Basic Otorhinolaryngology

A Step-by-Step Learning Guide

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Preface

This preface aims to provide you with some background information on this textbook: Who can benefit most from this book? What is the teaching approach that is used—i.e., how are the contents presented to make them easier to learn? Who are the members of the “textbook team”?

Approach

Who is the book written for? This book is both a textbook and a reference work. It is intended primarily for students, but is also written for physicians, especially those taking part in further training. True learning means understanding, and so teaching means explaining. An essential part of learning is understanding basic concepts and potentially complex interrelationships. Learning should also be interesting, and it should convey enjoyment of the material and its fascinating aspects. In this sense, this book aspires to be more than an exam review.

The capabilities of digital production technology have been used with the goal of creating an educationally compelling, graphically attractive, yet affordable textbook.

Structure: One of the main goals was to present the material in an easy-to-learn, user-friendly format. The result is a new kind of textbook in which the material is broken down into brief study units, which represent a cohesive learning unit. Subdividing the contents into manageable portions makes it possible to present thematic highlights that are usually not found in textbooks and would have been more difficult to incorporate into chapters with a traditional structure. Each study unit begins with a starter in boldface type. This states the topics that are covered in the unit and the way in which they fit into the overall scheme. Special points are noted, and the material is related to other study units. The starter is not a summary.

The topics in each study unit are presented on facing pages. For clarity, “open-book” logos are shown at the bottom corner of the right-hand page: the number of logos (from one to six) indicates the number of facing-page sets that are contained in the current study unit. The red-colored logo shows where you are in the unit.

Subject matter: This textbook conforms to the latest developments in otorhinolaryngology and head and neck surgery. All main information that is needed for the basic understanding of a topic is contained in the main text, figures, and tables.

1 Knowledge in depth

Boxes marked with this symbol provide information that goes beyond a basic understanding. This may include operating techniques, illustrative case descriptions, historical information, or repetitions from earlier study (e.g., in embryology). If you are in a hurry, you can skip the in-depth boxes and still understand the material in the main text.

Points of emphasis are meant to indicate “caution” or “take note,” and serve to direct attention to key points.

Terminology: Efforts have been made in recent years to establish a standard international nomenclature in various areas of otorhinolaryngology. The most up-to-date terms are used in the text, while older or less commonly used terms are noted as synonyms.

Fig. 1 Color code for flowcharts

[Diagram of flowchart with sections: Starting situation, Signs and Symptoms, History, observation, methods of examination, Diagnosis, differential diagnosis, Treatment, further management]
Acknowledgments

We have very much enjoyed working together on this book, and our collaboration on it has resulted in a text that is greater than the sum of the authors’ individual contributions. Ms. Richter, our project editor at Thieme Medical Publishers for the original German version of this book, made a substantial contribution to this outcome through her enthusiasm and tireless efforts in editing our manuscripts. Time and again, she pointed out areas that needed clarification, as well as passages that could be omitted. As a result, the book is her work as well as ours, and she has earned our sincere gratitude.

Terry Telger of Waco, Texas, prepared the English translation. Apart from approaching this project with his usual professionalism and tremendous skills, he revealed time and again points in need of clarification. Stephan Konny was our project editor at Thieme International for the English version. He managed the translation process, keeping in mind not only the language but the entire cultural aspects of such a process. Just as much as Ms. Richter for the German version, they earn our gratitude for the English version. We are also grateful to the publishers, Thieme, for promoting the project and fostering its development. Our thanks also go to Ms. Baum for her skillful artwork.

Special gratitude is owed to our families. A great deal of time that should really have belonged to them was spent preparing this project. Even so, they supported our lengthy work on the book with the encouragement that only a family can provide.

What Can We Improve?

Our goal was to tailor this book to meet readers’ needs. Only you, the reader, can judge whether we have accomplished this. We would therefore be delighted for you to contact us or the publishers regarding any changes that you would like to see in the next edition. We wish you much enjoyment and every success with this book.

R. Probst  G. Grevers  H. Iro

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Table 1  The textbook team

<table>
<thead>
<tr>
<th>Team members</th>
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Fig. 2  The editors

Left to right: Prof. H. Iro is the chairman of the Dept. of Otorhinolaryngology at Friedrich Alexander University Hospital in Erlangen, Germany; Prof. G. Grevers is a practicing otorhinolaryngologist in Stamberg, Germany; and Prof. R. Probst is the chairman of the Dept. of Otorhinolaryngology at the University Hospital Basle, Switzerland.
Anatomy, Physiology, and Immunology of the Nose, Paranasal Sinuses, and Face

1.1 Basic Anatomy of the Nose, Paranasal Sinuses, and Face
   - Facial Skin and Soft Tissues
   - The Facial Skeleton
   - External Nose
   - Nasal Cavities
   - Paranasal Sinuses
   - Vascular Supply
   - Nerve Supply
   - Functional Anatomy of the Ostialmeatal Unit

1.2 Morphology of the Nasal Mucosa
   - Respiratory Mucosa
   - Olfactory Mucosa

1.3 Basic Physiology and Immunology of the Nose
   - Physical Principles of Nasal Airflow
   - Conditioning of the Inspired Air
   - Protective Functions of the Nasal Mucosa
   - Speech Production
   - Olfaction
1.1 Basic Anatomy of the Nose, Paranasal Sinuses, and Face

The shape and appearance of the external nose affect not only the overall appearance of the face, but also the functional processes that take place inside the nose. The structural anatomy of the nose is important for both aesthetic and functional reasons, since the nose, as the gateway to the respiratory tract, performs a variety of physiologic functions.

Facial Skin and Soft Tissues

For the effective surgical treatment of soft-tissue defects in the face, whether of a traumatic or neoplastic nature, it is important to consider some distinctive features of the morphology and topographical anatomy of the face, since this is a highly conspicuous region in which the faulty or inadequate treatment of tissue changes will have obvious consequences. One such feature involves the tension lines of the skin (Fig. 1.1a), known also as the relaxed skin tension lines (RSTLs). Scars can be made less conspicuous by taking these tension lines into account when suturing facial skin injuries. The aesthetic units of the face are an important consideration in the treatment of larger soft-tissue defects (Fig. 1.1b). Failure to take these units into account will produce a poor cosmetic result.

The Facial Skeleton

Knowing the various components of the bony facial skeleton (Fig. 1.2) and their relationship to one another is important in trauma management and also in the diagnosis and treatment of inflammatory diseases of the facial skeleton and their complications. The upper jaw bone, or maxilla, houses the maxillary sinus and articulates laterally with the zygomatic bone (zygoma) via the zygomatic process (Fig. 1.2). The upper part of the maxilla borders the nasal bone, and its frontal process projects upward to the frontal bone. The zygoma also has a frontal process that connects superiorly with the frontal bone lateral to the orbit. The zygoma communicates posteriorly with the zygomatic arch.

External Nose

The shape of the external nose is defined by the nasal bones, a pair of rectangular bones in the upper nasal dorsum, and by the paired lateral cartilages (upper nasal cartilages) and alar cartilages (major alar cartilages) in the central and lower portions of the nose (Fig. 1.3). The lateral portions of the nasal alae also contain several small accessory cartilages, called the minor alar cartilages, which are embedded in the lateral soft tissues of the nose. The shape and stability of the alar cartilages, each of which consists of a medial and lateral crus, chiefly de-
termine the appearance of the nasal tip and the shape of the nares. As a result, they are also important in maintaining an effective nasal airway. Besides the medial crura, the inferior septal margin and the connective-tissue septum (columella) are also responsible for stabilizing the base of the nose (Fig. 1.4a). Subluxation of the inferior septal margin can also hamper nasal breathing by partially obstructing the nasal airway (Fig. 1.4b).

**Nasal Cavities**

The nasal cavities begin anteriorly at the nasal vestibule, which is bordered posteriorly by the internal nasal valve (limen nasi) located between the posterior border of the alar cartilage and the anterior border of the lateral cartilage. This valve area is the narrowest portion of the upper respiratory tract and, as such, has a major bearing on the aerodynamics of nasal airflow (see also 1.3, Basic Physiology and Immunology of the Nose, pp. 10–13). The anterior bony opening of the nasal cavity, called the piriform aperture, is bounded laterally and inferiorly by the maxilla and superiorly by the nasal bone (Fig. 1.2). The interior of the nose behind the nasal valve is divided by the nasal septum into two main cavities. The nasal septum is composed of an anterior cartilaginous part and two posterior bony parts. Abnormalities in the shape of the nasal septum (see also 3.2, Nasal Deformities, p. 30), which may consist of a deviated septum, tension septum, spurs or ridges, are a frequent cause of nasal airway obstruction. The choanae are the paired posterior openings through which the nasal cavities communicate with the nasopharynx.
The nasal cavity is bounded laterally by the lateral nasal walls, which are formed by the ethmoid bone and maxilla, and posteriorly by the palatine bone and the pterygoid process of the sphenoid bone. Several functionally important structures are located on the lateral nasal wall: the nasal turbinates and their associated passages (meati), sinus ostia, and the orifice of the nasolacrimal duct (Fig. 1.5).

The inferior turbinate consists of a separate bone that is attached to the medial wall of the maxillary sinus. The opening of the nasolacrimal duct is located in the corresponding inferior meatus (1.1). The middle and superior turbinates are part of the ethmoid bone. In rare cases, a rudimentary “supreme turbinate” is also present above the superior turbinate.

The middle turbinate has by far the greatest functional importance, because most of the drainage tracts from the surrounding paranasal sinuses open into the middle meatus (see also 1.3, Anatomy of the Ostiomeatal Unit, p. 7).

The nasal cavity is bounded superiorly by the cribiform plate of the ethmoid bone. This thin bony plate has numerous openings for the passage of the fila olfactoria and also forms the boundary of the anterior cranial fossa. The floor of the nasal cavity is formed mostly by the hard palate, which is formed in turn by the two palatine processes of the maxilla and the horizontal laminae of the palatine bone.

Paranasal Sinuses

The paranasal sinuses are air-filled cavities that communicate with the nasal cavities (Fig. 1.6). All but the sphenoid sinus are already present as outpouchings of the mucosa during embryonic life, but except for the ethmoid air cells, they do not develop into bony cavities until after birth. The frontal sinus and sphenoid sinus reach their definitive size in the first decade of life. The maxillary sinus is present at birth but remains very small until the second dentition, because the presence of tooth germs in the maxilla limit the extent of the sinuses. The maxillary sinus, frontal sinus, and anterior ethmoid cells drain into the nasal cavity through the middle meatus—i.e., below the middle turbinate (Fig. 1.5). The posterior ethmoid cells drain into the nasal cavity through the superior meatus. The ostium of the sphenoid sinus is located in the anterior wall directly above the choanae. The anatomical connections between the nasal cavity and paranasal sinuses are functionally important and play a key role in the pathogenesis of many rhinologic diseases that involve the paranasal sinuses (see also 1.3, Anatomy of the Ostiomeatal Unit, p. 7).

The maxillary sinus borders the nasal cavity laterally, and the orbital floor separates the upper part of the sinus from the orbit. Behind the maxillary sinus is the pterygopalatine fossa, which is traversed by the maxil-
1.2 Ethmoid roof and cribriform plate

The roof of the ethmoid labyrinth is formed mainly by the portion of the frontal bone that covers and closes the ethmoid cells superiorly. The ethmoid roof is continuous medially with the cribriform plate, the lateral lamina of which represents the continuation of the attachment of the middle turbinate and is very easily injured during surgical manipulations in this region (a ethmoid roof and anterior ethmoid at the level of the crista galli). The levels of the ethmoid roof and cribriform plate can vary considerably, even in the same patient, depending on the vertical extent of the lateral lamina. Computed tomography scans should be taken preoperatively to define the individual anatomy of the anterior skull base region (b–e coronal scans of the anterior skull base in a patient with conspicuous nasoethmoidal opacity caused by nasal polyps).
lary artery along with branches of the trigeminal nerve and autonomic nervous system. The floor of the maxillary sinus is closely related to the roots of the second premolar and first molar teeth. This creates a potential route for the spread of dentogenic infections, and a tooth extraction may create a communication between the oral cavity and maxillary sinus (oroantral fistula).

Superior and medial to the maxillary sinus are the ethmoid air cells—a labyrinthine system of small, pneumatized sinus cavities that are separated from one another by thin bony walls and extend posteriorly between the middle turbinate (medial border) and orbit to the sphenoid sinus. The orbital plate of the ethmoid bone, called also the lamina papyracea, forms the lateral bony wall that separates the ethmoid air cells from the orbit. Paranasal sinus inflammations can spread through this lamina to involve the orbit (orbital complications). The posterior ethmoid cells are closely related to the optic nerve. The ethmoid roof and cribiform plate (1.2) form the bony boundary that separates the ethmoid cells from the anterior cranial fossa. The surgeon who operates in this region must have a detailed knowledge of the relations of these structures to the ethmoid labyrinth.

The sphenoid sinus is located at the approximate center of the skull above the nasopharynx. Its posterior wall is formed by the clivus. It relates laterally to the cavernous sinus, the internal carotid artery, and cranial nerves II–VI, and it is very closely related to the optic canal.

The optic nerve and internal carotid artery may run directly beneath the mucosa of the lateral wall of the sphenoid sinus, without a bony covering.

The sphenoid sinus is bordered superiorly by the sella turcica and pituitary and by the anterior and middle cranial fossae.

