimulation [7] found improvement in 42.8% from electrical stimulation of the skin in front of the ears and 28.5% had improvement from placebo (sham stimulation with electrodes placed on the skin, but no stimulation). The study had 42 patients, 31 received stimulation, and 11 had electrodes placed on the skin, but no stimulation.

In a recent study of the effect of electrical stimulation through electrodes placed on the right and left C2 dermatome, Vanneste et al. found that such stimulation reduced the scores on a visual analog scale (VAS) of the strength of the tinnitus from a mean value of 6.16 (SD = 2.18) to 5.56 (SD = 2.42), thus 17.9% decrease [8]. Six participants had total disappearance of their tinnitus, and only 6% of the participants had a reduction in their tinnitus from the sham stimulation. The study had 240 participants with tinnitus (147 males and 93 females). Sixty-three of the participants had left-sided tinnitus, 44 had right-sided tinnitus, and 133 had bilateral tinnitus; 65 had pure tone tinnitus, 164 narrow band tinnitus, and 11 had both. The mean duration of their tinnitus was 6.19 years (SD = 7.92).

The stimulation lasted 10 min and consisted of constant current impulses at a rate of 6 per second, followed by 10 minutes stimulation at a rate of 40 impulses per second. The impulses (duration 250 µS (microseconds)) were presented at a subthreshold intensity. Sham stimulation was done with the same electrode placement and the stimulator turned on only for 30 seconds. The results were independent on the kind of tinnitus.

In a large study comprising 500 individuals with tinnitus, Steenerson and Cronin [9] found that 53% had improvement from electrical stimulation of the skin around the ears with at least two points on a subjective rating scale [10]. A month after the last treatment, 72% reported sustained benefit (when individuals with Ménière’s disease were excluded, 94% had sustained effect).

In a prospective descriptive study of 26 individuals with tinnitus, Herraz and coworkers [11] found that electrical stimulation of the skin around the ears improved the tinnitus in 46% of the participants (23% did not hear it anymore, and in another 23% its intensity was reduced). When using a VAS, scores improved from 6.5 to 6.0 after 2 weeks of treatment (p < 0.01). The participants used the electrical stimulation at home for 2 h, once per day for 2 weeks [alternating ramped burst, 150 pps, with pulse duration of 100 µS, the average intensity was 27 mA].

Intermittent “typewriter” type of tinnitus [described by Levine [12]] was the most responsive. The participants in the study were selected according to two criteria:

1. Their tinnitus was triggered by an acute somatic event.
2. The tinnitus could be modulated by orofacial movements or posture changes. Eight of the participants had “typewriter” tinnitus. This study shows the importance of selecting patients for treatment.

While these studies show that existing tinnitus can be modulated, tinnitus may also be elicted by the activation of the somatosensory system through touching the skin or through muscle contractions [13], change in gaze, etc. [14].

Mechanisms of Beneficial Effect from Stimulating the Skin Around the Ears

The investigators who placed stimulating electrodes on the skin behind the ears [2] were interested in stimulating the cochlea. However, electrical current from stimulation of the skin around the ears with tolerable intensities does not reach the cochlea with a strength that can possibly have any biological effect. Instead, such stimulation activates receptors or nerves in the skin. The skin behind the ears has dual innervations by spinal C2 and the upper branch of the trigeminal nerve (V1). The effect on the auditory nervous system, and thereby modulating tinnitus, is most likely caused by activating neurons in the trigeminal nucleus and the dorsal column nuclei that project to the dorsal cochlear nucleus (DCN) [4, 15–18] (see Chap. 9). This means that the beneficial effect on tinnitus from electrical stimulation of the skin was achieved by the effect the stimulation had on the neurons in the DCN. Animal studies by other investigators have found it likely that the DCN is involved in tinnitus [19, 20].

Recordings from cells (multunit recordings) in hamsters’ DCN show sound stimulation after the animals had been exposed to high intensity sounds of the kind that normally cause tinnitus. These animal studies [21] have shown that electrical stimulation of the pinna, and in the spinal trigeminal nucleus (Sp5), dorsal raphe nucleus (DR), and locus coeruleus (LC) increase activity in neurons in the DCN (number of Fos-positive neurons increases). The sound stimulation caused hyperactivity that lasted several weeks after exposure [22].
Electrical stimulation of the skin behind the pinna caused several different kinds of changes in the recorded unit or multiunit activity. Suppression—suppression, excitation—suppression, suppression—excitation, and excitation—excitation were observed. The results were interpreted to show DCN hyperactivity is a direct neural correlate of tinnitus and somatosensory electrical stimulation can modulate DCN hyperactivity.

Results of tracing experiments indicate that the DCN received inputs from the Sp5, DR, and LC. The above results suggest that the modulation of DCN activity through somatosensory electrical stimulation may involve both direct pathways via the Sp5 and indirect pathways via the DR and LC. The relief of tinnitus caused by somatosensory electrical stimulation may involve manipulations of both auditory and nonauditory functions. For example, the stimulation of trigeminal fibers that innervate the cochlea may affect blood flow in the cochlea [23, 24].

The nerves that are activated by electrical stimulation of the skin behind the ears or in the middle ear cavity contain fibers of different diameters. These different fiber types innervate different populations of cells in the trigeminal nucleus and the dorsal horn of the spinal cord and project to different structures in the brain. It is not known which fibers are most effective in modulating tinnitus, but it seems likely fibers that belong to the pain system and those that mediate innocuous stimulation (touch and vibration) may have a different ability to cause tinnitus suppression. Electrical stimulation activates all these different types of fibers, but to different degrees depending on the stimulus parameters.

Receptors in the skin are innervated by three kinds of nerve fibers, large myelinated fibers (Aβ), small diameter myelinated fibers (Aδ), and unmyelinated small diameter fibers [25]. These fibers innervate different kinds of cells in the spinal cord (see Chap. 15) and the trigeminal nucleus. These cells project centrally in different pathways [26]. The Aβ fibers mediate touch and vibration. The fibers terminate on cells in the spinal cord located in the dorsal column nuclei from which fibers cross the midline and form the medial lemniscus – thus, the classical somatosensory pathway. Aδ fibers terminate on cells located in layer I of the dorsal horn of the spinal cord. C fibers terminate on cells in layer II, which make connections to cells in layer I. From these connections, fibers cross the midline at segmental levels and form the anterior spinothalamic tract that mainly mediates pain as well as hot and cold sensations. C-fibers mediate slow burning pain. These fibers terminate on other cells in the spinal cord, the projections of which also join the anterior lateral system. The Aβ fibers have inhibitory influence on the cells that receive pain signals through Aδ and C fibers. Trigeminal nerve fibers are similar, and all fibers have synaptic contact with cells in the trigeminal nucleus. It is not known which fiber groups of cutaneous nerves mediate the effect on tinnitus from cutaneous electrical stimulation.

Nerve cells innervated by cutaneous receptors can also be activated by mechanical and by chemical stimulation. These forms of stimulation do not seem to have been used for the management of tinnitus, but may offer advantages or electrical stimulation which is perceived by many individuals as being unpleasant.

Little attention seems to have been paid to the parameters of the electrical stimulation used for pain and tinnitus control. Electrical stimulation of nerve fibers can activate or inactivate the target cells depending on the frequency of the stimulation. Low frequency stimulation typically activates the target nerve cells while high frequency stimulation may constantly depolarize cells, and thereby inactivates them.

It would be interesting to compare specific stimulations of these receptors, such as using capsaicin that activates pain receptors compared with vibrations used for pain control [27]. Such stimulation activates receptors innervated by larger myelinated fibers (Aβ) [23].

Acupuncture, which may be regarded as a form of electrical stimulation of cutaneous nerve fibers, has been used for pain control [27] and tinnitus [28].

**Stimulation of the Surface of the Cochlea**

Placing electrodes on the bony capsule of the cochlea has been used to treat tinnitus. It has been assumed that the observed effect was caused by electrical current reaching the hair cells in the cochlea [29]. The mucosa of the middle ear, and also the surface of the cochlear capsule, is innervated by several somatosensory structures such as the dorsal root ganglia of C2–4 and the trigeminal nucleus. The latter contributed the most [30]. Electrical stimulation applied to electrodes placed on the cochlear capsule may therefore activate the
somatosensory system in a similar way as stimulating the skin around the ears. There are also autonomous fibers in the middle ear mucosa and in the skin, stimulation of which may change blood flow in the cochlea [23] which may affect tinnitus.

Subcutaneous Stimulation of the Skin at Other Places of the Body

Other investigators placed stimulating electrodes on places of the body other than the skin around the ears and found that stimulation could affect some individuals’ tinnitus. Stimulation of the medial nerve at the wrist that innervates the skin of the hand can modulate tinnitus in some individuals [3].

Cacace and coauthors described [14] a form of tinnitus that could be evoked directly by cutaneous stimulation of the upper hand and fingertip regions in two adults after surgical removal of space-occupying lesions at the base of the skull and posterior fossa. Hearing and vestibular functions were lost in one ear and facial nerve paralysis was present after the operation. It was assumed that these abnormal sensations were caused by anatomical and physiological interactions between auditory and somatosensory structures in the brain.

Since somatic tinnitus can be elicited by muscle contraction, treatments that reduce contractions (botulinum toxin) have been tried as a treatment for some forms of tinnitus [31].

Anatomical Basis for Involvement of the Nonclassical Pathways in Modulation of Tinnitus by Cutaneous Stimulation

It was mentioned above that somatic input to the cochlear nuclei may be responsible for the effect on tinnitus from electrical stimulation of the skin around the ears. Increasing the somatic input to the nonclassical auditory pathways is another possible explanation for the beneficial effect of electrical stimulation of the skin around the ears. When the nonclassical auditory pathways are active, somatic stimulation may influence (modulate) the activity in the nonclassical auditory pathways. It is generally assumed that the nonclassical auditory pathways are involved in the cross-modal sensory effects between somatic and auditory senses. Such cross-modal interaction has been studied in connection with tinnitus [3, 32, 33] and in connection with the perception of physical sounds [3, 34].

Zhou and Shore [35] have shown in anatomical studies that the external nucleus (ICX) of the IC receives projections from the Sp5 (in the guinea pig). The ICX is known to belong to the nonclassical ascending auditory pathways [36] and receives projections from mostly contralateral DCN [35]. The DC and ICX receive somatic input mainly originating in somatosensory innervation of the upper part of the body [36]. Electrical and other forms of stimulation of the skin, joints or muscles can thereby influence (modulate) the nonclassical auditory pathways.

Conclusions

Electrical stimulation of the skin can affect tinnitus; for some individuals, it can relieve tinnitus. The exact mechanisms for the effect are complex and incompletely understood, but it may have similarities with the techniques that have been in use for many years in the management of pain such as TENS. New discoveries regarding anatomical and functional connections between the somatosensory system and cochlear nuclei have provided new insights into the mechanisms of electrical skin stimulation.

References

Chapter 92
Complementary Tinnitus Therapies

Manuela Mazzoli

Keypoints

1. Several different complementary therapies have been attempted, but few studies have been published regarding the efficacy of complementary treatments for tinnitus.