The frontal sinus is located in the frontal bone, its floor forming the medial portion of the orbital roof. The sinus, which is highly variable in its extent, is bounded behind by the anterior cranial fossa. Inflammations of the frontal sinus can give rise to serious complications because of its close proximity to the orbit and cranial cavity (orbital cellulitis, epidural or subdural abscess, meningitis).

### Vascular Supply

The external nose derives most of its blood supply from the facial artery, which arises from the external carotid artery, and from the ophthalmic artery, which springs from the internal carotid artery. The internal nose receives blood from the territories of the external and internal carotid arteries: the terminal branches of the sphenopalatine artery, which arises from the maxillary artery, and the anterior and posterior ethmoid arteries, which arise from the ophthalmic artery. A detailed knowledge of the vascular supply is particularly important in the management of intracranial epistaxis (nosebleed), which requires vascular ligation or angiographic embolization as a last recourse (see also 3.3, Epistaxis, p. 35). The venous drainage of the facial region is handled by the facial vein, retromandibular vein, and internal jugular vein.

The regional lymphatic drainage of the face and external nose is handled mainly by the submandibular lymph nodes, while the nasal cavity is additionally drained by the retropharyngeal and deep cervical lymph nodes.

### Nerve Supply

The facial skin receives its sensory innervation from terminal branches of the trigeminal nerve that enter the facial region through the supraorbital, infraorbital, and mental foramina (Fig. 1.2). Only the skin over the mandibular angle and the lower portions of the auricle are supplied by the great auricular nerve. The facial muscles are classified as mimetic or mastica-
tory, each of these groups receiving different motor innervation. While the mimetic muscles of the face develop from the blastema of the second branchial arch (the hyoid arch) and accordingly are supplied by the facial nerve, the masticatory muscles trace their embryonic development to the first branchial arch (the mandibular arch) and are therefore supplied by mandibular nerve branches arising from the trigeminal nerve.

Functional Anatomy of the Ostialmeatal Unit

The nose and paranasal sinuses are regarded as a functional unit. Many rhinologic disorders are transmitted from the nasal cavity into the paranasal sinus system. The ostiomeatal unit is the collective term for various anatomical structures located about the middle meatus. It represents the region on the lateral nasal wall that receives drainage from the anterior ethmoid cells, frontal sinus, and maxillary sinus (1.3). It is important to know the anatomical details of this region in order to understand the pathophysiology of acute and especially chronic paranasal sinus inflammations and the surgical procedures that are used in the causal treatment of these conditions.
1.2 Morphology of the Nasal Mucosa

Besides the anatomical structure of the external nose and nasal cavity, the nasal mucosa plays an essential role in numerous functions of the nose owing to its "gateway" location in the respiratory tract (see also 1.3, Basic Physiology and Immunology of the Nose, pp.10–13). This deals with the morphologic structure of the nasal mucosa. Understanding this structure is necessary for an understanding of functional processes.

The anterior part of the nasal cavity (the nasal vestibule), like the external nose, is covered by skin composed of a multilayer, keratinizing squamous epithelium. Anterior to the head of the inferior turbinate, this keratinized epithelium gives way to a nonkeratinized squamous epithelium, a nonciliated columnar epithelium, and finally a ciliated respiratory epithelium. Along with the submucous tissue, this ciliated epithelium forms the typical mucosal lining of the nasal cavity and paranasal sinuses (Fig.1.7). A small area on the upper nasal septum, superior turbinate, and part of the middle turbinate, located adjacent to the cribiform plate, is covered by olfactory mucosa and is called the olfactory region.

Respiratory Mucosa

Epithelium

The epithelium of the respiratory mucosa is composed of ciliary cells, goblet cells, and basal cells and provides an initial, mechanical barrier against infection. The ciliary cells dominate the surface of the respiratory epithelium. Each ciliary cell has approximately 150–200 cilia, which are composed of microtubules and are interlinked by "dynein arms." This cytoskeleton of the ciliary cells and the activity of dynein, a specialized protein, enable the typical, synchronous beating of the cilia in the respiratory epithelium. This ciliary action propels a blanket of mucous secretions (from the goblet cells) and serous secretions (from the nasal glands) toward the nasopharynx, mechanically cleansing the inspired air in a mechanism called mucociliary transport (see also 1.3, Basic Physiology and Immunology of the Nose, pp.10–13). The basal cells represent the morphologic connection between the columnar epithelium and goblet cells on the one hand and the epithelial basement membrane on the other. They are distinguished from the other epithelial cell types by an increased expression of certain adhesion molecules (e.g., intracellular adhesion molecule-1, ICAM-1) and increased cytokine synthesis (e.g., interleukin 1). Besides the four cell types mentioned, the epithelium also contains immunocompetent cells, mostly CD8-positive T cells, along with smaller numbers of mast cells, macrophages, and MHC-II-bearing dendritic cells, which function as antigen-presenting cells.

Lamina Propria

The lamina propria of the nasal mucosa is separated from the epithelium by a basement membrane. Some areas of the lamina propria, especially about the inferior turbinate, show a marked preponderance of vascular structures known as venous erectile tissue or sinusesoids. They consist of thin-walled and thick-walled venous capacitance vessels, which are important not only in warming the inspired air and producing secretions but also in controlling the tumescence of the nasal mucosa. Besides the venous capacitance vessels there are capillaries and, in deeper areas, arterial vessels. The lamina propria also contains numerous nasal glands, which mainly produce a serous secretion. The immunocompetent cells in the lamina propria consist of CD4-positive T lymphocytes along with CD8-positive cytotoxic cells and suppressor cells such as CD4-/CD8-negative T lymphocytes, mature B lymphocytes, Ig-plasma cells, mast cells, and macrophages. These cellular elements demonstrate the importance of the nasal mucosa, which acts in concert with local host reactions to mediate inflammatory and allergic responses in the nose (see also 1.3, Basic Physiology and Immunology of the Nose, pp.10–13).

Nerve Supply

Finally, the nasal mucosa is endowed with a rich nerve supply. It receives its sensory innervation from the trigeminal nerve and its autonomic innervation from the pterygopalatine ganglion. The parasympathetic fibers of this ganglion induce vasodilation and stimulate the secretory activity of the nasal glands, while the sympathetic fibers produce vasoconstriction and inhibit glandular secretions.

Olfactory Mucosa

Topography: The olfactory mucosa (see 1.4 for details on structure and function) covers the olfactory region, which occupies the anterior superior part of the nasal septum and adjacent areas of the lateral nasal wall,
including the side of the superior turbinate facing the septum and part of the middle turbinate. The junction of the olfactory mucosa with the respiratory mucosa is variable in its location.

**Stimulus processing system:** Although it covers an area of only a few square centimeters, the olfactory mucosa contains between 10 and 20 million bipolar sensory cells. The olfactory sensory cells have dendritic epithelial processes as well as basal axons that pass through the basement membrane between the supporting cells and basal cells and then join into bundles that are ensheathed by Schwann cells. These axon bundles, called the fila olfactoria, pass through foramina in the cribriform plate of the ethmoid bone and enter the cranial cavity. There they unite to form the olfactory nerve and pass to the olfactory bulb in the brain, the primary olfactory center. The latter is connected via the olfactory tract to the secondary olfactory center (olfactory cortex) in the temporobasal cortex, which is responsible for the perception of smells and their association with other sensory impressions. The secondary olfactory center also has projections to the limbic system that connect with the autonomic centers in the thalamus and hypothalamus; this creates a pathway that mediates the emotional and affective phenomena that are associated with smells. The olfactory cortex has connections with the tertiary olfactory centers (including the hippocampus, anterior insular region, and reticular formation), which are believed to have polysensory associative functions.

**1.4 Olfactory mucosa**

**Microscopic anatomy:** Besides receptor cells, the epithelium of the olfactory mucosa is composed of microvilli, supporting cells, and basal cells. The lamina propria additionally contains serous glands (olfactory glands) and vessels.

The function of the microvilli and of the olfactory glands located in the lamina propria of the olfactory mucosa is not yet fully understood.

The microvilli most likely represent extra chemoreceptors in the olfactory epithelium, which perform their function along with the classic receptor cells.

As for the olfactory glands, it is assumed that the secretions from these glands, released at the surface of the epithelium, also play a role in mediating the olfactory sense. Recent studies have shown that the secretion layer on the epithelium contains a specific protein that has a high affinity for most odorous substances, and thus could facilitate or even mediate their binding to the sensory cells.
1.3 Basic Physiology and Immunology of the Nose

To understand the pathologic processes that are important in inflammatory and allergic diseases of the nose, it is necessary first to understand the physiologic functions. As the threshold of the respiratory tract in humans, the nose is of major importance in conditioning the air before it reaches the lower airways. To understand this complex process, we must know something about the physics of nasal airflow, which also affects the warming and humidification of the inspired air. Due to its exposed position, the nasal mucosa is in constant primary contact with the environment and thus with a variety of potential pathogens. As a result, the nose is equipped with a variety of defense mechanisms (mechanical defenses, specific and nonspecific immune responses). As part of the supra-glottic vocal tract, the nose also contributes to speech production (see 18.1, pp. 386–389). Finally, the nose contains the olfactory sensory cells, giving it an essential role in olfaction (see p. 13).

Physical Principles of Nasal Airflow

During inspiration, the air stream enters the nasal vestibule in an oblique vertical direction. Aerodynamically, this air is in a state of laminar flow, meaning that there is no mixing of the different air layers. When the inspired air reaches the nasal valve located between the vestibule and nasal cavity, it passes through the narrowest site in the upper respiratory tract (limen nasi). Just past the nasal valve, the cross-section of the airway becomes greatly expanded, creating a “diffuser effect” that transforms most of the laminar flow of the inspired air into turbulent flow, in which different air layers are swirled together. Besides the velocity of the air, the degree of change in airflow characteristics at this stage is very strongly influenced by the specialized anatomy of the nasal cavity, which is subject to substantial individual differences. Septal deviation and cartilaginous or bony spurs on the septum can be as significant in this regard as turbinate hyperplasia or septal perforation. To a degree, the transition from laminar to turbulent flow within the nose is functionally desirable because it slows the flow velocity of the inspired air. This prolongs its contact with the nasal mucosa, contributing to olfaction and making it easier for the nose to clean, humidify, and warm the inspired air (see below).

Nasal Cycle

The “nasal cycle” is a physiologic phenomenon marked by an alternation between luminal narrowing and widening of the nasal cavities. This alternate congestion and decongestion of the nasal mucosa is effected mainly through reactions of the venous capacitance vessels of the inferior and middle turbinates, which are regulated by the autonomic nervous system (Fig. 1.8).

Conditioning of the Inspired Air

Inspired air is warmed and humidified in the nose before reaching the lower airways. Turbulent flow and other special physical conditions promote the necessary contact of the inspired air with the nasal mucosa. Moreover, the favorable relationship between the relatively small nasal cavity and the comparatively large mucosal surface area, which is further enlarged by the turbinates, also promotes the functionally important interaction between the inspired air and the mucosa. Humidification is accomplished by secretion and transudation from the nasal glands, the epithelial goblet cells, and the vessels of the lamina propria. Temperature regulation is controlled by the intranasal vascular system and especially the venous erectile tissue, which is particularly abundant in the inferior turbinates. The temperature in the anterior portions of the nasal cavity is lower than in the posterior regions. This temperature gradient produces a gradual warm-
ing of the inspired air, while on expiration, moisture and heat are returned to the nose through condensation. The warming capacity of the nasal mucosa is so efficient that even with ambient temperatures below zero, the temperature of the inspired air is raised by 25°C on entering the nasopharynx, with a relative humidity of over 90%.

Disturbances in the conditioning function of the nose can result from age-related drying of the mucosa due to involution of the goblet cells and glands. They can also result from chronic inflammatory changes or extensive resections of the mucosa during intranasal surgery.

Protective Functions of the Nasal Mucosa

Here the protective functions of the nose are separated into two parts to facilitate learning, although in life the various defense mechanisms are interrelated and should not be thought of as separate entities.

Nonspecific Defense Mechanisms

Mechanical defenses: The most important mechanical defense mechanism of the nasal mucosa is the mucociliary apparatus, which physically cleanses the inspired air. The mucociliary transport system consists of the cilia of the respiratory epithelium and a mucous blanket composed of two layers: a deeper, less viscid “sol layer” in which ciliary motion occurs, and a superficial, more viscid “gel layer” (Fig. 1.9). The physiology of ciliary movements is described in 1.5. Disturbances of mucociliary transport can have various causes, such as increased viscosity and thickness of the periciliary sol layer, hampering ciliary movements, or changes in the viscoelasticity of the gel layer resulting in ineffectual mucus transport. Finally, various pathogenic mechanisms can produce changes in the cilia themselves, regardless of the viscosity of the mucous blanket. For example, an acute viral infection of the upper respiratory tract can lead to desquamation of the epithelium, with a loss of ciliated cells. Also, certain micro-organisms can directly affect ciliary motility by reducing the beat frequency of the cilia. Finally, ciliary dyskinesia syndromes are congenital disorders based on morphologic changes in the cilia such as absence of the dynein arms. This results in uncoordinated, dyskinetic ciliary movements that prevent effective mucus transport (see also Paranasal Sinus Inflammations).

Nonspecific protective factors: The nasal mucosa also has a number of other, nonspecific defense mechanisms in the form of protective factors in the mucous blanket (Table 1.1).

Cellular defenses: The mucosa has nonspecific defense mechanisms at the cellular level as well. The predominant phagocytic cells are neutrophilic granulocytes, monocytes, and macrophages. They are accompanied by “natural killer cells” (NK cells), which comprise a small percentage of the peripheral lymphocytes and protect mainly against viral infections of the nasal mucosa.

Specific Immune Responses

Besides the nonspecific defense mechanisms of the nasal mucosa noted above, the nose possesses a specific immune system that can be viewed as a separate immunologic unit. It is made up of the nasal mucosa itself and the lymphoepithelial tissue of Waldeyer’s ring (see below). Recent discoveries indicate that the structures of Waldeyer’s ring, especially the pharyngeal and palatine tonsils, function as inductive components that are active in the absorption, processing, and presentation of antigens, whereas the nasal mucosa itself is purely an effector organ in which, for example, foreign material is phagocytized by immunocompetent cells.

The local, specific immune system of the nasal mucosa is based on the actions of antibodies, which are responsible for the humoral immune response, and of
The plasma cells also synthesize IgM and the less common IgG. When released, the immunoglobulins (especially IgA) are absorbed by the glandular cells of the lamina propria, provided with a secretory component, and re-released as secretory antibodies (sIgA).

**Humoral immune response:** Antibodies are formed in the paraglandular plasma cells. Most notably, IgA is an immunoglobulin that is characteristic of the respiratory mucosa and therefore of the nasal mucosa. The plasma cells also synthesize IgM and the less common IgG. When released, the immunoglobulins (especially IgA) are absorbed by the glandular cells of the lamina propria, provided with a secretory component, and re-released as secretory antibodies (sIgA).

**Cellular immune response:** Representatives of the cellular immune response of the nasal mucosa include mast cells, macrophages, various polymorphonuclear leukocytes (neutrophils, basophils, eosinophilic granulocytes), lymphocytes, and the cells of the reticuloendothelial system, which occur chiefly as dendritic (Langerhans) cells in the nasal mucosa. T lymphocytes are of special importance in the control and memory functions of the immune response, while B lymphocytes can differentiate into plasma cells and thus have a key role in the humoral immune response of the mucosa in connection with local antibody production. Eosinophilic granulocytes are found mainly in association with chronic sinusitis and nasal polyps. Their granules contain cystotoxic substances that can damage tissues by the lysis of cell membranes. Basophilic granulocytes are involved in immediate allergic reactions, although the mast cells are by far the most dominant cell type in this phase. The mast cells are also chiefly responsible for histamine release in the early phase of an allergic reaction. Basophilic granulocytes (the only representatives of polymorphonuclear leukocytes) and mast cells also have a specific receptor (FcR) for binding IgE. On contact with the corresponding allergenic substance, this can incite a devastating allergic reaction that may culminate in anaphylactic shock.

The epithelial cells of the nasal mucosa also have an immune function. In particular, the adhesion molecule ICAM-1, expressed by the epithelial cells, helps to prevent viral infections by acting as a receptor for more than 90% of rhinoviruses. Finally, the endothelial cells of the blood vessels play an important role in the specific immune responses of the nasal mucosa. The vascular endothelial cells are activated by various inflammatory mediators—e.g., interleukin 1, tumor necrosis factor-α (TNF-α), and they regulate the transendothelial diapedesis of immunocompetent cells into the surrounding tissue through the expression of various adhesion molecules (Figs. 1.10, 1.11).

**Speech Production**

Various organ systems are involved in the production of voice and speech. The anatomically separate functions of the respiratory tract, glottis, supraglottic vocal tract, and central nervous system must be coordinated in order to produce a normal voice sound. The “supraglottic vocal tract” refers to the air-containing regions located above the level of the vocal cords. The rigid portions of this tract, whose condition is subject to only minor variations under physiologic conditions (e.g., due to mucosal swelling), include the nose, paranasal sinuses, and portions of the nasopharynx. Their role in articulation is most apparent under pathologic conditions. “Hyponasal speech” (rhinophonia aperta) develops when these segments of the vocal tract contribute less to sound production as a result of partial or complete nasal obstruction or mass lesions in the nasopharynx. Conversely, “hypernasal speech” (rhinophonia clausa) occurs when these segments of the vocal tract contribute more than sound production as a result of various adhesion molecules.

**Olfaction**

The human olfactory system consists of the intranasal olfactory mucosa with its specialized olfactory epithelium and associated central pathways. The sensory cells consist of bipolar receptor cells whose proximal processes join to form the fila olfactoria, which are relayed through additional neurons and are distributed to the primary, secondary, and tertiary olfactory centers (see olfactory anatomy, p. 9). From a purely functional standpoint, an olfactory impression can be received only during inspiration, and only water-soluble and lipid-soluble substances are perceived. Even subtle changes in the chemical properties of a molecule can produce a clearly perceptible difference in the quality and quantity of the olfactory impression. The precise sequence of events that are involved in olfaction is still uncertain.
It is important clinically to differentiate between olfactory disturbances and taste disorders, because the senses of smell and taste are closely interrelated. Patients often believe that they have a dysfunction of both senses, even though an olfactory disturbance is the sole cause of the complaints in more than two-thirds of cases (see Table 2.2, p. 21).
Diagnostic Evaluation of the Nose and Paranasal Sinuses

2.1 History and Clinical Examination of the Nose
   - History
   - Clinical Examination

2.2 Special Rhinologic Tests
   - Testing Nasal Patency
   - Allergy Testing
   - Olfactometry

2.3 Imaging of the Nose and Paranasal Sinuses
   - Conventional Radiographs
   - Computed Tomography (CT)
   - Magnetic Resonance Imaging
   - Ultrasound
2.1 History and Clinical Examination of the Nose

Aside from special rhinologic tests, which are reviewed in 2.2 (see p. 19), the specific rhinologic history and clinical examination of the nose play a key role in further diagnostic and therapeutic decision-making.

History

Before the examiner asks about specific rhinologic symptoms, patients should be given an opportunity to describe their complaints “in their own words,” as in any history. The history should begin with questions about general, relatively nonspecific symptoms such as obstructed nasal breathing and nasal discharge. It is important to determine, for example, whether the nasal obstruction (“stuffy nose”) has been present for some time or is of recent onset, possibly in connection with trauma to the nose. Additional questions should elicit whether the complaints are unilateral, bilateral, or alternate between the sides and whether they are seasonal or present year-round.

In patients with nasal discharge, the consistency of the secretions should be assessed: is the discharge watery, mucopurulent, or blood-tinged (which may suggest a tumor)? To exclude allergic rhinitis, the patient should be questioned about sneezing attacks, itchy eyes (conjunctival irritation), cough, and respiratory complaints (evidence for allergic involvement of the lower respiratory tract).

If the history suggests that the disease may have an allergic cause, a specific allergy history should be taken. This includes the family and personal history (bronchial asthma, atopic dermatitis, food allergies) as well as details on the household and occupational environments, giving particular attention to pets, indoor plants, and potential allergen exposure at the workplace (e.g., in a bakery or hair salon).

Headaches may signify an accompanying paranasal sinus inflammation. Dryness of the nasal mucosa is a common finding in colds but can also result from changes in air quality, previous nasal surgery, or the chronic use of vasoconstricting nose drops or sprays that contain corticosteroids. Olfactory dysfunction is another possible symptom of rhinologic diseases, and the patient should always be questioned about this.

Clinical Examination

Inspection

The clinical examination begins with a visual inspection. Findings such as mouth breathing may direct the examiner to suspect nasal airway obstruction. The shape of the external nose may suggest intranasal abnormalities (e.g., a cartilaginous nasal deviation with a tension septum). It is particularly important to evaluate the nasal base (see Fig. 1.4), for which the patient’s head should be tilted back. In this position, the examiner can also test the stability of the nasal alae. If the alar cartilages are too soft, they will be indrawn even during normal, unforced inspiration. Skin changes such as erythema or swelling can occur with orbital complications of paranasal sinus inflammations (erythema and swelling of the upper and lower lids), in erysipelas (“butterfly”-shaped erythema of the midfacial skin), or with nasal furuncles, which present with circumscribed redness and swelling in the nasal vestibule.

Palpation

Palpation is most useful for detecting bony discontinuities. In patients with suspected neuralgias, it is also done to check for tenderness over the supraorbital, infraorbital, or mental foramina. In patients with a recent trauma history, palpation of the external nose will disclose any mobility or crepitus suggesting a fracture of the nasal pyramid. The midfacial bones (especially the bony orbital rim) are also palpated to check for step-offs indicating a fracture line. Soft-tissue swelling can limit the accuracy of this examination, however.

Anterior Rhinoscopy

The rhinologic examination itself begins with anterior rhinoscopy to evaluate the nasal vestibule and the anterior portions of the nasal cavity (Fig. 2.1).

Technique: The examiner holds the nasal speculum in the left hand and braces the index finger on the patient’s right nostril. The speculum is inserted into the nose with the blades closed. During the examination, the physician uses the right hand to position the pa-
tient’s head and gently opens the speculum to spread open the nostril to allow inspection of the nasal cavity. The speculum should not be opened too far, as this would cause discomfort. The head should be tilted slightly forward for evaluating the nasal floor, inferior turbinate, and the anterior portions of the septum. The head is tilted backward to obtain a limited view of the middle meatus and middle turbinate. Often this region cannot be adequately assessed by anterior rhinoscopy alone due to anatomical constraints. As a result, endoscopy is commonly used to examine this region as well as the posterior portions of the nasal cavity and the nasopharynx (see below). When anterior rhinoscopy has been completed, the speculum is carefully withdrawn with the blades slightly open to avoid avulsing hairs from the nasal vestibule.

In many cases the nasal mucosa should be decongested with vasoconstrictors prior to the examination, as this makes it easier to examine the interior of the nose. At the same time, it is also important to assess the “original” condition of the nasal mucosa, and so the nose should be examined before and after decongestion of the mucosa.

**Indication:** Anterior rhinoscopy is used not only for nasal examination but also for minor therapeutic procedures such as intranasal packing for epistaxis, foreign-body removal, and polypectomy.

**Children:** Smaller instruments (pediatric specula) are available for anterior rhinoscopy in children. Aural specula can also be used to examine the nose in infants or small children.

Decongestants should always be properly diluted when used in children.

---

**Posterior Rhinoscopy**

Posterior rhinoscopy was formerly done to evaluate the nasopharynx and posterior nasal cavity (choanae, posterior ends of the turbinates, posterior margin of the vomer). With the establishment of endoscopic examination techniques in rhinology, this procedure, which requires special patient cooperation, is now considered obsolete.

**Nasal Endoscopy**

Nasal endoscopy has become the most important and rewarding clinical examination method in rhinologic diagnosis.

**Prerequisites:** Nasal endoscopy requires practice because, unlike anterior rhinoscopy, it provides only close-up views of small intranasal areas. Besides rigid endoscopes, which are available in 4-mm and 2.8-mm diameters and assorted viewing angles (e.g., 0°, 30°, 120°), flexible endoscopes are also available for inspecting the nose and nasopharynx and exploring all of the pharynx and larynx in one sitting. Their main disadvantages compared with rigid scopes are their weaker light intensity and poorer image resolution. Also, it takes two hands to operate a flexible endoscope, while a rigid scope leaves one hand free for manipulating instruments.

The patient is seated for the examination (Fig. 2.2). As in anterior rhinoscopy, the preparations include decongestion of the nasal mucosa. A topical anesthetic should also be applied. Diagnostic nasal endoscopy is performed with a 4-mm 30° telescope. The 2.8-mm scope is used only in a very narrow nasal cavity or in children.
Technique: First the examiner advances the endoscope into the nasopharynx (Fig. 2.3) and inspects the eustachian tube orifice, torus tubarius, posterior pharyngeal wall, and roof of the nasopharynx (Fig. 2.4). While the transnasal nasopharyngeal inspection can provide very detailed views (e.g., for early detection of nasopharyngeal cancer), it should still be supplemented by transoral postrhinoscopic endoscopy (see 5.2, p. 104).

Nasal endoscopy is particularly useful for evaluating the ostiomeatal unit (see 1.3, p. 7), as this pathophysiologically important region generally cannot be adequately evaluated by anterior rhinoscopy alone. To inspect the middle meatus, the endoscope is first advanced toward the head of the middle turbinate. This should provide a good overview of the middle meatus (Fig. 2.5).

To advance farther into the ostiomeatal unit, the scope must negotiate the narrow passage between the uncinate process and the middle turbinate (asterisk in Fig. 2.5). Normally, this can be done only with a narrow-gauge scope (2.8 mm). The 4-mm endoscope can be used at this site only in patients who have had previous intranasal sinus surgery with resection of the uncinate process.

Direct endoscopic inspection of the paranasal sinuses is possible only to a limited degree. In some cases, the sphenoid sinus can be examined with a thin telescope passed through the natural ostium in the anterior sinus wall. If endoscopic exploration of the maxillary sinus is required (e.g., for a suspected tumor), it can be done either through the inferior meatus after perforating the lateral nasal wall or by a transfacial approach with incision of the maxillary sinus mucosa and perforation of its anterior wall.
2.2 Special Rhinologic Tests

While the examination methods described thus far are practiced routinely, special rhinologic test procedures are carried out only if there is specific evidence that suggests a particular disorder.

Testing Nasal Patency

Simple methods can be used for the preliminary assessment of nasal patency. One such method is to hold a reflective metal plate under the nose; the degree of fogging will give a crude impression of the patency of the tested nasal cavity. Nasal patency in infants can be tested subjectively by holding a wisp of cotton in front of each nostril.