2. Acupuncture seems to be mostly effective in acute and recent tinnitus as well as in somatic tinnitus. Other therapies that are in the area of musculoskeletal therapies, such as electrical stimulation applied to skin, the use of manipulations, or exercising, have been studied for somatic tinnitus.

3. There are strong indications that a metabolic component is involved in a subgroup of individuals with tinnitus, but only few studies have been performed of this aspect of tinnitus.

4. Training in mindfulness (including awareness), breathing techniques, meditation, and hypnosis are useful as complementary therapies for tinnitus that can reduce annoyance and fixation on the presence of tinnitus, improving sleep, anxiety, and the perceived quality of life.

5. Methodologies and study design for studies of efficacy of treatment are critical for the interpretation of the results. Difficulty to reach significance is not only an issue for complementary therapies, but also for drug trials or other kinds of therapies.

Keywords  Tinnitus • Complementary • Acupuncture • TENS • Manipulations • Nutrition • Metabolic • Mindfulness

Abbreviations

CIC  Chloride ion channels
CoQ  Coenzyme Q
MBSR  Mindfulness-Based Stress Reduction
MEG  Magnetic Electroencephalography
MSG  Monosodium glutamate
NADPH  Nicotinamide adenine dinucleotide phosphate
NGF  Neural growth factor
NMDA  N-methyl-D-aspartate
NOX-3  NADPH oxidase
OAE  Otoacoustic emissions
ROS  Reactive oxygen species
TENS  Transcutaneous electrical neural stimulation
THI  Tinnitus handicap inventory
TCM  Traditional Chinese medicine
TOAE  Transiently evoked otoacoustic emissions
TQ  Tinnitus questionnaire
VAS  Visual analogic scale
VMA  Vanillic mandelic acid

Introduction

So far, no drug or medication has been found to be clearly useful in treating tinnitus. There is, therefore, reason to study other and different complementary therapies, such as acupuncture and herbal therapy, as well as methods that have been found useful in the treatment of other conditions. Physical therapy, osteopathy, nutrition, and hypnosis or mindfulness have been proposed or sought after by patients for the treatment of tinnitus.

This should not be surprising since tinnitus mechanisms are complex and often associated with other symptoms; it
typically affects many aspects of a person’s life, such as sleep, mood, lifestyle, stress, perceived quality of life, etc. Before we judge these therapies as inadequate or unproven, we should consider that often in clinical practice there is wide use and acceptance of therapies that do not have any evidence-based efficacy from conventional surveys. For example, two surveys of the treatment used for Ménière’s disease showed large differences in the treatment used routinely by different physicians. In one study, Smith and colleagues [1] reported that 52% of ENT physicians treated patients with Ménière’s disease using many different medical and surgical therapies with little or no evidence of efficacy. The survey found that 94% of surgeons prescribed betahistine, 63% diuretics, and 71% advised salt restriction to their patients while 52% of surgeons continued to recommend endolymphatic sac decompression and 50% are still inserting a tympanostomy tube (PE tubes) in the ear drum despite suggestions these treatments have only a placebo effect [2].

Also the search for a treatment for tinnitus has been frustrating in all fields, including the more conventional pharmacology (see Chap. 42). This may be due to the lack of understanding of the pathology of tinnitus and of mechanisms underlying the different types of tinnitus in each patient. The fact that “annoyance” can be caused by symptoms such as fear, insomnia, etc., a nonhomogeneous design of the studies and the lack of more sensitive or objective measure for tinnitus and its annoyance are obstacles in the assessment of the efficacy of different treatments.

In this chapter, we review the most commonly used complementary therapies and different approaches aimed at reducing tinnitus annoyance. These are possibilities, at times promising, that should be explored in rigorous trials in order to understand if such treatments would benefit the tinnitus sufferers.

**Physical Treatment**

**Acupuncture**

Traditional Chinese medicine (TCM) is a complex and sophisticated diagnostic and therapeutic method used for the last 3,000–4,000 years, for which the Federal and Drug Administration (FDA) and the scientific community have approved the efficacy in the cure of several conditions [3]. According to TCM, the body is an intercorrelated system, and a healthy condition is obtained when all physiological functions are in a dynamic balance within the body and between body and environment (climate, food, physical activity, etc.). This concept in modern terms is defined as homeostasis. TCM treatment, that can include acupuncture or herbal pharmacology, is personalized for the patient since the same symptoms can arise from different types of unbalances.

According to classical textbooks of TCM, for acute tinnitus a cure can be attempted, it is more difficult to reduce chronic tinnitus (symptom of a long lasting unbalance in the body), with either acupuncture or herbal therapy [4].

Several studies have been done using a scientific approach to verify the efficacy of acupuncture in tinnitus therapy. Park and colleagues [5] reviewed 33 papers on the treatment of tinnitus with acupuncture. Most of these studies [6] did not report clinical trial, of the remaining, five were not randomized and only six included a control group in the trial. The results of these latter studies seem to indicate that acupuncture may be effective in some patients, although conclusions on the efficacy of acupuncture for tinnitus treatment cannot be drawn. In fact, in these studies, the participants who reported an improvement after acupuncture ranged between 8–84% [7–10], but in some studies the results were not significantly different from a control group [11, 12].

An interesting study by Podoshin and colleagues [10] compared the effect of three different treatments for tinnitus: acupuncture, biofeedback, and cinnarizine. Patients with known pathologic conditions, such as Ménière’s disease, vestibular schwannoma, or otosclerosis, were excluded. Sixty individuals with idiopathic subjective tinnitus were randomly divided into five groups to receive one of the three treatments mentioned above or placebo. Fifty-eight participants completed the study. Assessment was by subjective severity rating for tinnitus disturbance during activity and rest and by tinnitus matching. Although there was a nonsignificant trend toward the improvement in tinnitus disturbance in the acupuncture group, the percentage of participants who were improved with acupuncture (30%) was greater than the percentage of participants who were improved with cinnarizine (10%) or with either placebo biofeedback (0%) or placebo cinnarizine (10%), but less than with biofeedback (50%). Tinnitus matching showed no objective difference between either one of the three treatments and placebo.

In these studies, the design varied and methodologies of the treatments given to the participants were...
different (electrical, traditional, auricular acupuncture). In acupuncture, the treatment is often tailored to the participant’s individual needs. The tinnitus of the participants in these studies varied; the participants were not divided into subgroups according to such factors as their hearing status or whether their tinnitus was acute or chronic, with or without somatic components, etc. The measures of tinnitus discomfort were highly subjective – only visual analog scales (VAS) were used. However, tinnitus questionnaires were not used, nor were anxiety questionnaires used in the evaluation. It is therefore difficult to draw definite conclusions on the efficacy of acupuncture treatment for tinnitus based on these earlier studies.

More recently, papers describing studies that were more rigorously designed, using a more conventional approach to tinnitus research, have been published.

Jackson et al. [13] described a rigorously designed study with a small series of six participants with chronic tinnitus (longer than 6 months) using an individualized treatment approach to obtain the best treatment according to TCM principles. This study found improvement in most of the parameters analyzed for most of the participants (loudness, pitch, THI, the number of waking hours affected by tinnitus, and the quality of sleep). The oldest person in the group (79 years) who had had tinnitus for 20 years did not improve, while the person who improved the most, was the youngest in the group with shortest duration of tinnitus. This is similar to what has been found in other studies analyzing other types of treatment, mainly that individuals who have had tinnitus for a long period have poor outcomes of treatment (see Chap. 10). The sample in this study was too small to draw definite conclusions.

In a study by Zhou and colleagues [14] similar conclusions were recorded on a greater sample (140 participants who had both hearing loss and tinnitus). In this study, neural growth factor (NGF) was injected into selected acupoints on channels that, according to TCM, connect to the ears. The control group received intramuscular injections of vitamin B1 and B12. The study found an overall improvement regarding both hearing and tinnitus in 78.6% of the treatment group and in 31.8% of the controls ($p<0.05$). The effect of the treatment was especially good in patients with milder hearing impairment, those with recent onset of the symptoms and younger age, thus confirming other studies showing that tinnitus that has lasted a short time is more likely to get favorable results of treatment.

In a study by Okada et al. [15], acute tinnitus improved significantly more in the acupuncture treatment group compared to the sham acupuncture measured using a VAS scale. Average duration of symptom relief (90.24 ± 77.5 h) varied from 106.9 h in the study group to 72.3 h in controls. Eight participants (10.5%) reported improved quality of sleep – not only four in the study group, but also four among the controls – and the authors suggest the use of acupuncture for the relief of acute tinnitus.

Acupuncture indeed seems to have a direct effect on the function of the inner ear objectively measured with otoacoustic emissions (OAE). In a study by De Azevedo et al. [16], 38 patients with tinnitus were randomized between treatment and sham acupuncture. Transient otoacoustic emission (TOAE) were measured in the two groups before and after the treatment as well as contra lateral suppression. After the treatment with acupuncture, the amplitude of TOAEs increased significantly while in the sham group there was no significant increase. They also found higher suppression of the tinnitus after the acupuncture session compared with the sham group.

These later studies are promising and emphasize the importance of rigorous study design. The participants in the studies of treatment for tinnitus should be divided in appropriate subgroups to better identify the kinds of tinnitus that can respond best to acupuncture treatment. It is also worthy to consider that it has been widely demonstrated that many somatic complaints can be treated successfully with acupuncture [17–20]. Therefore, if there are somatic components to the onset or modulation of the tinnitus, such as neck or head trauma or whiplash, or if there are chronic inflammatory problems in the neck and upper back or head, acupuncture may be a treatment of choice.

**Physical Therapies**

Several studies have shown evidence that the somatosensory system of the upper cervical region and head is involved in some forms of tinnitus (see Chaps. 9 and 10). Studies have shown that tinnitus can arise directly from a disorder of the head and upper neck through the activation of the somatosensory system, which can trigger or modulate the tinnitus in 64–80%
of the participants [6, 21–28]. Other studies have shown that tinnitus can be evoked or modulated by pressure on painful trigger points in the upper back, neck, or shoulders [29].

**Transcutaneous Electrical Nervous Stimulation**

The transcutaneous electrical nervous stimulation (TENS) is a clinical form of electrical stimulation of the somatosensory system. It is the electrotherapy most commonly used in physiotherapy for pain, muscle contractions, and inflammation in several neural and osteo articular conditions as well as those affecting tendons and ligaments (see Chap. 15).

There are few reports on the application of somatosensory stimulation for treating tinnitus and the results of studies of the efficacy of this treatment are still controversial (see Chap. 14).

Kapkin et al. [30] reported a rate of tinnitus worsening after TENS therapy of 16.6% (7/42) with 42.8% (6/14) in the placebo, and the rate of improvement after therapy was 42.8% (18/42). However, an improvement was seen in 28.5% (4/14) of the controls. This is in agreement with other studies that showed electrical stimulation of the median nerve at the wrist could cause tinnitus to increase in some individuals and decrease in others [31].