Today the most standardized procedure for the assessment of nasal patency is **active anterior rhinomanometry** (Fig. 2.6). This procedure measures and graphically records the difference in pressure ($\Delta P$) from the naris (P2) to the nasopharynx (P1) and the respiratory air volume per unit time (V). One nostril is occluded for this test while the nasal air stream is measured on the opposite side. The accuracy of this test is most limited in patients with severe nasal airway obstruction, and the test cannot be performed when one nasal cavity is completely obstructed.

**Acoustic rhinometry** is described in 2.1. The differential diagnosis of nasal airway obstruction is outlined in Table 2.1.

Allergy Testing

While the history and nasal endoscopic findings can provide initial, relatively nonspecific evidence of an allergic etiology for rhinitis, allergy testing is used to verify and differentiate this condition. Various in-vivo and in-vitro methods are available for allergy testing.

Skin Tests

When a small amount of allergen is placed in contact with the skin, it can evoke a local or systemic (!) allergic reaction in a previously sensitized individual. The most widely used method is the **prick test**, in which the skin is superficially pricked with standard test substances that contain the suspicious antigens. The local skin reaction is compared with the reaction to a simultaneously applied positive control (histamine solution) and negative control (saline solution).

A positive skin prick test proves that sensitization has occurred but does not prove an allergic etiology for the rhinitis.

2.1 Acoustic rhinometry

Acoustic rhinometry is a measuring technique that is based on the principle of acoustic reflection and can be used to determine intranasal cross sections. Unlike rhinomanometry, it does not measure dynamic respiratory function but the cross sections of the nasal cavity at various sites, which are averaged together. The main advantages of this method over rhinomanometry are that it is faster and easier to perform and does not depend on patient cooperation. While these features are desirable in the examination of pediatric patients, it should always be considered that acoustic rhinometry measures static parameters and, unlike rhinomanometry, does not assess the patency of nasal airflow.

Table 2.1 Differential diagnosis of nasal airway obstruction

<table>
<thead>
<tr>
<th>Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute and chronic rhinitis (e.g., allergic, atrophic)</td>
</tr>
<tr>
<td>Sinusitis</td>
</tr>
<tr>
<td>Deviated septum (congenital, acquired)</td>
</tr>
<tr>
<td>Nasal pyramid fracture</td>
</tr>
<tr>
<td>Septal perforation</td>
</tr>
<tr>
<td>Nasal polyps</td>
</tr>
<tr>
<td>Cephalocele</td>
</tr>
<tr>
<td>Adenoids</td>
</tr>
<tr>
<td>Tumors of the nose, paranasal sinuses, and nasopharynx</td>
</tr>
<tr>
<td>Foreign bodies (especially in small children)</td>
</tr>
<tr>
<td>Drugs</td>
</tr>
<tr>
<td>Adverse effects: oral contraceptives, antihypertensive agents (e.g., reserpine, propranolol, hydralazine), antidepressants (e.g., amitriptyline)</td>
</tr>
<tr>
<td>Drug abuse: imidazoline derivatives (e.g., oxymetazoline hydrochloride, xylometazoline hydrochloride)</td>
</tr>
</tbody>
</table>

Serologic Tests

The total immunoglobulin E (IgE) assay—e.g., paper radioimmunosorbent test (PRIST)—can be used for the quantitative determination of nonspecific total IgE, and various tests are available for specific IgE determination—e.g., radioallergosorbent test (RAST), enzyme allergosorbent test (EAST), etc. Specific IgE testing is recommended because of the low sensitivity and specificity of the total IgE assay.
Nasal provocation test: This test is of greatest value in allergic rhinitis, as it is the only method in which a specified allergen is placed in direct contact with the nasal mucosa. The technique involves the selective application of an allergen solution to the head of the inferior turbinate. Rhinomanometry (see above), performed before and 20 minutes after application of the allergen, confirms the local allergenic effect of the test substance by showing a significant reduction of nasal patency due to reactive mucosal swelling.

Since provocative testing involves placing the allergen directly on the turbinate, it may incite a severe allergic response or even anaphylactic shock, and proper emergency equipment should be easily accessible in the examination room.
Olfactometry

Complaints of olfactory dysfunction become more frequent with ageing. They may be caused by a variety of underlying disorders (Table 2.2).

Because it is difficult for laypeople to differentiate between smell and taste, many patients initially complain of a taste disturbance when they actually have an olfactory disturbance.

Pure taste disorders are very rare. The relationship between taste and smell must be considered during the diagnostic work-up, and therefore both sensory modalities should be tested (see also 4.2, Taste Testing, p.76). After a detailed history has been taken, the patient is examined by anterior rhinoscopy or endoscopy to rule out anatomical and functional obstructions of nasal airflow to the olfactory groove.

Subjective Olfactory Testing

In subjective olfactory testing, various substances are held separately in front of each nostril before and after decongestion of the nasal mucosa. Several types of test substance are used: pure odorants that stimulate only the olfactory nerve (coffee, cocoa, vanilla, cinnamon, lavender), odorants with a trigeminal component (menthol, acetic acid, formalin), and substances that also have a taste component (chloroform, pyridine). Patients with a complete loss of smell (anosmia) cannot perceive pure odorants but can at least sense or taste the other substances.

Malingering should be suspected in patients who deny the perception of trigeminal stimulants.

Testing can also be performed with ready-to-use test kits (e.g., Sniffin’ Sticks). It is likely that these standardized tests will increasingly supplant the classic subjective smell tests, owing to their better reproducibility.

Objective Olfactory Testing

Objective olfactory testing is far more costly and is generally performed only at large centers. Pure odorants and trigeminal nerve stimulants are presented separately to the patient, and the responses are measured by the computer-controlled recording and analysis of olfactory evoked potentials. Objective olfactometry is used mainly in disability examinations.

<table>
<thead>
<tr>
<th>Table 2.2 Causes of olfactory disturbances</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Classification</strong></td>
</tr>
<tr>
<td>Transport of odorants</td>
</tr>
<tr>
<td>Nasal obstruction</td>
</tr>
<tr>
<td>Scar tissue occluding the olfactory groove</td>
</tr>
<tr>
<td>Perception: damage to the olfactory epithelium caused by:</td>
</tr>
<tr>
<td>Toxic substances</td>
</tr>
<tr>
<td>Drugs</td>
</tr>
<tr>
<td>Viral infections</td>
</tr>
<tr>
<td>Radiotherapy (rare)</td>
</tr>
<tr>
<td>Stimulus conduction and processing</td>
</tr>
<tr>
<td>Avulsion of fila olfactoria</td>
</tr>
<tr>
<td>Aplasia of the olfactory bulb (rare)</td>
</tr>
<tr>
<td>Injury to olfactory centers</td>
</tr>
<tr>
<td>Neurodegenerative diseases</td>
</tr>
<tr>
<td>Olfactory hallucinations</td>
</tr>
</tbody>
</table>
2.3 Imaging of the Nose and Paranasal Sinuses

Imaging procedures are an important tool in the diagnostic work-up of rhinologic diseases. Besides conventional sinus radiographs, the most important imaging modalities at present are computed tomography and magnetic resonance imaging.

Conventional Radiographs

Indications

Standard paranasal sinus radiographs in the occipitomental projection (Fig. 2.7a, b; Water projection) and occipitofrontal projection (Fig. 2.7c, d; Caldwell projection) are still routinely obtained, particularly in cases of acute inflammation. They are also obtained to evaluate midfacial fractures.

Diagnostic Value

The value of sinus radiographs is inherently compromised by the presence of superimposed structures. If previous surgery has been done on the paranasal sinuses, roentgen interpretation is further hampered by scar tissue, which can mimic sinus opacity. It is sometimes difficult to evaluate the sphenoid sinus in the occipitomental projection. If there is a high index of suspicion for sphenoid sinus involvement, a lateral sinus projection should be added to the study. The craniocaudal extent of the frontal and maxillary sinuses can also be evaluated with this technique.

Computed Tomography (CT)

Indications

Besides an occasional malformation, the main indications for CT scanning of the nose and paranasal sinuses are chronic sinusitis, trauma (especially frontobasal fractures), and tumors. CT sinus scans are compromised by metal-bearing dentures, which cause beam-hardening artifacts that can significantly degrade the image quality.

Scan Planes

Computed tomography can provide nonsuperimposed primary images of the paranasal sinuses in coronal (Fig. 2.8) and axial planes of section (Fig. 2.9). Sagittal images can be reconstructed secondarily from the axial or coronal scans, but they are of poorer quality. Coronal planes are used mainly for CT examination of the paranasal sinuses, and axial scans can be added for special investigations.

Fig. 2.7 Standard radiographic projections of the paranasal sinuses

a,b The occipitomental projection demonstrates the maxillary sinus and gives a limited view of the sphenoid sinus.

c,d The occipitofrontal projection is better for evaluating the ethmoid cells and frontal sinus.
Scan Acquisition

Scans can be acquired using the sequential, single-slice technique (conventional CT) or a continuous spiral technique (spiral or helical CT). The advantages of spiral CT are complete coverage with no interslice gaps (“volume scan”) and a shorter examination time (about 20 seconds), making the images less susceptible to respiratory and motion artifacts.

Documentation

CT images documented on radiographic film should occupy the whole frame, displaying only the structures that are relevant for making an interpretation.

Interpretation

Normally aerated paranasal sinuses exhibit air density on CT scans—i.e., they appear black. The normal mucosal lining of the sinuses is not visualized. The bony sinus walls appear hyperdense (white).

Magnetic Resonance Imaging

Indications

Magnetic resonance imaging (MRI) has fewer indications than CT in patients with paranasal sinus disease. This is primarily because MRI is markedly inferior to CT in defining the bony boundaries of the sinuses. The strength of MRI lies in its superior soft-tissue discrimination (Fig. 2.10). MRI is indicated in diseases...
that involve the paranasal sinuses in addition to the cranial cavity or orbit (e.g., tumors and congenital malformations such as encephaloceles). It can also supply information that is useful in differentiating soft-tissue lesions within the paranasal sinuses (mucoccele, cyst, polyp), and it can distinguish between solid tumor tissue and inflammatory perifocal reaction.

**Contraindications**

Before ordering an examination, the physician should consider the basic physical principle of MRI—i.e., the utilization of magnetic fields and radiofrequency energy.

At present, MRI is contraindicated in most patients with electrically controlled devices such as a cardiac pacemaker, insulin pump, cytostatic pump, or cochlear implant. By contrast, modern internal fixation materials such as titanium are usually nonmagnetic and therefore MRI-compatible.

**Method**

The standard imaging protocol employs a $T_1$-weighted spin-echo sequence before and after intravenous contrast administration in addition to a proton- and $T_2$-weighted turbo spin-echo sequence.
Slice Thickness

The slice thickness should not exceed 3–4 mm, and the slice increment should be no greater than 0.6 mm. Imaging of the frontal skull base, orbit, parapharyngeal space, and pterygopalatine fossa requires the highest possible spatial resolution with a thin slice thickness (3 mm).

Imaging Planes

Primary scan acquisition in MRI can be done in three planes: axial, coronal, and sagittal (Fig. 2.10). Plain, unenhanced T1-weighted images are excellent for defining normal craniofacial anatomy.

Ultrasound

The paranasal sinuses can also be visualized with ultrasound (A and B mode).

Advantages: Ultrasound is particularly useful in the follow-up of acute inflammatory processes, as it can eliminate the need for extra radiographic views. It is also used in children and pregnant women for the same reason.

Disadvantages: Ultrasound yields much less detailed images than CT and MRI, and it cannot provide three-dimensional rendering.

Indications

The frontal and maxillary sinuses are most easily accessible to ultrasound imaging. The anterior ethmoid cells can be scanned via the medial canthus of the eye, but it should be added that these cells can be examined from this site only by using a small A-mode transducer or a more costly, specialized B-mode transducer; a large linear array (7.5 MHz) cannot be used. Scanning the middle and posterior ethmoid cells by the transocular route is extremely challenging and requires a highly experienced examiner. The sphenoid sinus is inaccessible to ultrasound imaging because of its location.
# Diseases of the Nose, Paranasal Sinuses, and Face

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- Malignant Tumors 64
3.1 Malformations of the Nose, Paranasal Sinuses, and Face

Malformations involving the nose may be caused by developmental abnormalities of the nasal floor, palate, nasal roof, and internasal region. A variety of disorders can result, depending on the affected anatomic structures.

Choanal Atresia

Epidemiology: Choanal atresia has an incidence of one in 5000 to one in 10,000 births and is more often unilateral than bilateral. The atresia is bony in 90% of cases and membranous in only 10%.

Symptoms: Bilateral choanal atresia is an acutely life-threatening emergency because the neonate, except when crying, is an obligate nasal breather until about the sixth week of life. As a result, the infant experiences episodes of asphyxia at rest when its mouth is closed, especially during periods of sleep, and also during feeding. The resulting hypoxia is manifested by cyanosis, bradycardia, and an erratic respiratory rate with the mouth open or closed. Cyanosis that is present at rest and improves with exertion is called paradoxical cyanosis because of its opposite pattern relative to cyanosis with a cardiac cause.

Unilateral choanal atresia may be manifested by a purulent nasal discharge on the affected side. Choanal atresia may be associated with various other anomalies, with fully developed cases presenting as the CHARGE syndrome (coloboma; heart disease; atresia of the choanae; retarded growth, development and/or central nervous system anomalies; genital hypoplasia; ear anomalies or deafness).