Herraiz et al. [155] reports the improvement of tinnitus in 46% of their sample of 26 individuals with tinnitus receiving 2 h treatment daily (alternated stimuli, 150 pps, pulse duration of 10 μs, 0–60 mA amplitude, mean amplitude 27 mA) for 10 days. If tinnitus was intermittent and not associated with other symptoms, results were more consistent. In this study, participants were selected having some clinical clues to somatic influence on tinnitus, such as painful trigger points and modulation of tinnitus with head and neck or jaw movements.

In a study by Aydemir et al. [32], after TENS treatment the subjective improvement of tinnitus measured by VAS scale was only marginally significant (p=0.059). However, after electrical stimulation, there was statistically significant improvement regarding tinnitus severity scores, tinnitus handicap inventory scores, NHP fatigue, social isolation, and emotional problems scores. Many parameters were measured by the SF-36 (p<0.05), such as physical functioning, general health, vitality, social functioning, role limitations due to emotional problems and mental health.

**Manipulations**

Many of the manipulative treatments used for musculoskeletal disorders, such as chiropractic manipulations, osteopathy, and massages may be considered in tinnitus treatment not only in patients with somatic tinnitus, but also in some other patients because these therapies may elicit reflex effects on nonmusculoskeletal symptoms.

In published data, chiropractic manipulations are used mainly for musculoskeletal disorders, but the improvements of nonmusculoskeletal symptoms after chiropractic manipulations have been described to occur in 2–10% of all patients treated and by 3–27% of those who complained to have nonmusculoskeletal problems [33, 34]. The success of spinal manipulative therapy, particularly of the atlanto-occipital joint, can be up to 82% of patients with dizziness (46% total relief, 36% high improvement). In contrast, only 10% of patients with tinnitus showed an improvement according to one study (p<0.001) [35].

In some cases, cervical problems, such as cervical degeneration or cervical instability, can present with symptoms mimicking Ménière’s disease: dizziness, fluctuating hearing loss, and tinnitus [36–38]. Cervical problems in the generation of tinnitus should be taken into account, especially in the elderly with a later onset of symptoms.

Osteopathy is a well-known system founded by Dr. A.T. Still (1828–1917), focusing on the diagnosis, treatment, prevention, and rehabilitation of musculoskeletal disorders and the effects of these conditions on a patient’s general health. Osteopathy is based on the principle that the body has the ability to heal. Osteopathic care focuses on strengthening the musculoskeletal systems to treat existing conditions and prevent illness. This holistic approach ensures that all treatment is tailored to the individual patient. According to osteopathic textbooks, the therapy for tinnitus aims at the identification of structural problems to correct; the relaxation of muscles especially in the neck, upper back, and TMJ; and the improvement of lymphatic local circulation.
In a randomized study comparing osteopathic with electrical stimulation of the skin over the neck, shoulders, and upper back (dynamic TENS, InterX®) for tinnitus, we have found the treatments reduced tinnitus annoyance measured as THI scores and VAS (for perceived loudness, percentage of time of annoyance and perceived quality of life) in 60% of the participants who were treated with osteopathic manipulations and in 46% of those treated with electrical stimulation, with a longer duration of the effects in the osteopathic treated group (Mazzoli et al.: in preparation). The benefit from both the osteopathic treatment and the electrical stimulation was more evident in patients with associated postural or somatic problems. No benefit was seen in the participants with noise-induced tinnitus.

The use of osteopathy could be useful not only in somatic tinnitus, but also in individuals with tinnitus that mimics chronic pain since osteopathic treatment can interfere with the mechanisms of modulation of pain [39].

Many individuals with tinnitus can modulate or evoke their tinnitus by the manipulation of myofascial trigger points [40, 41]. Trigger points are small hypersensitive areas of skeletal muscles in which a tight muscular band is often present. These small areas can be painful either spontaneously or after mechanical stimulation and can raise local or referred pain [42]. In the presence of painful trigger points, tinnitus can be an isolated symptom or be part of the so called myofascial syndrome, usually affecting the upper back or neck, and accompanied by several physical symptoms and clinical findings, such as sleep disturbance, lacrimation, vertigo, and skin reddening which all greatly affect the quality of life [43]. Trigger points can also be found in individuals who report no modulation of their tinnitus with the manipulation of trigger points. The presence of chronic pain in the areas of trigger points should be regarded as a characteristic that seems to correlate with modulated tinnitus compared to nonmodulated tinnitus [28], pointing once more to common mechanisms of action between pain and tinnitus (see Chap. 14).

Voluntary Exercise

If there is clinical evidence for somatic modulation of tinnitus, especially in noncontinuous tinnitus triggered, in particular, by some movements, exercising those movements can lead to habituation and a reduction of the tinnitus annoyance or to the disappearance of the symptom as described by Sanchez et al. [6]. In their study, they evaluated 38 individuals with tinnitus triggered or exacerbated by specific movements of the head and neck, as well as the shoulders or jaw. The participants were then instructed to exercise that movement for 10 min for 2 months and report the changes in their tinnitus and whether the exercise changed the pattern of tinnitus modulation. The participants were tested in different visits and the test retest was reliable for the modulation.

Influence on Tinnitus from Nutrition

Tinnitus can be triggered by drugs and is listed among the undesired effect of several medications (http://www.t-gone.com/tinnitus/drugs.asp) such as aspirin, diuretics, pain medications, etc. Despite this, there is little scientific evidence regarding metabolic or dietary treatments for tinnitus.

There are indications that the ear is particularly sensitive to nutritional and metabolic factors, specifically, several dietary changes are recommended for Ménière’s disease, such as reducing salt intake or eliminating allergenic foods [44–46]. Some individuals with Ménière’s disease may experience the increase of symptoms when drinking coffee or eating chocolate and other foods – similar to individuals with migraine [47] – and diet alone can work well in reducing the symptoms.

Even though up to now there are few scientific studies on metabolites or nutritional factors in tinnitus, we can try to analyze some substances that might be involved in triggering or modulating tinnitus.

Sodium Chloride

It is also known that salt intake may affect relapses in Ménière’s disease [48]. Some individuals with tinnitus report that after excessive salt intake their tinnitus increases. Sodium chloride may influence the inner ear in several ways: through its action on the blood pressure, inducing vasoconstriction in the cochlea,
increasing renal fluid retention, or acting on specific ion channels changing the composition of endolymph and therefore affecting the inner ear function. There is considerable relationship between impaired renal function and hearing loss [49]. Some inherited renal diseases are accompanied by hearing disorders, such as Alport syndrome and Bartter syndrome [50]. Also, the incidence of hearing loss is higher among patients with chronic renal failure than in the general population [51]. The renal adverse effects of some drugs (e.g., aminoglycosides and loop diuretics) may be accompanied by ototoxicity. Kidney and inner ear tissues are related immunologically, biochemically, and functionally. For example, the stria vascularis and the tubular epithelium in the kidney have similar ion transport processes [51] and the chloride ion channels of the ClC-K family are expressed exclusively in the kidneys and ears [52] and are involved in NaCl renal tubular reabsorption.

Monosodium Glutamate

Monosodium glutamate (MSG) is a sodium salt of the nonessential amino acid glutamic acid. It is used as a food additive and in its free form is commonly marketed as a flavor enhancer. It has the HS code 29224220 and E number E621. It is also included under the denomination “natural flavorings.” It is thought to cause the “Chinese restaurant syndrome” the symptoms that may include headache, throbbing of the head and ears, dizziness, tinnitus, lightheadedness, a feeling of facial pressure, tightness of the jaw, burning or tingling sensations over parts of the body, chest pain, and back pain, although mechanisms are yet unclear and the reality of the syndrome is controversial. [53, 54].

Glutamate is an excitatory neurotransmitter and is released in high quantity when hair cells are damaged by noise, or when ototoxic medications or infections affect the inner ear. An excess amount of glutamate is released, which leads to cell and neuron death mediated by high Ca++ flux into cells [55, 56].

No studies have been published that evaluate the real effect of MSG taken orally on the inner ear or auditory pathways, but damages to other organs have been consistently described in animal models [57–59]. In particular, it can induce diabetes [60], which is associated with increased risk for inner ear disorders and tinnitus.

Glucose

Several studies report a higher incidence of tinnitus and hearing loss in up to 27–76% of individuals who have diabetes, subclinical diabetes, or abnormal glucose metabolism, such as hyperinsulinemia (insulin resistance) or hypoglycemia [61–65]. The improvement of tinnitus after a diabetic-like diet has been reported [63].

Glucose metabolism can influence the inner ear in several ways. The inner ear, like the brain, does not have energy reserves. Its metabolism depends directly on the supply of oxygen and nutrients, including glucose from the blood supply. Alterations in glucose metabolism, therefore, have the potential to disturb the workings of the inner ear. Altered glucose metabolism can cause damage to several systems leading to peripheral neuropathy or microvascular disease. Fukushima et al. [66] described histological findings of the cochlea in individuals with diabetes type 2 that showed the walls of the vessels of the basilar membrane and stria vascularis in all turns were significantly thicker in individuals treated with either insulin or oral hypoglycemic than those of controls. The stria vascularis was atrophied in most turns of the cochlea in the insulin group and in the lower and middle turn of the oral hypoglycemic group. The difference was significant compared to the controls. The loss of cochlear outer hair cells was significantly greater in individuals treated with either insulin or oral hypoglycemic than those of controls. The stria vascularis was atrophied in most turns of the cochlea in the insulin group and in the lower and middle turn of the oral hypoglycemic group. The difference was significant compared to the controls. The loss of cochlear outer hair cells was significantly greater in the lower and upper basal turns in both diabetic groups while no significant difference was found in the number of spiral ganglion cells or inner hair cells between groups.

High levels of circulating glucose and hyperinsulinemia can also influence renal sodium reabsorption leading to salt sensitive hypertension [66–68], which could be mediated by oxidative stress factors [69, 70].

High levels of insulin are influenced by carbohydrate intake. It is therefore important to identify individuals at risk, that may include overweight patients with a craving for sugar or carbohydrates, who get lightheaded when fasting. Also blood testing could present hyperglycemia, although most frequently fasting glycemia can be normal while abnormal levels of glucose load curve response, hyperinsulinemia, hyperlipidemia, increased plasma aldosterone concentrations, or microalbuminuria, and sometimes chronic renal disease can be found.
**Artificial Sweeteners**

Discovered in 1965, aspartame is a low-calorie sweetener with a sugar-like taste but is approximately 200 times sweeter than sucrose. Aspartame is unique among low-calorie sweeteners in that it is completely broken down by the body to its components – the amino acids aspartic acid, phenylalanine, and a small amount of methanol. The safety of this aspartame is still controversial [71–74].

It has been reported that the consumption of aspartame could cause neurological and behavioral disturbances in certain individuals [73, 74]. Headaches, insomnia, and seizures are also some of the neurological effects that have been encountered, and these may be accredited to changes in regional brain concentrations of catecholamines (norepinephrine, epinephrine, and dopamine) [73, 74]. Consumption of large doses of aspartame in a single bolus dose will have an effect on some biochemical parameters, including plasma amino acid levels and brain neurotransmitter levels. The data from extensive investigations in humans into the possibility of neurotoxic effects of aspartame, in general, do not support the hypothesis that aspartame in the human diet will affect nervous system function, learning, or behavior [72].

Nevertheless, the use of aspartame or other sweeteners should be careful in specific patients, such as individuals with migraine, insomnia, dizziness, and possibly tinnitus. Even for diabetic patients, who seem to be the ideal consumers of the product, there are some concerns since aspartame intake has been associated to increase in baseline glucose and insulin levels in diabetes type 2 patients [71, 75].

**Antioxidants**

The suggestion that antioxidants should be used for the prevention or the repair of damage to the labyrinth comes from the studies on ototoxicity, although so far no clinical trials on patients with tinnitus have been published.

The most common ototoxic drugs in clinical use are aminoglycosides antibiotics, platinum-based chemotherapeutic agents (cisplatin and carboplatin), loop diuretics, macrolide antibiotics, and antimalarials. It is well established that oxidative reaction and free radicals in the cochlea are involved in causing damage to the cochlea from drugs and acoustic trauma [75–81].

Both aminoglycosides and cisplatin ototoxicity appear to involve the production of reactive oxygen species (ROS) in target tissues in the inner ear by activating the nicotinamide adenine dinucleotide phosphate (NADPH) oxidase, NOX-3, an enzyme unique to the cochlea. ROS can then deplete cochlear tissues of antioxidant protective molecules, for example, glutathione and antioxidant enzymes. This leads to a cascade resulting in oxidation of lipids, increased calcium influx, and apoptosis in cells of the cochlea [75, 78, 79]. The upregulation of endogenous protective mechanisms in the cochlea or the treatment with exogenous compounds reduces ototoxicity. Experimental studies in animals have shown that a variety of antioxidants, including aspirin, *Ginkgo biloba*, inhibitors of caspase-3, and caspase-9 can attenuate the ototoxicity of either aminoglycosides or cisplatin [79–82].

Similar mechanisms involving free radicals mediated damage can occur also in the nervous system at the level of the dorsal cochlear nucleus as well as inferior colliculus [83, 84]. Also, there is some evidence that psychological stress may cause oxidative damage in vivo, both in animal models and in humans [85–88].

A combination of antioxidant agents (vitamins A, C, and E) acts in synergy with magnesium to effectively prevent noise-induced trauma [87]. Neither the antioxidant agents nor magnesium used alone seems able to reduce noise-induced hearing loss or sensory cell death. In combination, however, they are highly effective in reducing both hearing loss and cell death, even with treatment initiated just 1 h prior to noise exposure [77, 89, 90]. These mechanisms could be involved also in aging and/or vascular problems associated with diabetes or inflammation generating tinnitus.

Preliminary results of a study by [91] on the effect of administration of Nanoquinone, a source of Coenzyme Q10 (CoQ10), on chronic tinnitus found no general improvement of tinnitus after increased CoQ10 levels. However, patients with a low CoQ10 level before treatment and with a significant increase in the CoQ10 level afterward showed a decrease of the total tinnitus score and of all its dimensions, except for emotional distress. CoQ10 is a 2, 3-dimethoxy, 5-methyl, 6-polysoprene parabenzoquinone, and is located in all membranes throughout the cell. The highest concentrations are
found in the heart, the liver, the kidneys, and the pancreas. It is an endogenously synthesized substance involved in a variety of essential processes, such as influence on the mitochondrial electron transport chain. However, it also appears to have membrane-stabilizing properties and to act as an antioxidant in conjunction with vitamin E [92].

**Zinc**

One of the few nutrients that have been studied in relation to tinnitus is zinc. Deficiency can be related to tinnitus, especially in the elderly [93]. Zinc deficiency affects many organ systems where it plays an essential role in numerous biochemical pathways, including the integumentary, gastrointestinal, central nervous system, immune, skeletal, and reproductive systems and is needed for maintaining DNA integrity [94, 95]).

The brain has the highest zinc content in the body [96] and is implicated in the function of glutaminergic neurons [97]. It has also been reported that hypozincemia activates the N-methyl-D-aspartate (NMDA) receptor, of the glutamate family, which may play an important role in the induction of epileptic discharge [97, 98]. Interestingly, it has been found that behavioral stress can modulate cellular influx of zinc changing the pattern of neural elements firing, especially in the hippocampus [96]. For these reasons, zinc has been proposed as a likely modulator of tinnitus.

Ochi and colleagues [99] have found that hypozincemia was related to the perceived loudness of tinnitus. However, in their study the average hearing sensitivities of patients with hypozincemia did not differ significantly from those of patients with normal serum zinc levels, suggesting zinc deficiency is likely related to tinnitus originating more centrally rather than hearing loss due to a peripheral disorder. On the other hand, the authors excluded people older than 59 from the study [99], and zinc deficiency increases over the age of 60 [93].

Most authors suggest administration of 50–66 mg of elemental zinc daily for individuals with hypozincemia [99–101]. The method by which zinc levels are measured can affect the definition of hypozincemia and may influence clinical decisions. Less than 2% of zinc in the organism is found free in plasma and most of the zinc in the body is located outside cells [93, 102]. Zinc level in the serum is therefore not a good measure for assessing zinc balance in the organism. In plasma, zinc is primarily bound to albumin and copper, and zinc is reciprocal in the serum. Because of this reverse and competitive correlation between zinc and copper, serum copper and albumin levels should also be assessed to prevent the risk that individuals given zinc supplementation develop hematologic abnormalities that occur in individuals with low copper levels [100].

**Herbal Therapy**

In TCM, there is a complex and sophisticated pharmacopeia where herbal remedies are knowledgeably mixed in combinations not only to obtain the desired synergic effect, but also to reduce side effects or correct the effect of specific ingredients. In TCM, the symptom is cured by reducing the functional unbalance that is assumed to cause the symptoms, and this can be different from patient to patient. A few randomized control studies on the treatment of tinnitus have been reported in the Chinese literature regarding the use of herbal drugs, claiming improvement in 40–55% of the participants, [103, 104]. Similar studies have been reported in behavioral animal models [105]. These studies have so far not been replicated at an international level.

Herbal medications should be considered as drugs since they contain pharmacological active substances in the form of phytocomplexes, some of which can be beneficial while others are toxic. Most modern medication derive from the isolation of the active substances (e.g., digitalis), and the herbal medications are often as effective as their synthetic counterpart [106, 107]. Nevertheless, investigations of the effects of herbal remedies on tinnitus, like other potential drug treatments, have often suffered from the lack of useful animal models and systematic clinical trials employing double-blind and placebo-controlled designs.

**Ginkgo biloba**

*G. biloba* leaves have been used therapeutically by the Chinese for centuries for the treatment of asthma and bronchitis and have also been used for tinnitus
relief. The active ingredient has been isolated as EGb-761 containing 24% flavonoids, 7% proanthocyanidins, and 6% terpenoids. The flavonoids are mainly flavonol-glycosides with antioxidant properties, while the terpenoid fraction contains ginkgolides, sesquiterpene, and bilobalide. Ginkgolide B, in particular, has potent platelet-activating factor (PAF) receptor antagonist properties. Many of the CNS effects of EGb-761 have been attributed to the combination of its antioxidant and PAF receptor antagonist actions. It is also a vasodilator and along with its antioxidant properties these are the reasons for thinking that it would be useful in the management of tinnitus [108].

Despite the claims that G. biloba extracts have some efficacy in treating tinnitus [109–111], there is very little objective evidence to support this. Hilton and Stuart [112] reviewed the clinical evidence relating to the use of G. biloba by individuals with tinnitus and concluded that there were no reliable data on which to base a conclusion, due to the methodological shortcomings of the available studies. In particular, very few studies have employed double-blind, placebo-controlled designs, where possible experimenter bias and patient expectation can be controlled. Where these sorts of controls have been used, the results have usually been negative [113, 114].

The use of G. biloba extracts can lead to potential undesired effects. Due to the fact that these extracts have vasodilator effects, combined with drugs, such as aspirin, they could potentially increase bleeding [114].

**Black Cohosh**

Black Cohosh (Cimicifuga racemosa), a buttercup plant grown in North America, is a popular preparation for the treatment of menopausal and other symptoms, including fatigue, neuralgia, rheumatism, sore throat, asthma, bronchial spasms, bronchitis, and whooping cough [115]. Black Cohosh has been used for centuries by women to stimulate menstrual flow, ease the strains of childbirth, and confer relief from premenstrual syndrome and menopause. With its mildly sedative and relaxing effect, Black Cohosh has been used also as a tinnitus herb to treat anxiety, nervousness, and chronic tinnitus. Black Cohosh may indeed have a dopaminergic effects and serotonin-binding properties in the brain. As a central nervous system depressant, Black Cohosh directly inhibits vasomotor centers involved with inner ear balance and hearing. As such, Black Cohosh has been used clinically for relief of tinnitus [116], although clinical trials on the therapeutic effect for tinnitus are missing.

There are few known health concerns regarding Black Cohosh, and consuming large amounts (over 5 g per day) is known to cause dizziness, vomiting, lowered blood pressure, limb pain, and can damage the liver [117].

**Ligustrum**

*Ligustrum* (*Ligustrum lucidum*) has been advocated by traditional herbalists for the management of tinnitus. It is considered, without scientific proof, to have a powerful liver and kidney protecting function; it supports adrenal function and has been found to have hypoglycemic, hypolipidemic, and antioxidant efficacy, and its use has been suggested for individuals with diabetes [118]. The recommended dosage for tinnitus is 400 mg three times per day. In this dosage, there are no known side effects from administration of this herb. No clinical trials have been published that have assessed its efficacy in tinnitus.

Other herbs have been suggested as possible remedies for tinnitus including: Mullein (*Verbascum densiflorum*); *Pulsatilla*; *Lycium* fruit (*Lycium barbarum* or *Lycium chinense*); *Cornus* (*Cornus officinalis*), only in association with Chinese fox glove root and Chinese yam; *Cuscuta chinensis* seeds are used alone and in combination with astragalus seeds (*Astragalus complanatus*), but no scientific studies of the effect on tinnitus of these herbs have been published for these herbs.

**Unconventional Treatment**

**Mindfulness**

Mindfulness involves bringing one’s awareness to focus on experience within the mind at the present
moment (from the past, the future, or the mechanical stream of consciousness). Mindfulness, such as meditation, yoga, Tai Chi, breathing techniques, etc., has been used in practice of spiritual healing in parts of the world for more than 5,000 years. During the last 40 years, the practice of meditation has become increasingly popular in Western countries as a complementary mind–body therapeutic strategy for a variety of health-related problems. By paying close attention to the present experience, practitioners begin to see both inner and outer aspects of reality as aspects of the mind in a nonjudgmental way, learning to observe without the continuous internal commentary or judgment. However, mindfulness does not have to be constrained to a formal meditation session. Mindfulness is an activity that can be done at any time and can be learned through several practical techniques that help reconnecting to the present moment each time the mechanical stream of thoughts drives us into our subjective mental “virtual reality.”