Diagnosis:

Both choanae in newborns should be routinely catheterized in the immediate postnatal period (e.g., with the suction catheter) to exclude choanal atresia.

The clinical suspicion of choanal atresia can be confirmed by examination with a rigid or flexible endoscope (Fig. 3.1).

Treatment: The acute care of choanal atresia in asphyxia consists of intubation followed by perforation of the atresia plate. Recurrent stenosis is prevented by inserting a stent and securing it with a suture (to prevent aspiration). The definitive surgical repair of bilateral choanal atresia is performed during the first weeks or months of life. Surgery for unilateral atresia can be postponed until school age, when the anatomy of the region is more similar to that encountered in adults.

Frontobasal Dysraphias

The incidence of dysraphias involving the anterior skull base is approximately one in 20,000 to one in 40,000 births. The familial pattern of occurrence suggests a genetic component to the disease. For further details on the embryology, see 3.2.

Manifestations: Congenital dysraphias of the anterior skull base can have various manifestations that include dorsal nasal fistulas (3.3), dermoids (3.4), frontonasal extracerebral gliomas, and cephaloceles (3.5).

Fig. 3.1 Choanal atresia

Postrhinoendoscopic view shows partial closure (stenosis) of the left choana accompanied by complete closure (atresia) of the right choana (contrast with normal-appearing choanae in Fig. 2.4, p. 18).
3.2 Embryology of frontobasal dysraphias
The skull base may be affected by congenital closure defects, analogous to a dermal sinus or spina bifida involving the lumbar portion of the spine. Dysraphic anomalies of the anterior skull base are caused by exposure to teratogenic agents during the second or third week of embryonic development, when the neural tube is forming from the neural plate, or during the fourth week, when the cerebral ventricles and central canal are forming and the central nervous system is separating from the epidermis and migrating to a deeper level.

3.3 Dorsal nasal fistula
Morphology: A dorsal nasal fistula consists of a fistulous tract that is lined by keratinized squamous epithelium and forms a tiny opening on the dorsum or tip of the nose. The fistula may terminate blindly or even extend into the cranial cavity, creating an open communication with the subarachnoid space.
Symptoms: Fistulas that terminate blindly are usually manifested clinically at an older age due to inflammation around the fistulous opening. If the fistula communicates with the subarachnoid space, it can lead to severe complications such as cerebrospinal fluid leakage, meningitis, or brain abscess.
Diagnosis: The diagnosis is established by computed tomography or magnetic resonance imaging. Diagnostic catheterization or contrast injection is contraindicated due to the risk of intracranial complications.
Treatment: Treatment consists of complete removal of the fistulous tract, which may include excising the dural defect and repairing it by duraplasty. Incomplete removal of the fistula will predispose to recurrent infections.

3.4 Nasal dermoid
Nasal dermoids, like dorsal nasal fistulas, are lined by keratinized squamous epithelium. Sites of predilection are the dorsal nasal midline and nasal flank, where the lesions present as cystic protrusions. A nasal dermoid may coexist with a dorsal nasal fistula in rare cases. Abscesses may develop as an inflammatory complication. Diagnosis and treatment are the same as described in 3.3.

3.5 Cephalocele
Cephaloceles are herniations of intracranial contents through a bony defect in the skull. Several types are distinguished according to the structures involved: meningocele (congenital protrusion of the leptomeningeal), meningoencephalocele (leptomeningeal and brain tissue), and meningoencephalocystocele (leptomeningeal plus portions of the ventricular system).
Etiology: Most cephaloceles are congenital, but rare cases are post-traumatic (e.g., after a frontobasal fracture).
Classification: Cephaloceles of the anterior skull base are classified into two groups. Sincipital cephaloceles are located near the glabella, forehead or orbit, while basal cephaloceles are found mainly in the nasal cavity or nasopharynx.
Presentation: Most cephaloceles are manifested clinically during childhood. The sincipital forms appear as a pulsating mass near the glabella, often associated with a broad nasal dorsum and hypertelorism. Fig. a illustrates these features in a girl with an extensive meningocele. The basal forms present as an intranasal mass, typically with associated nasal airway obstruction. They closely resemble intranasal polyps and should be considered in the differential diagnosis of children with suspected nasal polyps, which are rare in this age group.
Diagnosis: Computed tomography (CT) and magnetic resonance imaging (MRI) can supply information on the location and extent of the mass and the associated bony defect. In the case shown in Fig. a, coronal CT with a bone window setting (Fig. b) defines the extensive bone defect in the ethmoid roof (arrows), while CT with a soft-tissue window (Fig. c) more clearly distinguishes the cephalocele from adjacent structures.
Treatment: Treatment is always surgical and consists of removing the cephalocele and repairing the dural defect. Any associated anomalies of the orbit and facial skeleton should also be corrected.
3.2 Nasal Deformities

A basic distinction is drawn between deformities of the external nose and intranasal deformities. They are frequently combined, however, as deformities of the external nose are generally associated with a variable degree of nasal septal curvature and may even be caused by them (e.g., cartilaginous nasal deviation with a deviated septum, humped nose with a tension septum). For learning purposes, however, septal deviation is classified as an intranasal deformity and is described separately from the various external deformities.

**Septal Deviation**

**Definition:** A congenital or traumatically acquired bending or bowing of the nasal septum.

**Symptoms:** Almost everyone has some degree of bowing, spurring, or ridging of the cartilaginous or bony nasal septum. Mild forms do not cause symptoms and have no pathologic significance (Fig. 3.2a).

More pronounced degrees of septal curvature can obstruct nasal breathing and may also cause olfactory impairment due to inadequate ventilation of the olfactory groove. Deficient nasal airflow can also lead to paranasal sinus sequelae such as headaches and recurrent sinusitis. A large septal spur that comes into contact with the nasal turbinates can cause epistaxis (Fig. 3.2b).

**Diagnosis:** Septal subluxation is a special form in which the anterior septal margin is displaced from the median plane (Fig. 1.4b, p.3). This condition is readily identified by external inspection of the nasal base. Further clinical examination consists of anterior rhinoscopy or endoscopy (Fig. 3.2a, b), which can verify the morphologic changes in the nasal septum. The degree of nasal obstruction can be objectively evaluated by rhinomanometry (see p.19). For medico-legal reasons, olfactory testing should always be done prior to surgical treatment (see p.21).

**Treatment:** The treatment of choice is surgical straightening of the deviated septum (septoplasty). This procedure involves removing the deviated cartilaginous and bony portions of the septum along with any spurs and ridges and reimplanting them as needed until the septum occupies a tension-free position in the median plane.

The indication for septoplasty is basically any septal deviation that is causing subjective complaints with functional impairment of nasal breathing.

The indication for septoplasty should be weighed very carefully in children and adolescents under 15–17 years of age, and a very conservative approach should be taken in patient selection. Indiscriminate use of this procedure in younger patients can damage the growth zones of the septum, causing long-term problems.
Deformities of the External Nose

**Causes and forms:** Deformities may be congenital or traumatically acquired. Virtually any bony and cartilaginous structures of the external nose may be affected. An accompanying septal deviation is present in many cases. The most common deformities are a crooked nose, humped nose, saddle nose, and broad nose, which may occur separately or in combinations (Fig. 3.3).

**Diagnosis:** Besides inspection of the external nose, in which the affected cartilaginous and bony structures are identified, the diagnostic workup should include anterior rhinoscopy or endoscopy to evaluate the shape and position of the nasal septum.

Photographic documentation should always be obtained preoperatively for medicolegal reasons.

**Treatment:** Since deformities of the external nose are frequently associated with intranasal changes, most cases have both a functional and an aesthetic indication for corrective surgery.

The treatment of choice is “functional septorhinoplasty,” with correction of the nasal septum and external nose. In most cases the bony nasal skeleton has to be osteotomized at multiple sites in order to achieve the desired nasal shape and position. The humped nose additionally requires dorsal hump removal. Saddle nose is corrected by filling the dorsal concavity with a cartilage graft taken from the septum, auricle, or rib.

The preoperative appearance of various nasal deformities is shown on the left side, the postoperative appearance on the right side.
3.3 Nosebleed (Epistaxis)

Nosebleed is a relatively common, usually harmless symptom that may reflect a number of diseases of variable severity. By knowing the potential causes, the physician can react appropriately in threatening cases.

Causes

Nosebleed may have a local or systemic cause. Possible local causes (Table 3.1) include mucosal hyperemia due to an acute inflammation (rhinitis), allergies, and ambient conditions that dry the mucosa, increasing the fragility of the intranasal vessels (e.g., air conditioning). Local manipulations (nose picking) can also cause a nosebleed, usually in Kiesselbach’s area (a richly vascularized area of septal mucosa at the junction of the nasal cavity and vestibule, Fig. 3.4). Other possible local causes of epistaxis are congenital or acquired abnormalities of the nasal septum, such as pronounced septal spurs or ridges (see Fig. 3.2b, p. 30). Finally, nosebleed can result from a septal perforation (Fig. 3.5). A perforated septum can have several causes, including a septal fracture with a superinfected septal hematoma (septal abscess), autoimmune disease (e.g., Wegener granulomatosis, see 3.10, p. 50), or a previous septoplasty (see p. 30) leading to mucosal perforation and cartilage necrosis.

Epistaxis may also be symptomatic of an underlying systemic disease (Table 3.2). Besides vascular and circulatory diseases, typical examples are the various forms of hemorrhagic diathesis (e.g., Osler disease, see 3.7, p. 35), infectious diseases, and endocrinopathies.

Diagnosis

Nosebleed requires a simultaneous, coordinated protocol of diagnostic and therapeutic actions. One possible algorithm is shown in Fig. 3.6.

The diagnostic work-up begins with blood pressure measurement. Except in very minor cases, the Hb should also be determined, and a coagulation disorder should be excluded by determining the platelet count, bleeding time, thromboplastin time (formerly: Quick), partial thromboplastin time (PTT), and thrombin time.

Bleeding site: The nasal cavity is inspected by anterior rhinoscopy or endoscopy following decongestion and local anesthesia of the mucosa. In most cases the bleeding site is in Kiesselbach’s area (Fig. 3.4). It can be difficult to locate the bleeding source, however, when there is profuse bleeding from the posterior parts of the nasal cavity, which are less accessible to inspection.

Table 3.1 Local causes of epistaxis

<table>
<thead>
<tr>
<th>Classification</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Change in the nasal septum</td>
<td>Perforation (Fig. 3.5): traumatic, iatrogenic, inflammatory; spurs or ridges (Fig. 3.2b, p. 30)</td>
</tr>
<tr>
<td>Mucosal or vascular injury</td>
<td>Foreign bodies, rhinoliths, trauma (including nose picking), allergy, acute rhinitis, traumatic aneurysm of the internal carotid artery (very rare)</td>
</tr>
<tr>
<td>Neoplasia</td>
<td>Benign and malignant neoplasms of the nose, paranasal sinuses, and nasopharynx</td>
</tr>
<tr>
<td>&quot;Idiopathic&quot;</td>
<td></td>
</tr>
</tbody>
</table>
**Treatment**

**General measures:** The intensity of the bleeding and risk of aspiration can be reduced before the cause and location have been established. The nostrils are compressed against the nasal septum, and the patient is told not to swallow blood running down the pharynx. The patient is kept in an upright posture to reduce blood flow to the head and inhibit the swallowing of blood. An ice bag can be placed on the back of the neck to induce reflex vasoconstriction (see Fig. 3.7). An intravenous line should be placed if bleeding is severe.

**Silver nitrate cautery:** Mild epistaxis from Kiesselbach’s area can often be controlled by selective local cauterization of the bleeding site with silver nitrate. Opposing sites on the nasal septum should not be cauterized due to the risk of septal perforation.

**Nasal packing:** For severe epistaxis, the anterior nasal cavity can be packed with ointment-impregnated gauze strips (see Fig. 3.7) or with ready-made foam packs that expand on contact with fluid. Both nasal cavities should always be packed in order to produce adequate counterpressure.

Alternatives to anterior nasal packing are shown in 3.6. Intranasal packing should not remain in place for more than 2–3 days. Balloon catheters should be progressively deflated starting on the second day; otherwise they may cause irreversible tissue necrosis. Long-term mucosal hygiene should be maintained after the packing is removed.

---

**Table 3.2 Systemic causes of epistaxis**

<table>
<thead>
<tr>
<th>Classification</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vascular and circulatory diseases</td>
<td>Atherosclerosis, arterial hypertension</td>
</tr>
<tr>
<td>Infectious diseases</td>
<td>Influenza, measles, typhus</td>
</tr>
<tr>
<td>Endocrine changes or diseases</td>
<td>Pheochromocytoma, pregnancy, diabetes mellitus</td>
</tr>
<tr>
<td>Hemorrhagic diathesis</td>
<td></td>
</tr>
<tr>
<td>• Coagulopathies</td>
<td>Congenital: e.g., hemophilia A and B, Willebrand disease</td>
</tr>
<tr>
<td>• Platelet disorders</td>
<td>Acquired: e.g., anticoagulant therapy, hepatocellular insufficiency</td>
</tr>
<tr>
<td>• Thrombocytopenias</td>
<td>Idiopathic thrombocytopenic purpura, platelet proliferation disorders, platelet distribution disorders</td>
</tr>
<tr>
<td>• Thrombocytopenias</td>
<td>Acquired: uremia, dysproteinemia, adverse effects of dextran and acetylsalicylic acid (ASA) therapy</td>
</tr>
<tr>
<td>• Vasopathies</td>
<td>Schönlein–Henoch purpura, Osler disease</td>
</tr>
</tbody>
</table>

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**3.6 Alternatives to anterior nasal packing and complications**

A double-lumen balloon catheter is introduced and inflated with water to produce local compression in the nasal cavity and nasopharynx. If the bleeding persists, a posterior nasal pack (Belloq pack) can be inserted, but it should be used with caution due to the risk of aspirating the pads in the nasopharynx. Systemic complications of anterior and posterior nasal packing:

- Arterial hypoxia: fall of oxygen partial pressure with pulmonary dysfunction due to an impaired nasopulmonary reflux mechanism.
- Toxic shock: focal staphylococcal infection develops within 24 h after nasal packing, with generalized shock symptoms caused by bacterial toxins.