Presumably, via activation of limbic system parasympathetic pathways [119], mindfulness techniques have been shown to shift the balance between sympathetic and parasympathetic activation toward the parasympathetic in activity as studied in short- and long-term practitioners compared to controls. This included a reduction in heart, respiratory and pulse rates; of systolic blood pressure and oxygen metabolism; of urinary vanillic mandelic acid (VMA); and increases of skin resistance because of reduced sympathetic activity [120–122]. These physiological alterations are indicators of increased parasympathetic and decreased sympathetic activation [123] and therefore physiological relaxation that has been related to stress relief and may have a role in the prevention of stress-related illness, such as respiratory illness and hypertensive cardiovascular disease [124].

Also, mindfulness training has been found to improve chronic pain and associated symptoms [125–130], and given the similarities between tinnitus and chronic pain (see Chap. 14) [131], a beneficial effect of mindfulness in tinnitus patients can be expected.

Furthermore, there is evidence that mindfulness-based stress reduction (MBSR) programs are useful not only in treating chronic depression and anxiety problems [132, 133], but also in reducing depression relapses [134–136] and improving the measures of sleep quality or duration [137, 138]. Affective symptoms may increase the level of tinnitus annoyance. Also, there is evidence that the acceptance of the symptom obtained with mindfulness training has a therapeutic effect on tinnitus perceived annoyance [139, 140].

A marked reduction in alpha waves (8–12 Hz) at magnetic electro encephalography (MEG) recordings and an increase in delta waves (1.5–4 Hz) have been observed in individuals with tinnitus, especially when recorded from the temporal and left frontal areas. These anomalies have been associated with distress [141, 142, 156]. Meditation, on the other hand, causes increased theta coherence as well as increased alpha power [143]. The emergence of the slow (delta) waves in the attention-related frontal regions that occur during meditation provides strong support for the hypothesis that meditative states are not the same as relaxation states and that attentional processing is involved [144, 156]. This indicates that MBSR can be useful for the treatment of tinnitus since they have significantly higher everyday cognitive failures than nontinnitus patients, and this is related to the control of attentional processes [146]. Mindfulness training indeed improves attention skills possibly also reducing tinnitus-induced cognitive insufficiencies [147].

**Hypnosis**

Erickson’s hypnosis [148, 149] includes relaxation and emotional management techniques. The hypnotic trance leads to a modified state of consciousness characterized by a shift of hemisphere dominance from the logic dominant to the analogical dominant. The purpose of hypnosis, according to Erickson, was that of having access to the subconscious potential and natural learning ability while avoiding limited conditioned schemes [150].

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1 Erickson’s hypnosis: Hypnosis is a mental state (state theory) or set of attitudes (non-state theory), usually induced by a procedure known as a hypnotic induction, which is commonly composed of a series of preliminary instructions and suggestions. Milton H. Erickson, M.D. (1901–1980) was one of the most influential post-war hypnotherapists. He wrote several books and journal articles on the subject. During the 1960s, Erickson was responsible for popularizing a new branch of hypnotherapy, which became known as “Ericksonian hypnotherapy” Ericksonian hypnotherapy, eventually characterized by, amongst other things, the absence of a formal hypnotic inductions, the use of indirect suggestion, a “metaphor” (actually they were analogies, rather than “metaphors”), confusion techniques, and double binds (Erickson, 1977; Barker, 1986).
A few studies have been published regarding the treatment of tinnitus with hypnosis and the results seem promising. Ross and colleagues [151] reported the results of a study with 392 participants who were treated for 28 days with hypnosis. The tinnitus questionnaire (TQ) scores decreased in 90.5% of the participants who had subacute tinnitus and in 88.3% of those with chronic tinnitus. The improvement of the TQ score at the end of therapy was 15.9/14.1 points, which was highly significant. Effect sizes in the treatment groups (0.94/0.80) were superior to those in the waiting-list controls (0.14/0.23). The TQ score remained stable in the 1-year follow-up controls. Significant improvement in the quality of life has been observed after the treatment but depends on initial level of tinnitus severity.

In another study of hypnosis treatment on tinnitus, Maudoux et al. [152] showed that the THI score improved from 60.23 before hypnosis therapy to 16.9 at the end of the treatment, which was highly significant (p ≤ 0.005).

Other studies have shown that greater success of hypnotherapy was achieved in individuals who did not have hearing loss associated to their tinnitus [153]. This means that a subgroup of individuals would benefit more than others. The paper by Cope [154] provides a review of peer reviewed studies of the efficacy of hypnosis in the treatment of tinnitus.

Conclusions

Many of the complementary therapies described in the literature for treating tinnitus are promising, but the mechanisms of the beneficial effect are not known. Many of the studies that have been published indicate the importance of identifying subgroups of different types of tinnitus that would respond better to specific treatments. Methodology and study design are important to obtain clear indications of the efficacy of the therapies that are studied. Many of the tests that are used are not sensitive enough for tinnitus annoyance since THI questionnaires do not correspond to loudness or pitch of tinnitus, and are not sensitive enough. These issues contribute to the difficulty of defining “improvement” after tinnitus therapy and to compare the results between studies.

References

81. Yamasoba, T, Schachta, J, Shojia, F and Miller, JM, Attenuation of cochlear damage from noise trauma by an iron chelator, a free radical scavenger and glial cell line-derived neurotrophic factor in vivo Brain Res, 1999 815:317–325


103. Yang, DJ, Tinnitus treated with combined traditional Chinese medicine and Western medicine Zhong Xi Yi Jie He Za Zhi, 1989 9(5):270–271

104. Tan, KQ, Zhang, C, Liu, MX and Qiu, L, Comparative study on therapeutic effects of acupuncture, Chinese herbs and Western medicine on nervous tinnitus Zhongguo Zhen Jiu, 2007 27(4):249–251

105. Wang, H, Jiang, S, Yang, W and Han, D, Evaluating effects of some medicine on tinnitus with animal behavioral model in rats Zhonghua Er Bi Yan Hou Ke Za Zhi, 2000 35(5):331–334


111. Morgenstern, C and Biermann, E, Ginkgo-Spezialextrak Gb 761 in der Behandlung des Tinnitus aurium Fortschr der Medizin, 1997 115:7–11


144. Weisz, N, Dohrmann, K and Elbert, T, The relevance of spontaneous activity for the coding of the tinnitus sensation Prog Brain Res, 2007 166:61–70


Keypoints

1. Despite nearly 20 years of experience with low-level laser therapy (LLLT) for tinnitus concerns remain as to its effectiveness as a treatment modality for tinnitus.
2. Only a few reports show that LLLT is an effective treatment for tinnitus and other inner-ear conditions.
3. Many conflicting reports show no benefit whatsoever.
4. This chapter provides an outline of the biological basis for LLLT and reviews findings of controlled clinical studies of the use of LLLT in tinnitus treatment.

Keywords Tinnitus • Low level laser • Therapy • Biostimulation • Photostimulation

Abbreviation

LLLT Low level laser therapy

Introduction

The last 30 years have seen an enormous increase of research work in the clinical application of laser technology in Otorhinolaryngology. Surgical removal of tumors of the larynx and pharynx was revolutionized by the many advantages offered by “hard” or “hot” surgical lasers (such as Carbon dioxide, Erbium yttrium aluminum garnet, and Neodymium yttrium aluminum garnet). The ability to remove soft-tissue lesions under microscopic control in combination with excellent control of bleeding, due to the coagulative capacity of these lasers, results in good functional outcome even after extensive surgery [1]. The perforation of the footplate in stapes surgery represents another use of these different types of lasers [2]. Some otologists prefer to use a laser for stapes surgery because they do not have to touch the footplate manually.

At the other end of the spectrum of available lasers are semiconductor diode lasers, or combined helium–neon and gallium–arsenide lasers, which are sometimes referred to as “cold” or “soft” lasers. These lasers have only about one hundredth of the power of a surgical laser. In clinical medicine, diode lasers have been predominantly used to accelerate the healing of injured peripheral nerves [3, 4] and soft-tissue injury [5], and to reduce inflammation [6] and pain [7]. The clinical effectiveness for these applications termed low-level laser therapy (LLLT) or low-intensity laser irradiation or “biostimulation” is still controversial.

Biological Effects of LLLT

Low-Level Laser Therapy

Low-level laser therapy (LLLT) uses a light source for treatment. This light source is usually red to near infrared (wavelengths in the range of 630–904 nm) and is obtained from diode lasers or a combination of helium–neon and gallium–arsenide lasers. LLLT produces no noticeable heat, sound, or vibration. Instead, LLLT may act via non-thermal or photochemical reactions in the cells that are also referred to as “photobiology” or
“biostimulation.” The red visible and near-infrared laser wavelength used may be mainly absorbed in proteins, but the identity of the photoreceptors responsible for the biological effects of LLLT is unknown [8]. The laser light may act on the mitochondrial cytochrome system, endogenous porphyrins in the cell, or the energy-absorbing chromophores in LLLT [9]. In vitro studies using cell cultures have demonstrated stimulatory effects of such laser radiation on fibroblasts, immune cells, epithelial cells, neurons, and the blood vascular system (for review see [8]).

Clinical Application

Clinical applications of LLLT show some potential effectiveness in treating soft-tissue injury [10], chronic pain [11], and wound healing [12]. Other studies have failed to demonstrate similar effectiveness [13, 14]. Optimal wavelength, dose, dose-rate effects, tissue penetration, the role of coherence and peak power, and repetition rates for the different applications are still unknown in clinical use of these lasers. Earlier studies have shown that LLLT can prevent neuronal degeneration, promote improved neuronal function and repair, and enhance neural growth [15]. Based on these studies, LLLT was proposed for treatment of tinnitus and sensorineural hearing loss more than one decade ago [16].

Use of LLLT to Treat Tinnitus

Low-level laser radiation has been tried for treatment of tinnitus [16–18]. For that purpose, the laser light is applied through the ear canal. It was assumed that low-intensity laser irradiation was capable of penetrating soft tissue to reach the cochlea, but it remains unclear if the intensity is adequate to affect cochlear hair cells when applied through the ear canal. Multiple scattering of the laser energy by erythrocytes and microvessels has to be taken into consideration [19]. Physical measurement performed on human petrous bones showed that only the transmeatal application provided sufficient light to reach all parts of the cochlea, whereas mastoidal application did not provide sufficient energy of light in the cochlea [20]. Several biological effects in the cochlea were assumed to occur: LLLT could possibly increase cell proliferation, synthesis of ATP and collagen; affect the release of growth factors; promote the local blood flow in the inner ear; and activate repair mechanisms in the inner ear through photochemical and photophysical stimulation of the hair-cell mitochondria (“mitochondrial energy transfer”) [18, 21]. There is some experimental support for this theory from basic science [22, 23]. However, experimental data from the ear are rare. One animal study showed suppression of the compound action potential of the eighth nerve from low-level laser irradiation [24].

Though the exact peripheral mechanism of tinnitus is still uncertain, it is generally accepted that the conscious perception of tinnitus must involve the cerebral cortex. Interestingly, a study of Siedentopf et al. [25] demonstrated functional activation after transmeatal LLLT of healthy human subjects in different auditory and non-auditory structures of the brain by means of fMRI, which indicates that LLLT may affect central mechanisms.

Clinical Studies of LLLT in Patients with Tinnitus

Few reports on laser therapy of tinnitus have been published [16–18, 26–32]. Both positive and negative effects have been reported. Different wavelengths, pulsing, dosage, target of irradiation, and treatment schedule have been used. The outcome criteria and placebo control vary among the published studies making it difficult to assess the efficacy of laser treatment for tinnitus.

Earlier studies have used a combination of intravenous application of Ginkgo biloba extracts with mastoidal laser irradiation [26–28]. The rationale for the combination was the assumption in synergistic effects, as anecdotal reports supposed that Gingko extracts should increase cerebral blood flow, accelerate oxygen supply, and therefore improve tinnitus complaints [33]. Later studies changed the target to the external auditory meatus und used the laser as a monotherapy [16, 17, 29–32]. Tinnitus improvement rates varied from 15 to 67% of patients [16]. Some studies summarized that LLLT showed no efficacy in tinnitus treatment [17, 26, 27, 30, 32]. Others found LLLT to be useful in tinnitus treatment and were encouraging for further investigations [16, 29, 31]. Clear positive results were demonstrated by Wilden and Dindinger.
[18] in a treatment study without placebo control. However, a systematic review of randomized controlled clinical trials of LLLT treatment found no statistically significant difference between laser and placebo [34]. Only Wilden and Ellerbrock [35] described improvement of hearing thresholds after LLLT in more than 80% of treated subjects, whereas other studies did not observe any significant changes in hearing thresholds [16, 17, 26, 28, 32]. Nakashima et al. [30] reported on one patient who suffered from acute hearing deterioration after the third laser irradiation. All other studies could not observe any severe complications or side effects.

**Conclusion**

As the exact treatment mechanisms remain unclear and multiple placebo-controlled clinical studies failed to demonstrate significant efficacy, further studies are needed before this treatment modality can be recommended for routine clinical use.

**References**

Chapter 94
Similarities Between Treatments of Tinnitus and Central Pain

Dirk De Ridder and Aage R. Møller

Keypoints

1. Neurobiology, pathophysiology, neuroimaging, and clinical presentation share many common aspects between neuropathic pain and tinnitus.
2. Similar treatments for pain and tinnitus exist, but pharmacological methods are more successful for treatment of pain than for treatment of tinnitus.
3. Peripheral and intracranial ablative neurosurgical treatments yield common results and complications for pain and tinnitus.
4. The most promising analogous treatments for pain and tinnitus are non-invasive and invasive methods for neuromodulation, such as various forms of brain stimulation using transcranial magnetic stimulation (TMS) and transcranial direct current stimulation (tDCS).
5. TENS, for stimulation of peripheral nerves, and neurofeedback, have beneficial effects on both pain and tinnitus.
6. Invasive neuromodulatory treatments such as cochlear implants, dorsal column stimulation/auditory brainstem implant, subthalamic nucleus stimulation, and sensory cortex stimulation are beneficial for both tinnitus and pain.

Keywords
Tinnitus • Central pain • Treatment • Cortical stimulation • Peripheral nerve stimulation • Transcranial magnetic stimulation • Transcranial direct current stimulation

Abbreviations
ABPI  Auditory brainstem implant
CI  Cochlear implant
DBS  Deep brain stimulation
DCN  Dorsal cochlear nucleus
DCSCS  Dorsal column spinal cord stimulation
DLPFC  Dorsolateral prefrontal cortex
EEG  Electroencephalography
fMRI  Functional MRI
MRI  Magnetic resonance imaging
MSI  Magnetic source imaging
MVD  Microvascular decompression
PET  Positron emission tomography
rTMS  Repetitive TMS
STN  Subthalamic nucleus
tDCS  Transcranial direct current stimulation
TENS  Transderm electric nerve stimulation
TMS  Transcranial magnetic stimulation
TMS  Trans cranial magnetic stimulation
VTA  Ventral tegmental area

Introduction

Similarities between pain and tinnitus were discussed in Chap. 14. Similarities Between Tinnitus and Pain. In this chapter, we will discuss similarities in treatment of neuropathic pain and some forms of tinnitus. Apart from the developmental and reorganizational analogy, a clear clinical analogy exists between phantom pain and tinnitus [1–4]. Both symptoms are wholly subjective sensations, events that may change in character and quality. Both can be masked and relieved by electrical stimulation with a residual inhibition. Transection
of an afferent nerve usually does not help relieve tinnitus or chronic pain. In both systems, the ascending system is modified by a descending counterpart. This leads to similar characteristic symptoms in both tinnitus and phantom pain [2–4].

A normal stimulus to the skin in individuals with phantom pain can create a painful sensation (allodynia) in the same way tinnitus patients can perceive a sound as unpleasant or painful. A painful stimulus often generates an explosive and prolonged reaction to the stimulus (hyperpathia) in individuals with phantom pain similar to the hyperacusis seen in tinnitus patients [5]. The wind-up phenomenon, a worsening of pain sensation with repeated stimuli of the same intensity, is also present in some individuals with tinnitus, where it is described as an increasingly unpleasant sensation on repeating the same sound [2, 3]. Furthermore, a feeling of anxiety, nausea, and a clear stress response is often encountered both in individuals with phantom pain and tinnitus [2, 3] (see also Chap. 14).

There are at least two distinct forms of pain: normal physiological pain, by activation of nociceptors in a normally functioning somatosensory system, and neuropathic pain, which is the result of deafferentation and activation of a hereby pathologically functioning somatosensory system. There is no physiological tinnitus that is analogous to physiological pain, and therefore there are no similarities for the treatment of tinnitus to the common analgesics that are quite efficient for acute physiological body pain. Many different kinds of medications for physiological pain are readily available and have few side effects. There are also medications for neuropathic pain. Medications such as gabapentin and pregabalin are effective in treatment of central neuropathic pain. Similar medication that has a generally beneficial effect on central tinnitus does not exist either. Several other treatments are used for neuropathic pain with varying results including other pharmacological treatments [6], epidural treatments [7], regional nerve blocks [7], destructive lesions [8], treatment with calcitonin [7], transcutaneous electrical nerve stimulation [7], motor cortex stimulation (MCS) [9–11], and thalamic stimulation [12–14]. Existing treatments for various forms of pain are far more efficient than treatment of tinnitus. So even between the pathophysiology of neuropathic deafferentation pain and deafferentation tinnitus, there have to be some fundamental differences.

### Medication

Some medications are used for both neuropathic pain and tinnitus, for example, clonazepam [15, 16] and gabapentin (and this only in acoustic trauma related tinnitus) [17]; however, most pain medication will not benefit tinnitus patients. For a detailed analysis of pharmacological approaches to tinnitus, the reader is referred to Chap. 78.

### Destructive Procedures

#### Nerve Sections

The auditory information is brought to the brain via the auditory or cochlear nerve, and feedback from the cortex to the cochlea is mediated via the vestibular nerve [18, 19]. The inferior vestibular nerve connects to the auditory nerve via a small nerve fiber bundle [20]: Oort’s bundle which contains about 360 myelinated and 1,000 unmyelinated axons [21]. As always, there is some variability, but vestibulocochlear anastomoses can be found in 80% of the population [22]. Based on this anatomical knowledge, both cochlear and vestibular nerve sections have been performed in an attempt to cure tinnitus.

In a recent review paper on vestibular nerve section performed for tinnitus [23], the proportion of patients in whom tinnitus was exacerbated postoperatively ranged from 0 to 60%, with a mean of 16.4% (standard deviation 14.0). The proportion of patients in whom tinnitus was unchanged was 17–72% (mean 38.5%, standard deviation 15.6), and in whom tinnitus was improved was 6–61% (mean 37.2%, standard deviation 15.2). In the majority of patients undergoing vestibular nerve section, ablation of auditory efferent input (and thus total efferent dysfunction) to the cochlea was not associated with an exacerbation of tinnitus [23]. Therefore, if a nerve section is elected, vestibular nerve section is to be preferred to cochlear nerve section in which the success rate in abolishing tinnitus is disappointing and the results generally unpredictable [24]; an important part of the patients (55%) report no effect or a worsening of their tinnitus [25]. Only one paper reports good results with cochlear nerve section for tinnitus [26]: two-thirds
completely relieved, 28% improved, and only 5% non-responders, without a single patient worsening (see Chap. 39).

Section of the auditory nerve is controversial and now regarded contraindicated because it involves causing deprivation of signals to the auditory system, which is known to promote plastic changes. Despite a long history of ablative procedures in neurosurgery for pain control, the evidence supporting destructive procedures for benign pain conditions remains limited to class III evidence (retrospective studies) [8]. The fact that nerve lesioning is worse than non-destructive treatments [e.g., microvascular decompression (MVD)] in pain is demonstrated in trigeminal neuralgia where MVDs are better than destructive treatments such as rhizotomies or gamma knife surgery. MVD has the highest rate of long-term patient satisfaction with the lowest rate of pain recurrence [27, 28].

After surgical removal of vestibular schwannoma with resection of the auditory nerve, most patients have a small improvement of their tinnitus, but 50% of the people who do not present with tinnitus develop it after the surgery [29].

**Frontal Lobotomies**

Tinnitus and pain distress have both been linked to a neural network consisting of the anterior cingulate, frontal cortex, and insula [30–33]. These brain areas are also implicated in the distress perceived by people with posttraumatic stress disorder [34, 35], as well as asthma-related dyspnea [36], suggesting that these areas may constitute a “general distress network”. In the 1930–1940s frontal lobotomies were performed both for pain [37, 38] and tinnitus [39, 40]. The net results of these treatments were the persistence of the perception of pain and tinnitus, but the affective component related to the pain and tinnitus disappeared. For treatment of pain, the frontal lobotomies have now been refined and restricted to anterior cingulotomies. Except for a decline in focused attention performance [41–43], other neurocognitive functions (including language, memory, motor, visual-constructional, and intellectual functions) remained unaffected after the anterior cingulotomies [43]. The decreased attention modulates (decreases) the emotional experience of pain that was related to self-perceived tension and which was expressed by anger before the treatment, which also improved mood and decreased psychasthenia [44]. Cingulotomy also reduced behavioral spontaneity, expressed as a decrease in self-initiated action [42]. When performing cingulotomies for intractable pain, 72% of patients report improvements in their pain, 55% no longer take narcotics, 67% note improvement in their family life, and 72% note improvement in their social interactions. Fifty-six percent of patients report that the cingulotomy was beneficial and 28% return to their usual activities or work [45]. No reports have been published on the use of cingulotomy for treatment of tinnitus.