---

**Fig. 3.5 Septal perforation**

The bleeding in this case is from the edges of a septal perforation. The endoscope was introduced into the right nasal cavity. The left middle turbinate and right inferior turbinate are visible in the background.
Fig. 3.6 Flowchart for the diagnosis and treatment of epistaxis

Epistaxis

General measures:
- Upright posture
- Ice bag on back of neck
- Compress the nostrils
- Tell the patient not to swallow blood
- Place an i.v. line

Blood pressure?

Elevated
- Antihypertensive agent
- Decongest and anesthetize nasal mucosa
- Inspect nasal cavity by anterior rhinoscopy or endoscopy
- Kiesselbach’s area
- Light bleeding
- Silver nitrat cautery of bleeding site

Normal or decreased
- Volume replacement
- Lab tests:
  - Hb
  - Platelet count
  - Clotting time
  - Coagulation: TPT, PTT, TT

If necessary, give platelet concentrate or fresh frozen plasma

Changes in nasal septum (perforation, spur, etc.)

Posterior sites
- Bilateral anterior nasal pack or balloon tamponade
- Heavier bleeding
- Surgical ligation of bleeding vessel (Fig. 3.8) or angiographic embolization

Insults the anterior nose
- Local surgical treatment (e.g., septoplasty)

Lab tests:
- Hb
- Platelet count
- Clotting time
- Coagulation: TPT, PTT, TT

Blood pressure?

Elevated
- Antihypertensive agent
- Decongest and anesthetize nasal mucosa
- Inspect nasal cavity by anterior rhinoscopy or endoscopy
- Kiesselbach’s area
- Light bleeding
- Silver nitrat cautery of bleeding site

Normal or decreased
- Volume replacement
- Lab tests:
  - Hb
  - Platelet count
  - Clotting time
  - Coagulation: TPT, PTT, TT

If necessary, give platelet concentrate or fresh frozen plasma

Changes in nasal septum (perforation, spur, etc.)

Posterior sites
- Bilateral anterior nasal pack or balloon tamponade
- Heavier bleeding
- Surgical ligation of bleeding vessel (Fig. 3.8) or angiographic embolization

Insults the anterior nose
- Local surgical treatment (e.g., septoplasty)
**Vascular ligation or embolization:** The most common source of bleeding from the posterolateral part of the nasal cavity is the sphenopalatine artery (branch of the maxillary artery), which can be coagulated or clipped under endoscopic control. The ligation or angiographic embolization of a larger arterial trunk may be considered as a last recourse. When this is done, the source of the bleeding must be accurately identified since the nasal lining is supplied by various arteries (Fig. 3.8).

Prevention of recurrent bleeding: Besides the conservative treatments noted above, some causes of epistaxis require surgical treatment since nasal packing alone is of only temporary, symptomatic benefit (3.7).

**3.7 Surgical prevention of recurrent epistaxis**

The main indications for surgery are changes in the nasal septum such as septal spurs (Fig. 3.2b, p. 30), ridges, and perforations (Fig. 3.5). Treatment consists of straightening the nasal septum (septoplasty, see p. 30) or closing the septal perforation (e.g., by implanting an auricular cartilage graft and using local mucosal flap advancement).

In diseases that are associated with vascular changes, such as Osler disease, telangiectatic areas on the septal mucosa can be treated with a surgical laser. The figure shows numerous punctate telangiectasias in the left nasal cavity in Osler disease.

If laser treatment is inadequate, other surgical options are available. In a Saunders dermoplasty, for example, the telangiectatic septal mucosa is resected and replaced with a free skin graft (e.g., from the supraclavicular area).

![Fig. 3.7 Anterior nasal packing](image)

Treatment of a patient with epistaxis. Ointment-impregnated gauze strips are layered into both nasal cavities. An ice bag is placed on the back of the patient’s neck.

![Fig. 3.8 Vascular ligation for severe epistaxis](image)

Depending on the bleeding source, various vessels can be ligated through a cervical approach, by the transnasal endoscopic route, or by a transmaxillary route in the pterygopalatine fossa.
3.4 Soft-Tissue Injuries and Plastic Surgery

Facial soft-tissue injuries are still a common occurrence in recreational and traffic accidents. When improperly managed, they can result in disfiguring scars and deformity. Poor cosmetic results are particularly objectionable in this very conspicuous region. This deals with necessary diagnostic measures and also reviews the most important techniques of facial plastic surgery.

**Diagnosis**

Before a traumatic facial wound is treated, possible coexisting fractures should be excluded by clinical examination and, if necessary, by imaging studies such as biplane skull films, standard sinus projections (see Fig. 2.7, p. 22), and computed tomography (CT) scans. Especially with bite wounds, a smear should be taken for microbiologic examination.

Every patient with facial injuries should be asked about tetanus immunization.

**Treatment**

Prior to surgical treatment, measures are taken to reduce micro-organism counts and prevent infection (tetanus, rabies), especially in patients with bite wounds.

In the interest of maximum tissue preservation, only tissues that are definitely necrotic should be debrided from facial wounds.

Wound margins should never be reapproximated under tension, as this would result in aesthetic and functional deficits such as incomplete eyelid closure.

In most soft-tissue injuries to the nose, adequate treatment consists of primary reapproximation and suturing of the wound margins.

**Scar Camouflage**

Two fundamental local principles in facial soft-tissue surgery are the relaxed skin-tension lines (RSTLs, see Fig. 1.1a, p. 2) and the aesthetic units (see Fig. 1.1b). The RSTLs are a particularly important consideration when there are no soft-tissue or skin defects and only a direct closure is required, since scars are easier to camouflage when they are oriented along the RSTLs.

The techniques described below are also useful for revising a functionally and/or aesthetically objectionable result, such as lengthening a heavily contracted scar.

Z-plasty (Fig. 3.9a, b): When a wound margin runs perpendicular to the RSTLs, it can be reoriented with a single or multiple Z-plasty and lengthened in the direction of the scar axis.
Fig. 3.10 Local flap techniques

a Horizontal advancement flap

b Bilobed flap

c Rhomboid flap

d Island flap

a Small “Burow triangles” are excised at the ends of the incisions, allowing the two rectangular flaps to be advanced for defect closure.
b The bilobed flap is a butterfly-shaped advancement flap used to close a defect.
c The rhomboid flap can be used on the nasal flank, as illustrated, or on the cheek.
d The skin between the defect and superficial flap is undermined, and the island flap is pulled into the defect on its subcutaneous pedicle.
W-plasty (Fig. 3.9c): The principal effect of this technique is to lengthen the scar.

Broken-line closure (Fig. 3.9d): The effect of this technique is to “optically disperse” the scar, making it more irregular and less noticeable.

Repair of Tissue Defects

Soft-tissue defects (traumatic or post-tumor resection, see p. 62) often cannot be adequately managed by primary wound closure with reapproximation of the skin margins.

Smaller tissue defects can be repaired with local flaps such as a sliding flap, bilobed flap, rhomboid flap, or island flap (Fig. 3.10, p. 37). A larger defect of the nose can be covered by turning down a forehead flap (Fig. 3.11), if necessary after first reconstructing the nasal skeleton with cartilage and bone grafts.

Local flaps are often inadequate for more extensive defects of the external nose (e.g., tumor, dog bite), which may require more complex reconstructive procedures using an autologous transfer. This may consist of a pedicled flap (e.g., the myocutaneous pectoralis major flap, Fig. 3.12) or a microvascular free flap. In a microvascular free transfer, the autologous tissue is removed with its supply vessels, which are anastomosed to corresponding arteries and veins at the recipient site. Very extensive defects in the nasal region, like those created by a tumor resection, are repaired in multiple sittings using more complex flap transfers from the scalp (Fig. 3.13).

If the graft must also provide a degree of stability (e.g., alar cartilage of the external nose), this can be accomplished with a composite graft harvested from the auricle (Fig. 3.14) or from the costal or septal cartilage.

Larger defects of the nose can be reconstructed with a forehead flap based on the supratrochlear arteries. a The skin surrounding the defect is circumscribed, mobilized, and turned downward to provide intranasal lining. b Next the forehead flap is raised and partially backed with cartilage (composite graft) for coverage of the external defect. c The forehead defect can usually be closed directly.

Fig. 3.11 Forehead flap

Fig. 3.12 Pedicled myocutaneous flap

The myocutaneous pectoralis major flap is frequently used in the head and neck and is useful for repairing large defects in the facial region. Based on the thoracoacromial artery, the flap is composed of skin, subcutaneous soft tissue, and portions of the pectoralis major muscle. It is mobilized, swung into the tissue defect, and sutured into place.
Fig. 3.13 Scalp flap

The Converse scalp flap is usually based on the superficial temporal artery. Owing to its size and width, one of its applications is for total nasal reconstruction. Backing material (e.g., costal or auricular cartilage) should be added to the flap when it is mobilized and inset in order to obtain proper nasal height.

Fig. 3.14 Composite graft

The nasal alar defect is repaired with a composite graft of cartilage and skin from the auricle to restore adequate stability to the nasal vestibule.

b, c A woman with a nasal alar defect caused by a dog bite, shown before (b) and after (c) debridement of the wound margins.

d Appearance after inset of the auricular composite graft.

e Appearance 6 months later. The healed graft shows an excellent color and texture match with its surroundings.
3.5 Fractures of the Nasal Pyramid and Lateral Midface

Bony injuries to the nasal pyramid and midface are still common in sports and traffic accidents. Most of the injuries are closed fractures. The initial findings may be deceptive, due to hematoma-induced soft-tissue swelling.

Nasal Pyramid Fracture

The nasal pyramid is predisposed to fractures because of its exposed location. The fractures are classified as open or closed on the basis of concomitant soft-tissue injuries.

Diagnostic procedure: Inspection may show obvious deviation of the external nose (Fig. 3.15a) or a simple depression of the lateral nasal wall. Swelling of the surrounding soft tissues is also present, usually caused by a hematoma. Intranasal inspection by anterior rhinoscopy or endoscopy is done to check for concomitant mucosal injuries and especially to evaluate the nasal septum, which also may be fractured. Crepitus noted on palpation confirms the suspicion of a fracture. Further diagnostic measures include radiographs of the nose in the lateral projection (Fig. 3.15b) and standard sinus projections to exclude bony involvement of the lateral midface (see below).

Complications: When a septal fracture is covered by an intact soft-tissue envelope, there is a danger of subperichondrial hemorrhage with hematoma formation (Fig. 3.16a). The hematoma may become infected, giving rise to a septal abscess (Fig. 3.16b), which in turn can lead to cartilage necrosis with loss of the nasal septum and dorsal saddling. Alternatively, the infection may spread to the cranial cavity by the vascular route, causing meningitis.

Treatment: Surgical treatment is generally indicated due to the potential for permanent nasal deformity. An open fracture requires immediate surgical care accompanied by tetanus prophylaxis or a tetanus booster. If the fracture is displaced and closed, it can be safely reduced during the initial week after the injury. The displaced or depressed bone fragments can be reduced manually or with the aid of a special instrument (elevator) (Fig. 3.17). After the reduction, the nasal cavities should be packed to provide “internal splinting,” and a plaster cast is applied externally.

Fig. 3.15 Nasal pyramid fracture

a The clinical appearance, showing deviation of the external nose with an intact soft-tissue envelope.

b The lateral radiograph of the nose demonstrates a fracture line (arrow).
Lateral Midfacial Fractures

Lateral midfacial fractures are usually caused by blunt trauma to the side of the face. Affected structures of the bony facial skeleton are the maxillary sinus, orbit, and the zygoma or zygomatic arch (see Fig. 1.2, p. 2 and Fig. 1.6, p. 6).

An isolated fracture of the orbital floor with a partial herniation of the orbital contents into the maxillary sinus is a special type of lateral midfacial fracture called a blow-out fracture (see Fig. 3.20 b).

Symptoms: A depressed fracture of the zygoma presents clinically with facial asymmetry. Depression of the zygomatic arch frequently causes limited mouth opening.

Fractures of the orbital floor can cause diplopia on upward gaze due to entrapment of the inferior rectus muscle.

Sensory disturbances involving the cheek, ipsilateral upper lip, and lateral nasal wall suggest a direct or indirect fracture-induced lesion of the infraorbital nerve, which enters the buccal soft tissues below the infraorbital margin and is commonly involved by fractures of the orbital floor.

Diagnosis:

Inspection: Swelling is usually present due to subcutaneous hemorrhage (peri orbital or “monocle” hematoma, Fig. 3.18). Asymmetry of the affected facial half is most likely to occur with depression of the zygoma or zygomatic arch, depending on the location and extent of the fracture (Fig. 3.19).

Enophthalmos signifies involvement of the orbital floor (herniation of orbital contents).