**Thalamic Lesions**

Thalamic lesioning has been used for both pain and tinnitus suppression based on the idea of thalamocortical dysrhythmia [46] as unifying pathophysiological mechanism of tinnitus and pain [47]. However, the experience is very limited up to now; so no definitive conclusions can be drawn of the value of this treatment for tinnitus suppression.

**Lesioning of Autonomic Nervous System**

It is well known that the sympathetic system influences both pain and tinnitus perception [2, 3]. Both pain and tinnitus tend to worsen under stressful situations. Therefore, interfering with the sympathetic system has been performed both in pain and tinnitus [48–51]. If tinnitus responds to a stellate block, a complete suppression of the tinnitus was possible in 31%, in 50% a partial response, and in 19% no response was obtained by surgical sympathectomy [51]. In Ménière’s disease, the patients who did not improve their tinnitus intensity were no more distressed by their tinnitus [51]. The patient should be warned that 24 h after operation the deafness and tinnitus may be slightly worse, possibly as the result of irritation of the sympathetic nerve trunk; it may take a week or 10 days to settle down [51]. It can be expected, however, that cervical sympathectomies for tinnitus relief might only yield a temporary benefit, in a couple of months, similar to what is known for sympathectomies at C2 and C3 for occipital neuralgia [52].
Neuromodulation

Cortex Stimulation

The neurobiological, pathophysiological, and clinical analogies between deafferentation tinnitus and deafferentation pain [1–4, 53] suggest that the resulting phantom symptoms of central pain and central tinnitus are caused by cortical hyperactivity/reorganization. Therefore, it can be assumed that the same basic strategy for treating these two conditions can be applied.

The basic strategy can be summarized as follows:

1. The hyperactivity/reorganization that is associated with central pain and some forms of tinnitus can be demonstrated by functional neuroimaging techniques such as PET scan, fMRI, or MSI (magnetic source imaging).
2. The anatomical area of hyperactivity/reorganization can then be influenced by (neuronavigated) transcranial magnetic stimulation.
3. If successfully suppressed by TMS, an electrode can be permanently implanted extradurally over the anatomical area of cortical hyperactivity/reorganization.

The details of this approach are presented in the chapter on cortex stimulation for tinnitus (Chap. 90). In summary, a selection criterion of more than 50% transient tinnitus improvement, lasting only a few seconds, on two separated placebo-controlled TMS sessions was used for implanting cortical stimulation electrodes.

Deep Brain Stimulation (DBS)

Subthalamic nucleus (STN) stimulation is capable of both improving pain [54, 55] and tinnitus [56] in patients with Parkinson’s disease, but the mechanism is unknown. STN stimulation also modulates olfactory [57] and visual [58] function suggesting that the STN has a general modulatory action on sensory processing. Stimulation of the auditory cortex, which does not send direct projections to the subthalamic nucleus, induces only late excitatory responses in the STN via the indirect cortico-striato-pallido-subthalamic pathway [59]. Many cells in the STN respond to both motor and auditory cortex stimulation as well as to frontal cortex stimulation [59]. Therefore, it is possible that DBS of the STN improves tinnitus via its influence on the motor–auditory integration cells in the STN or indirectly via the frontal cortex. Another possibility is that it occurs via an indirect pathway involving the medial forebrain bundle. Activation of connections between the medial (limbic) STN and the medial forebrain bundle has been proposed as a mechanism for the emotional and motivational influences of STN stimulation [60]. The medial forebrain bundle connects the ventral segmental area (VTA) to the nucleus accumbens, which has been implicated in tinnitus as well [61].

Transcutaneous Electrical Nerve Stimulation and Cochlear Implants

Neuropathic pain and tinnitus are both related to deprivation of sensory input to the brain (deafferentation symptoms). One way of compensating for the effect of deafferentation is by supplying the missing information through direct electrical activation of the peripheral receptors or the sensory nerves. Electrical stimulation of the peripheral somatosensory nerves, transcutaneous electrical nerve stimulation (TENS), and the auditory nerve [cochlear implants (CI)] has been used to suppress hyperactive clinical states of the respective system, which develop as a result of the deafferentation. Neuropathic pain can be modulated by TENS [62]. The effect on pain from such stimulation of the skin or peripheral nerves is mediated by the inhibitory influence from Aβ fibers on neurons in the spinal cord that receive nocuous input from C and Aδ fibers (see Chap. 14). TENS may also affect central pain, probably through activation of neural plasticity [1].

In the auditory system, peripheral nerve stimulation is performed by CI (see Chap. 77). The use of CI for tinnitus has shown promising results with regards to tinnitus suppression [63–68]. TENS is commonly used in the treatment of pain but has been used in tinnitus as well [69–75]. TENS modulates tinnitus most likely via somatosensory–auditory interactions at the level of the cochlear nuclei [76–78] or the inferior colliculus [79] (see Chap. 9). The DCN has been implicated in the pathophysiology of tinnitus [80, 81] (see Chaps. 9 and 31), and therefore modulating its activity could be useful in some forms of tinnitus (see Chap. 31). Using
c-fos studies, it was recently shown that electrical stimulation of the skin around the ear modulates dorsal cochlear nucleus activity through both direct pathways via the trigeminal system and indirect pathways via the dorsal raphe and the locus coeruleus [82]. When auditory input to the DCN is diminished, an increase in somatosensory influence on auditory neurons occurs, which could be due to cross-modal reinnervation or increased synaptic strength [83]. This favors the use of TENS in auditory deafferentation tinnitus, even though clinical data not always support the use of TENS for tinnitus [84]. Selecting who benefits from TENS and who does not will be important for the future clinical application of this method.

**Dorsal Column Stimulation and Auditory Brainstem Implants (ABI)**

Electrical stimulation of the second neuron in the somatosensory system is known as dorsal column stimulation (DCS) and is used in the management of chronic, intractable neuropathic pain [85]. The method is based on the “gate–control” theory presented by Melzack and Wall [86], who postulated that activity in large diameter cutaneous fibers (type Aβ) inhibits the transmission of noxious information to the brain. Electrical stimulation of these large afferents by an electrode placed dorsomedially in the epidural space elicits a tingling sensation (paresthesia) in the corresponding dermatomes. To obtain successful treatment of chronic, neuropathic pain by DCS, the stimulation-induced paresthesia has to cover the anatomical areas of pain completely [87, 88].

Electrical stimulation of the cochlear nucleus in the auditory brainstem yields suppressive effects on tinnitus in 80% of patients who use their auditory brainstem implants (ABI) daily [89]. This is supportive of the theory that the DCN is critically involved in tinnitus [80, 81] (see Chap. 9).

**Transcranial Direct Current Stimulation**

Transcranial direct current stimulation (tDCS) involves stimulation by a weak constant current (between 0.5 and 2 mA) flow through the cerebral cortex via scalp electrodes. Anodal tDCS typically has an excitatory effect on the local cerebral cortex by depolarizing neurons, while the opposite occurs under the cathode electrode through a process of hyperpolarization [90]. This effect of tDCS lasts for an hour or longer after a single 20–30 min treatment session [90–93].

With the anode electrode placed over the dorsolateral prefrontal cortex, tDCS can modulate both pain [94] and tinnitus (see Chap. 89), possibly via a similar mechanism, most likely a top-down modulation of auditory [95] and somatosensory [96] processing.

For pain, cathodal tDCS stimulation of the somatosensory cortex contralateral to the side to which the pain is referred [97] and left-sided anodal tDCS over the auditory cortex [98] can influence pain and tinnitus, respectively, via a more direct effect than tDCS applied through electrodes placed on the frontal part of the scalp (anode right side, cathode left side).

**Transcranial Magnetic Stimulation**

TMS is a non-invasive method of inducing electrical current in the brain [99]. It uses a coil placed on the scalp that generates magnetic pulses of very short duration (100–300 μs) at approximately 1.5–2.0 T in strength [100]. Because magnetic fields pass largely undistorted through the scalp and skull, TMS is powerful enough to cause neuronal depolarization in the cortex. TMS originally delivered single impulses. Further development of TMS equipment allowed repetitive magnetic impulses (rTMS) to be delivered, which are more effective than single impulses. The area of the brain that is stimulated and the intensity of the electromagnetic field depend on physical properties and rapidly decrease with the distance to the coil. It was estimated that a “figure of eight coil” stimulates an area of approximately 3×2 cm at cortical surface, but the induced current falls to near zero at a depth of 3 cm [101].

TMS has been used as a putative prognostic tool for cortex implants at the auditory cortex for treatment of tinnitus [102, 104] and for implants on the somatosensory cortex [103, 105] and motor cortex [106] for treatment of neuropathic pain. Details can be found in the chapter on cortex stimulation for tinnitus (Chap. 90).

Repetitive sessions of TMS (rTMS) have also been used as a treatment for pain [107, 108] and tinnitus [101, 109–112]. Details can be found in Chap. 88.
Neurobiofeedback

Tinnitus and pain are associated with abnormally coupled low and high frequency synchronous oscillatory activity in the brain [31, 46, 113–117]. If this abnormal oscillatory activity is related to the auditory and somatosensory phantom percept, a logical attempt to treat these symptoms is by normalizing this abnormal activity. Neurofeedback is a biofeedback technique using electroencephalographic (EEG) or fMRI signals for training individuals to alter their brain activity via operant conditioning. This has been used for both tinnitus [118–119] and fibromyalgia pain [120]. A detailed description of this technique is given in Chap. 87.

A better understanding of the spectral and connectivity changes, as well as alterations in independent components in tinnitus and pain, combined with new software development for source-analyzed neurofeedback training is expected to permit this technique to become a more powerful tool in treatment of both tinnitus and pain.

Conclusion

Tinnitus does not seem to respond to medication used for physiological or neuropathic pain. This means that pharmacological treatment does not seem to benefit from the neurobiological, pathophysiological, neuroimaging, and clinical analogy between tinnitus and pain, and pharmacological treatment [122], in general, has had little success in treatment of tinnitus.

Methods such as ablative neurosurgical approaches consisting of nerve sections or intracranial destructive lesions have found use in treatment of both tinnitus and pain.

Different kinds of invasive and non-invasive neuromodulation seem to be more promising analogous treatments. For invasive stimulation implanted electrodes on the auditory and somatosensory cortex, deep brain stimulation of the subthalamic nucleus and thalamus, TENS/cochlear implants, and dorsal column stimulation/auditory brainstem implants most likely use similar mechanisms to improve pain and tinnitus. Non-invasive neuromodulation techniques such as cortical transcranial magnetic stimulation, transcranial direct current stimulation, transcutaneous electrical nerve stimulation, and neurofeedback appear to be analogous in their effect on pain and tinnitus as well.

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Chapter 95
Treatment Strategies of Temporomandibular Joint and Masticatory Muscle Disorders in Patients with Tinnitus

Ralf Bürgers, Michael Behr, and Martin Gosau

Keypoints

1. Temporomandibular joint disorders (TMD) are often accompanied by tinnitus.
2. Improvement or total remission of tinnitus from treatment of TMD is a secondary effect.
3. Treatment of TMD is suggested for patients with impaired range of jaw motion, painful jaw movements, pain in masticatory muscles or temporomandibular joint (TMJ), or oral parafunction and masticatory muscle hyperactivity.
4. Treatment of TMD includes:
   a. Intraocclusal stabilization appliances (occlusal splints);
   b. Selective adjustment of the occlusal surface of teeth and artificial dentition;
   c. Reassurance and counseling
   d. Medication, physical therapy or physiotherapy
   e. EMG biofeedback with progressive relaxation, hypnosis, and acupuncture;
   f. Joint injections (hydrocortisone), injection of local anesthetics into muscle trigger points;
   g. TMJ surgery.

Keywords Temporomandibular disorders • Tinnitus • Treatment

Abbreviations

TMD Temporomandibular disorder(s)
TMJ Temporomandibular joint

Introduction

In dentistry, the most common approach to treating patients with symptoms of temporomandibular disorders (TMD) and concomitant tinnitus targets on the therapy of the TMD, more or less independent of the severity and quality of the tinnitus symptoms [1–3]. This approach is based on the assumption that appropriate TMD treatment will also eliminate or at least reduce tinnitus symptoms [1, 4–8]. This “dental approach” to the treatment of TMD with concomitant tinnitus may be adequate for patients who suffer from tinnitus as a secondary complaint due to a primary dysfunction of the temporomandibular system. In contrast, sole treatment of TMD is inadequate in patients who suffer from TMD as an implication of the tinnitus (as occurs in individuals who process their tinnitus symptoms through nightly grinding) or in whom both TMD and tinnitus symptoms may be caused by a third “collective trigger,” such as mental pressure or a specific medication [9–11]. In such patients, sole treatment of tinnitus with an isolated TMD therapy is doomed to failure from the very start. This fact emphasizes the importance of interdisciplinary therapeutic concepts for the clinical treatment of individual patients with tinnitus.

This chapter will mainly consider such treatments of TMD that are beneficial on tinnitus when it accompanies temporomandibular problems. For general coverage of treatment of TMD, the reader is referred to the considerable literature covering this topic.
Treatment of TMD

Indication for TMD Therapy

Therapeutic intervention is beneficial to patients with TMD if one of the following symptoms is present:

1. Impaired range of jaw motion (limitation of mouth opening, deviation, or deflection)
2. Painful jaw movements
3. Pain in masticatory muscles, temporomandibular joint (TMJ), or trigger zones that occurs when palpated (myofascial pain or arthralgia1)
4. Oral parafunction2) and masticatory muscle hyperactivities (bruxism3), clenching, and rocking of teeth

Usually, TMD treatment does not benefit patients with painless TMJ sounds or symptom-free occlusal interferences. Nevertheless, conservative (non-surgical) TMD therapy might alleviate patients with TMD who also have tinnitus if the tinnitus is related to TMD symptoms (e.g., ipsilateral occurrence of TMJ clicking and tinnitus).

TMD Therapy

Therapeutic interventions for TMD are complex because of the diversity of symptoms that may be caused by many different disorders of the TMJ (arthogenic), the masticatory muscles (myogenic) or both. TMD treatment options include intraocclusal stabilization appliances (occlusal splints), selective adjustment of the occlusal surface of teeth and artificial dentition, reassurance and counseling, medication, physical therapy or physiotherapy, EMG biofeedback with progressive relaxation, hypnosis, acupuncture, joint injections (hydrocortisone), injection of local anesthetics into muscle trigger points, and TMJ surgery. However, it would be negligent to give only one general recommendation or regime for the therapy of TMD or TMD-related tinnitus [12].

Approximately 30% of patients with TMD (with myogenic or arthogenic pain) report persisting pain after TMD therapy [13]. In most patients, TMD will abate without active professional intervention [14]. Two kinds of therapy are available: conservative treatments, consisting of intraoral splints, counseling, physiotherapy, and medications; and invasive treatments (TMJ surgery), ranging from arthrocentesis and lavage to disectomy. Most patients with TMD respond to conservative treatment and such reversible therapy does not change the structures of the masticatory system [14]. Invasive treatment is irreversible and can cause minor or more extensive structural changes. In this chapter, only the most important and generally approved treatment modalities will be presented, focusing on treatment modifications for patients with TMD and tinnitus.

It has been the main goal of TMD therapy to eliminate or at least reduce pain and discomfort and achieve normalization of the mandibular range of motion [12], but attention on tinnitus in patients with TMJ problems has been increased, and improvement or total remission of tinnitus has now become an important therapeutic goal of TMD therapy.

Counseling

Patients with TMD and TMD-related tinnitus benefit from receiving background information on their disorder and its implications [15–18]. Detailed explanation of the possible correlation between tinnitus and TMD, as well as the special features of a combined therapy, can have beneficial effects on both the pain and the tinnitus [14].

Medication

Some of the medications usually prescribed for TMJ pain have been reported to cause tinnitus and hearing loss [19, 20].

Treatment of TMD-Related Tinnitus

Improvement or relief of tinnitus has been reported after TMJ surgery [21–25].

In general, no difference exists between the dental therapy of TMD and the therapy of TMD-related tinnitus. In some patients with TMD, the intensity of the
tinnitus can be modulated by mandibular movements, pressure on the TMJ, or biting (mostly by enhancement) [12, 17, 26, 27]. If the quality and severity of tinnitus change during passive and active movement of the jaw, the chances for improvement by dental therapy might be higher [17, 26].

Many patients with tinnitus, who are referred to a dentist, have chronic tinnitus. The success of treatment of TMD-related tinnitus decreases with the duration of the tinnitus [28]. While therapy of acute TMD-related tinnitus may be aimed at total elimination, or significant reduction, of the tinnitus, the likelihood of complete remission of tinnitus is small in patients who have had their tinnitus for a long time [28]. Patients with objective tinnitus related to the TMJ or masticatory muscles (clicking and crepitation) are distinguished from those with subjective tinnitus. Treatment of objective tinnitus should focus on localizing and eliminating the source of the sound. Some investigators have questioned the significance of TMJ sounds and their clinical and pathological relevance [12].

**Prognosis**

Conflicting evidence of the effect of TMD therapy on TMD-related tinnitus exists. Various studies about the effect of different TMD treatment strategies on tinnitus have been ruled out, and an astonishing high percentage of improvement or elimination of tinnitus symptoms has been reported in some investigations (see Table 95.1) [3, 6, 20, 22, 26, 29, 30, 31]. No significant differences were found between various TMD therapies (splint vs. physical therapy vs. self-observation, etc.) [28].

Few prospective controlled randomized studies on the efficacy of TMD therapy for TMD-related tinnitus have been published, and many studies are only descriptive. Most studies that included a control group found that the placebo effect on tinnitus was considerable [26]. The great individual variation in the tinnitus of patients with TMD makes it difficult to compare the stated efficacy of treatments used in different studies [12, 28, 32].

**Table 95.1 Effect of TMD therapy on tinnitus: literature overview**

<table>
<thead>
<tr>
<th>Source</th>
<th>Total no. of patients</th>
<th>TMD therapy</th>
<th>Complete remission or improvement (%)</th>
<th>Complete remission (%)</th>
<th>Improvement (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bernstein et al. [33]</td>
<td>28</td>
<td>Adjusting deflective tooth contacts, counseling</td>
<td>75%</td>
<td>21 (75%)</td>
<td></td>
</tr>
<tr>
<td>Bürgers (unpublished)</td>
<td>25</td>
<td>Occlusal splints, physiotherapy</td>
<td>(44%)</td>
<td>2 (8%)</td>
<td>9 (36%)</td>
</tr>
<tr>
<td>Bush [6]</td>
<td>35</td>
<td>Oral splint, physiotherapy, surgery, medication</td>
<td>86%</td>
<td>11 (31%)</td>
<td>19 (55%)</td>
</tr>
<tr>
<td>Dolowitz [34]</td>
<td>43</td>
<td>Muscle exercises</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gelb et al. [1]</td>
<td>26</td>
<td>Treatment “to establish maxillomandibular balance”</td>
<td>96%</td>
<td>25 (96%)</td>
<td>17 (65%)</td>
</tr>
<tr>
<td>Hankey [2]</td>
<td>6</td>
<td>Various stomatognathic treatment</td>
<td></td>
<td>(50%)</td>
<td></td>
</tr>
<tr>
<td>Ioannides et al. [35]</td>
<td>2</td>
<td>Counseling, physiotherapy, restoration of occlusion</td>
<td></td>
<td>1 (50%)</td>
<td></td>
</tr>
<tr>
<td>Kelly et al. [5]</td>
<td>46</td>
<td>Occlusal splints (cast metal overlays)</td>
<td>80%</td>
<td>37 (80%)</td>
<td>20 (43%)</td>
</tr>
<tr>
<td>Koskinen et al. [7]</td>
<td>8</td>
<td>Selective grinding, occlusal splints, thermotherapy, muscle relaxants, muscle exercises</td>
<td>63%</td>
<td>5 (63%)</td>
<td>2 (25%)</td>
</tr>
<tr>
<td>Linsen et al. [8]</td>
<td>22</td>
<td>Distraction splints</td>
<td></td>
<td>17 (77%)</td>
<td>8 (36%)</td>
</tr>
<tr>
<td>Rubinstein et al. [3]</td>
<td>57</td>
<td>Occlusal splints, occlusal adjustment, muscle exercises</td>
<td></td>
<td></td>
<td>26 (46%)</td>
</tr>
<tr>
<td>Tullberg et al. [36]</td>
<td>73</td>
<td>Counseling, jaw exercises, occlusal bite splints</td>
<td></td>
<td></td>
<td>31 (43%)</td>
</tr>
<tr>
<td>Wright et al. [29]</td>
<td>93</td>
<td>Splints, self care instructions, medication</td>
<td>86%</td>
<td>80 (86%)</td>
<td>52 (65%)</td>
</tr>
</tbody>
</table>
Summary and Conclusions

1. General TMD treatment recommendations are inadequate for patients with TMD-related tinnitus. There is no typical TMD patient and no typical TMD therapy, just as there is no typical tinnitus patient and no typical tinnitus therapy.

2. Different TMD treatment options are in use for TMD-related tinnitus, but no significant differences have been found between different TMD therapies regarding tinnitus.

3. Complete remission of tinnitus symptoms remains an unrealistic therapeutic goal. At best, the dental approach can provide an additional therapeutic option in individual patients with tinnitus. TMD treatment options contribute to inter- and multidisciplinary approaches to the clinical management of the symptoms of tinnitus.

References

11. Laskin, DM, Block, S, Diagnosis and treatment of myofascial pain-dysfunction (MPD) syndrome J Prosthet Dent 1986, 56:75–84
17. Vernon, J, Griest, S, Press, L, Attributes of tinnitus that may predict temporomandibular joint dysfunction Cranio 1992, 10:282–287
27. Peroz, I, Dysfunctions of the stomatognathic system in tinnitus patients compared to controls HNO 2003, 51:544–549
35. Ionnides, C.A., Hoogland, G.A., The disco-malleolar liga-
36. Tullberg, M., ernberg, M., long-term effect on tinnitus by treat-
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