It is prudent to seek ophthalmologic consultation in these cases.
Palpation: Concomitant soft-tissue swelling can make it difficult or impossible to palpate sites of bony discontinuity or displacement. The following areas should be examined:

- Frontozygomatic suture (upper part of the lateral orbital rim)
- Infraorbital margin (anterior bony margin of the orbital floor)
- Zygomatic arch (often difficult to evaluate due to soft-tissue swelling)

Sensory testing: Wisps of cotton can be used to test sensory function on the healthy and affected sides.

Radiographs: Whenever a lateral midfacial fracture is suspected, standard sinus radiographs should be obtained (occipitomental and occipitofrontal projections, see Fig. 2.7, p. 22) to define the extent of the bony discontinuity or displacement (Fig. 3.20a, b).

Treatment: Surgical treatment is unnecessary for undisplaced, asymptomatic fractures, but it is indicated for displaced fractures or fractures that are causing symptoms such as sensory deficits in the distribution of the infraorbital nerve, diplopia on upward gaze, enophthalmos, restricted jaw opening, or facial asymmetry. Treatment consists of reduction and fixation of the bone fragments using miniplates, interosseous wiring, or both (Fig. 3.21a and Fig. 3.22). In all cases the patient should be cautioned to avoid nose blowing.
Fig. 3.21 Displaced lateral midfacial fracture on the left side

The sinus radiographs show a displaced lateral midfacial fracture on the left side with separation of the frontozygomatic suture and a displaced orbital floor fracture before (a) and after (b) reduction and stabilization with miniplates.

Fig. 3.22 Internal fixation of an orbital rim fracture

A fracture of the left infraorbital margin has been stabilized by wire and miniplate fixation. The orbital contents are retracted upward with a spatula.

Fig. 3.23 Depressed fracture of the zygoma

Depressed fracture of the zygomatic arch (arrows), demonstrated by the bucket-handle view (a) and axial CT scan (b).

Fig. 3.24 Depressed fracture of the zygomatic arch

a Sinus radiograph of a depressed zygomatic fracture on the right side (the patient had been kicked by a horse).
b Axial computed tomogram in the same patient shows the displaced body of the zygoma (arrow), which has penetrated the orbital cone from the lateral side.
3.6 Fractures of the Central Midface and Anterior Skull Base

Although fractures of the central midface and anterior skull base (frontal skull base, rhinobase) are separate entities, they are on a clinical continuum and are therefore discussed in the same.

Classification:
Central midfacial fractures: The Le Fort classification (Fig. 3.25) describes various midfacial fracture patterns ranging from isolated detachment of the alveolar process (Le Fort I, Fig. 3.26) to separation of the midfacial bones from the anterior skull base (Le Fort III).

Frontobasal fractures: These are bony injuries to the anterior skull base and adjacent paranasal sinuses (frontal and sphenoid sinuses, ethmoid labyrinth). The Escher classification distinguishes four types of frontobasal fractures based on the location and extent of the fracture lines (Fig. 3.27).

Etiopathogenesis: Fractures of the central midface and frontal skull base generally occur in multiply injured patients (usually vehicular accidents), but they can also result from "trivial" trauma or even a surgical procedure (e.g., endoscopic sinus surgery), since some bony portions of the anterior skull base are quite thin (e.g., the cribriform plate of the ethmoid bone). Frontobasal fractures occupy a special place among skull fractures because they are usually an "indirectly open" injury that creates a communication between the cranial cavity and the environment.

Ascending infection can occur via the adjacent paranasal sinuses in frontobasal fractures and can lead to life-threatening intracranial complications (e.g., meningitis, brain abscess).

The dura along the fracture line tends to become torn at sites where it is firmly adherent to the bone of the skull base (e.g., cribriform plate, sphenofrontal suture, sellar tubercle, spheno-occipital synchondrosis).

Symptoms: Most patients present with a unilateral (see Fig. 3.18) or bilateral periorbital hematoma, depending on the nature and direction of the traumatizing force. A dish face is seen in combined fractures (Le Fort II–III, Escher III) where the midface has been separated from the skull base and displaced inward.

Cerebrospinal fluid (CSF) rhinorrhea is one of the few reliable signs of an anterior skull base fracture with associated dural injury. It can also occur with petrous bone fractures, in which case CSF leakage occurs via the eustachian tube.

Particularly in fresh head injuries, CSF leak from a dural tear may be obscured by heavy bleeding or may be contained by bone fragments, prolapsed brain tissue, swollen mucosa, or foreign bodies.

Severe craniocerebral trauma can also result in vision loss caused by ocular destruction or injury to the optic nerve (nerve contusion, rupture of the nerve in an intact sheath due to sagittal brain motion within the skull). Diplopia due to oculomotor palsy from damage to the third, fourth, or sixth cranial nerve is somewhat rare and occurs only if the fracture line runs through the cavernous sinus. Extensive injuries with sites of bone dehiscence can lead to cerebral prolapse, with
brain tissue herniating externally or into the nasal cavity. Anosmia can result from a fracture of the cribriform plate with avulsion of the fila olfactoria, or it may signify damage to more central structures in the setting of a cerebral concussion or contusion.

**Diagnosis:**

Patients with craniocerebral trauma require interdisciplinary care.

Before an ENT examination is performed, the patient’s vital functions should be stabilized by the anesthesiologist or neurosurgeon. Facial soft-tissue injuries are often extensive but may provide little information on possible associated bony injuries at the level of the skull base. A basic impression is gained by palpation of the facial bones (Fig. 3.28) and inspection of the nasal cavity by rhinoscopy or endoscopy. Profuse bleeding from soft-tissue injuries, including intranasal lesions, often makes it difficult to adequately evaluate the injury in the acute stage, and it may not be possible at this time to confirm or exclude a CSF leak in multiply injured patients, many of whom are intubated. The next step after nasal inspection is to inspect the oral cavity and oropharynx. Otoscopy or otomicroscopy should also be performed to exclude a concomitant petrous bone fracture.

A clear, watery nasal discharge should raise suspicion of a CSF leak (Fig. 3.29), which requires differentiation from ordinary nasal mucus. Various CSF tests (e.g., glucose test, β2-transferrin assay) are available for this purpose (3.8).

Intracranial complications (especially bleeding) should also be excluded.

**Computed tomography** supplies additional important information on the location and extent of injuries. Neurosurgeons order CT scans with a soft-tissue window as an initial study for excluding intracranial bleeding, hematoma, and pneumocephalus. This type of study is not useful for the assessment of bony lesions. While indirect signs such as the presence of intracranial air (Fig. 3.30 c) suggest a strong likelihood of dural injury with a frontobasal fracture, only a bone-window CT scan can reliably evaluate or exclude a fracture. So whenever a frontobasal fracture is suspected, high-resolution CT scans (2-mm slice thickness, bone window) of the paranasal sinuses should be obtained in the axial and coronal projections. The
axial scans are for evaluating the anterior and posterior walls of the frontal sinuses (Fig. 3.30 a) and spheno-noid sinus (Fig. 3.30 b), while the coronal scans more clearly define the ethmoid roof and cribriform plate (Fig. 3.30 c, d).

The preliminary testing of hearing and balance is possible only if the patient is conscious and responsive to verbal commands.

Olfactory testing to exclude anosmia often cannot be performed in the acute stage but should be done at a later time. For medicolegal reasons, it should always precede surgical treatment (see Olfactometry, p. 21).

**Treatment:** Every confirmed fracture of the anterior skull base should be treated surgically in operable patients, regardless of whether or not a CSF leak has been detected. The patient should also be instructed not to blow the nose. With an isolated central midfacial fracture that does not involve the anterior skull base, surgical treatment of the maxilla should be provided by a maxillofacial surgeon to ensure the restoration of normal occlusion. The urgency of surgical intervention for a frontobasal fracture is shown in Table 3.3.

Three main surgical approaches to the anterior skull base are available (Fig. 3.31). The choice depends on the individual situation and is made in consultation with the other involved specialties (e.g., neurosurgeon, maxillofacial surgeon, ophthalmologist).
Axial computed tomography scans demonstrate a fracture of the anterior and posterior walls of the frontal sinuses (arrows) (a) and a clivus fracture (arrow) that extends anteriorly into the sphenoid sinus (b). The coronal scans show air in the cranial cavity (c) and a fracture of the ethmoid roof (d) (arrow).

Table 3.3  Indications for the surgical treatment of frontobasal fractures

<table>
<thead>
<tr>
<th>Vital indications (operate immediately)</th>
<th>Absolute indications (operate as soon as possible)</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Life-threatening rise of intracranial pressure due to intracranial hemorrhage</td>
<td>• Open brain injury</td>
</tr>
<tr>
<td>• Bleeding from the nose or sinuses that is refractory to conservative treatment</td>
<td>• Dural tear from an indirectly open head injury</td>
</tr>
<tr>
<td>• Bleeding from an open skull injury that is refractory to conservative treatment</td>
<td>• Penetrating foreign bodies and impalement injuries</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Relative indications (operate in 1–2 weeks)</th>
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</thead>
<tbody>
<tr>
<td>• Displaced bone fragments</td>
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<tr>
<td>• Fractures involving the drainage tracts of the paranasal sinuses (“ostiomeatal unit”)</td>
</tr>
<tr>
<td>• Acute or chronic sinusitis at the time of the injury</td>
</tr>
<tr>
<td>• Post-traumatic sinus inflammation, mucopyocele formation</td>
</tr>
<tr>
<td>• Supraorbital nerve injury due to an adjacent fracture</td>
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</table>
3.7 Inflammations of the External Nose, Nasal Cavity, and Facial Soft Tissues

Inflammatory diseases of the nasal skin usually have a bacterial cause and may be manifested on the exposed skin and the dermal appendages. Although these diseases fall primarily within the scope of dermatology, the otolaryngologist is frequently faced with inflammations of this type that can lead to life-threatening complications unless adequately treated. Inflammation of the nasal cavity (rhinitis) predominantly involves the nasal mucosa. While it can have a variety of causes, it always exhibits a more or less pronounced combination of characteristic symptoms. Although rhinitis and sinusitis are on a continuum owing to the special anatomical and physiological relationships of the nose and paranasal sinuses (see 1.1–1.3, pp. 2–13), the diseases are covered separately in this unit for learning purposes. The various forms of rhinitis are discussed according to their etiology, noting that the great majority of cases seen in clinical practice are mixed forms that have more than one cause.

Purulent Inflammations of the Hair Follicles

Pyodermas of the hair follicles are common purulent inflammations that can occur at almost any age. The main causative organisms are staphylococci. If the disease is confined to the hair follicles, it is termed folliculitis. If the infection spreads to deeper tissues and forms a central core of purulent liquefaction, it is called a furuncle.

Symptoms: Nasal furuncles present as painful, tender, erythematous swellings about the nasal tip and nares (Fig. 3.32). There may be concomitant edematous swelling of the upper lip. The changes are confined to the outer skin and do not involve the mucosa. Fever is sometimes present.

Treatment: The treatment of choice is high-dose parenteral administration of an antibiotic that is active against staphylococci, such as flucloxacillin sodium or dicloxacillin sodium, combined with the local application of an antibiotic-containing ointment (chlortetracycline HCl). Also, the upper lip should be moved as little as possible, and so the patient should be placed on a liquid or semisolid diet and speak as little as possible.

An essential goal of these measures is to prevent the potentially lethal complication of intracranial spread.

Complications: Inadequate treatment or manipulations of the nasal furuncle itself can result in hematogenous spread to intracranial structures, since the veins of the nose and upper lip drain via the angular and ophthalmic veins into the cavernous sinus. If tenderness in the medial canthus of the eye raises a sus-
picion of thrombophlebitis of the angular vein, the vessel should be surgically ligated and divided.

**Erysipelas**

_Etiopathogenesis:_ The principal causative organisms are beta-hemolytic group A streptococci. Less common pathogens are streptococci of other groups, _Staphylococcus aureus_, and gram-negative rods (e.g., _Klebsiella pneumoniae_), which gain entry to the skin through minor injuries (usually on the face and limbs). The inflammation spreads diffusely in the skin and subcutaneous tissue.

_Symptoms:_ Facial erysipelas (Fig. 3.33) usually begins with a high fever and a feeling of tension in the soft tissues, followed rapidly by broad areas of erythema and swelling, which are sharply demarcated from unaffected skin. The tissue is warm to the touch, and small blisters occasionally form.

If facial erysipelas spreads lateral to the nose and about the eyelids, there is a risk of intracranial involvement by hematogenous spread of the causative organisms, as with a nasal furuncle.

**Treatment:** The treatment of choice is the parenteral administration of penicillin. Moist compresses soaked in an antiseptic solution can also be applied locally.

**Inflammations of the Nasal Cavity**

**Acute Rhinitis**

_Epidemiology:_ Acute rhinitis (common cold) is the most prevalent infectious disease. Given its frequency and the fact that the disease does not confer postinfection immunity, acute rhinitis has assumed major epidemiologic and economic significance.

_Etiopathogenesis:_ Rhinoviruses and coronaviruses comprise almost half of the causative organisms of acute viral rhinitis. Other pathogens are influenza viruses and adenoviruses. The infection is transmitted by the airborne route (droplet infection). Cold exposure and other environmental factors can increase the susceptibility of the host to infection. The incubation period is 3–7 days.

_Symptoms:_ The disease begins with an initial dry stage characterized by malaise (lethargy, headache, fever) and local discomfort in the nose and nasopharynx (burning, soreness). This is followed by a catarrhal stage marked by a watery, initially serous nasal discharge and nasal obstruction due to mucosal swelling, which mainly involves the turbinates. The viruses damage the mucociliary transport system, which hampers the normal clearing of secretions. With a profuse nasal discharge, inflammatory changes often develop about the nasal vestibule. Viral damage to the epithelium promotes bacterial colonization, which alters the consistency of the clear nasal discharge, causing it to become mucopurulent (Fig. 3.34). The local and systemic symptoms usually subside in about a week.

_Treatment:_ Treatment consists of supportive measures to relieve nasal obstruction and prevent sinusitis and other sequelae by the use of decongestant nose drops. Nose drops should be used no longer than absolutely necessary (generally no more than one week) due to the risk of tachyphylaxis (see also rhinitis medicamentosa, p. 11) with severe rebound swelling of the nasal mucosa.

Various other options are available for relieving the discomfort of acute rhinitis, including chamomile steam inhalation, “light baths,” and infrared therapy.

3.9 Differential diagnosis of facial soft-tissue swelling

_Lupus erythematosus_ (LE), the most common form of cutaneous LE, is an inflammatory dermatosis that frequently affects the face, spreading in a butterfly-shaped pattern over the cheeks, forehead, and nose. The differential diagnosis also includes _allergic contact dermatitis_, which may be induced by cosmetics, toilet articles, sun creams, or exposure to airborne plant pollens. In strongly sensitized patients who wear a face mask, even a single contact can incite a severe, acute allergic reaction with erythema and edematous swelling of the facial soft tissues. Finally, the differential diagnosis should include _angioedema_, which is also associated with facial swelling that chiefly affects the eyelids and lips (see also 4.4, p. 87).
Antibiotics may also be prescribed in patients with bacterial superinfection or paranasal sinus involvement.

**Nonspecific Chronic Rhinitis**

*Etiopathogenesis:* Chronic inflammation of the nasal mucosa can have various underlying causes. Besides recurrent acute inflammations with progressive damage to the mucosa, nonspecific chronic rhinitis can develop due to anatomic changes (e.g., marked septal deviation, septal spur) or other lesions of the nasal cavity (polyps, tumors) and nasopharynx (adenoids, see 5.3, p.108). Environmental factors such as sustained extreme temperatures or air pollutants can also bring on this condition.

*Symptoms:* Patients present clinically with obstructed nasal breathing and a mucous nasal discharge. They also complain of frequent throat clearing and occasional hoarseness.

*Treatment:* The most important step is to eliminate the cause by removing chronic irritants from the environment or by surgically correcting any intranasal pathology (e.g., septoplasty, p.30). Supportive measures such as decongestant nose drops or nasal irrigation with saline solution are of only temporary benefit.

**Specific Chronic Rhinitis**

This category includes inflammations of varying causes that may be manifested in the nose (3.11).

---

**Allergic Rhinitis**

Diseases of the nose and paranasal sinuses have undergone a disproportionate rise during the past few decades. Meanwhile, the spectrum of rhinologic diseases has changed from a qualitative standpoint as well. As allergic diseases have become more prevalent, there has also been a notable rise in the incidence of allergic rhinitis.

*Etiopathogenesis and classification:* Allergic inflammation of the mucosa is triggered by an immediate, IgE-mediated reaction of the immune system to any of a number of foreign substances, particularly pollens and animal allergens.

Allergic rhinitis is classified as seasonal (hay fever) or perennial according to the presence of the allergen in the environment.

**Seasonal allergic rhinitis** in Central Europe is caused mainly by pollens from alder, hazel, birch, grasses, rye, mugwort, and plantain. Clinical symptoms appear between February and September, depending on the individual allergen spectrum of the patient, and disappear at the end of the pollen season.

By contrast, **perennial allergic rhinitis** is caused by year-round allergen exposure that incites a permanent inflammation of the nasal mucosa. The predominant causative allergens are house dust, pet dander, and molds. The disease may also be caused by certain foods (e.g., strawberries, nuts, eggs, fish) as well as occupational exposure to allergens (e.g., bakers and hairdressers). A new occupational allergen, especially prevalent in health workers, is latex, which is used to manufacture disposable gloves.

*Symptoms:* The clinical manifestations include obstructed nasal breathing and sneezing attacks, a watery nasal discharge, and itching of the nose and eyes (conjunctivitis).
This chronic inflammatory disease is extremely rare and is caused by the gram-positive anaerobe *Actinomyces israelii*. Tuberculosis can involve the nasal mucosa as a primary infection following the inhalation of infectious droplets, forming a primary complex approximately 6 weeks after infection. Another manifestation is lupus vulgaris, the most common postprimary form of cutaneous tuberculosis. Ulcerative cases are marked by increasing necrosis within the tubercular granulomas, which can cause a mutilating destruction of nasal skin and cartilaginous structures. Lupus vulgaris may also arise from the nasal mucosa itself, causing the contiguous destruction of nasal structures from the inside.

**Sarcoidosis**
Sarcoidosis is a common, granulomatous systemic disease of unknown etiology that predominantly affects women under 40 years of age. Although the lymph nodes, lungs, joints, and skin are chiefly affected and involvement of the upper respiratory tract is relatively rare, nasal symptoms may be the initial manifestation of the disease. Involvement of the external nose is called lupus pernio because the characteristic skin changes resemble chilblains (pernio). Involvement of the nasal mucosa mainly affects the septum and inferior turbinate, which develop yellowish, submucous nodules that have the gross appearance of intramucosal granulomas.

**Rhinoscleroma**
This chronic inflammatory disease is extremely rare and is manifested in the nose, oral mucosa, and upper respiratory tract. It is transmitted by *Klebsiella pneumoniae* (subspecies rhinoscleromatis). Rhinoscleroma presents the features of atrophic rhinitis (see below): fetid nasal discharge, dry mucosa, and crusting. The nasal mucosa shows inflammatory infiltrates that may progress to granulations and also involve the nasal vestibule.

**Actinomycosis**
The gram-positive anaerobe *Actinomycyes israelii* can cause this disease, usually in immunocompromised patients. Symptomatic involvement of the nose and paranasal sinuses is somewhat uncommon. Changes in these regions may include firm infiltrates in the nasal mucosa that resemble a nasal furuncle. There have also been sporadic reports of granulations forming on the nasal mucosa and in the paranasal sinuses. Untreated, the inflammation can spread and cause severe tissue destruction with a fatal outcome.

**Syphilis**
Nasal involvement by syphilis occurs mainly in the tertiary stage of the disease. It is manifested by the appearance of isolated gummatas or by diffuse gummatous infiltration of the nasal cavity. Untreated, the disease causes progressive destruction of the surrounding tissue, and eventual bone destruction can occur. Infants with congenital syphilis contracted in utero may also manifest nasal changes that include a purulent and sometimes bloody nasal discharge, which may be mistaken for "normal" infant rhinitis.

**Malleus**
Malleus is a rare infectious disease (causative organism: *Pseudomonas mallei*) that is transmitted to humans from horses and occasionally from house pets. When the nasal mucosa is the portal of entry, it exhibits inflammatory swelling, pustule formation, and ulceration with a viscous nasal discharge containing blood and pus.

**Fungal infections**
Aspergillosis is the most common fungal infection causing chronic specific rhinitis, with fungal colonization occurring mainly in the paranasal sinuses (see Sinusitis in 3.8, pp. 54–57). In rare cases and especially in immunocompromised patients, the infection may take an aggressive, fulminating course with the destruction of surrounding structures, resulting in a very high mortality.

**Mucormycosis** resembles aspergillosis in its symptoms and course. The invasive form of mucormycosis still has a relatively high mortality rate despite the availability of systemic antifungal agents, but it mainly affects patients with a weakened immune status.

**Rhinosporidiosis** is another very rare disease caused by the spore-forming fungus *Rhinosporidium seeberi*. Highly vascular, friable granular lesions develop in the anterior portions of the nose and may spread to involve the paranasal sinuses and nasopharynx.

### 3.12 Nasal hyperreactivity

Because the nasal mucosa is reactive to physical, chemical, and pharmacologic stimuli, which may take the form of allergens, pollutants (cigarette smoke, dust, fumes), or even position changes or exertion, the term “nasal hyperreactivity” (analogous to “bronchial hyperreactivity”) has lately been coined as a collective term for a heightened reactivity of the nasal mucosa to these agents. Some of the diseases that lead to nasal hyperreactivity are marked by signs of inflammation and others chiefly by disturbances of autonomic nervous regulation, with an altered response of the associated receptors on vessels, nerves, and glands of the mucosa.

**Diagnosis:** The diagnostic workup should include a detailed allergy history (do the symptoms present year-round or only during contact with certain animals or plants? do they disappear during vacation?). With seasonal allergic rhinitis, inspection of the nasal cavity typically reveals a bluish-purple discoloration of the mucosa (Fig. 3.35). With perennial rhinitis, the mucosa is bright red and shows inflammatory changes. Careful allergy testing (see 2.2, p.19) is necessary to identify the antigens involved.
Treatment:

The best treatment strategy is to avoid contact with the allergen or eliminate allergenic irritants from the environment.

Since the avoidance strategy is not always possible for practical reasons, most patients with allergic rhinitis receive pharmacologic treatment. The various drugs and their properties (mast-cell stabilizers, local and systemic H1 antihistamines, local steroids) are reviewed in Table 3.4. Further information on immunotherapy (hyposensitization) and surgical options can be found in 3.13.
Vasomotor Rhinitis

Vasomotor rhinitis resembles allergic rhinitis in its clinical features, but there is no evidence that the patient has been previously sensitized. The pathogenesis of vasomotor rhinitis is believed to involve neurovascular autonomic disturbances in regulating the tonus of the nasal mucosal vessels. The symptoms consist of obstructed nasal breathing, watery nasal discharge, and sneezing. The history shows that the symptoms are related to a temperature change, the consumption of hot liquid or alcohol, or less specifically to “emotional stress.” On inspection, the appearance of the nasal mucosa is similar to that in allergic rhinitis. Medical therapy includes the use of antihistamines or corticosteroid-containing nasal sprays. In the Kneipp system of therapy, ice-cold water is sniffed up the nose as a way of “training” the neuroautonomic regulation of the blood supply to the nasal mucosa. The last recourse for intractable vasomotor rhinitis is surgical reduction of the turbinates by electrocautery, laser ablation, or mucotomy, especially in cases with pronounced inferior turbinate hyperplasia. If significant septal deviation is present, a septoplasty should be performed.

Atrophic Rhinitis

Symptoms: Atrophic rhinitis is characterized by pronounced dryness of the nasal mucosa. Severe cases, especially with secondary bacterial colonization, are marked by a fetid nasal odor that is not perceived by the patient due to degeneration of the olfactory epithelium.

The etiology of primary atrophic rhinitis is unknown. Secondary forms can have various causes including an extensive prior tumor resection, the excessive use of nose drops, drug abuse (cocaine), or previous radiotherapy for nasal and sinus tumors. Iatrogenic causes include a botched septoplasty or an excessive turbinate reduction (conchotomy). Endoscopic examination reveals a broad nasal cavity lined with dry, crusted mucosa.

Treatment should begin with conservative, symptomatic measures (saline “nasal douche,” soothing mucosal ointments). Under no circumstances should decongestant nose drops be used, as the vasoconstriction would exacerbate the patient’s symptoms. If conservative treatments prove inadequate, an attempt can be made to reduce the nasal cavity surgically by the submucous implantation of cartilage grafts. This creates a relative increase in surface area in relation to the volume of the nasal cavity.

Hormonal Rhinitis

Synonym: pregnancy-associated rhinitis

Hormonal rhinitis occurs mainly during pregnancy and is believed to be caused by estrogen-induced swelling of the mucosa with nasal airway obstruction. The symptoms diminish as term approaches and disappear after the delivery.

Rhinitis Medicamentosa

This disease occurs mainly as a side effect from the long-term use of decongestant nose drops. It can also result from the use of certain antihypertensive drugs—e.g., rauwolfia alkaloids, beta-blockers, angiotensin-converting enzyme (ACE) inhibitors—and from oral contraceptive use, in which case the rhinitis is attributed to a vasoactive estrogen effect. Clinical symptoms consist of obstructed nasal breathing, dry mucosa, and occasional olfactory disturbances.
3.8 Sinus Inflammations

Sinus inflammations (sinusitis) generally develop in association with rhinitis, and so the term “rhinosinusitis” is often applied to these disorders. Despite the continuum that exists between rhinitis and sinusitis, they are discussed as separate entities in this textbook for teaching purposes. Inflammations that are confined chiefly to the nasal cavity are covered in the previous . This unit deals with acute and chronic sinusitis in addition to nasal polyposis, mucoceles, pyoceles, and rhinosinogenic complications—diseases in which clinical symptoms arising from the paranasal sinuses are the dominant features.

Acute Sinusitis

Etiopathogenesis: While acute sinusitis in children predominantly affects the ethmoid cells due to incomplete pneumatization of the other sinuses (see 1.1, p. 4), acute sinusitis in adults affects the following sinuses in descending order of frequency: maxillary sinus, ethmoid cells, frontal sinus, and sphenoid sinus. The inflammation may involve one, several, or all of the paranasal sinuses (pansinusitis). Acute sinusitis generally results from the spread of an intranasal inflammation (rhinitis), since the mucosa of the paranasal sinuses communicates with that of the nasal cavity (rhinogenic sinusitis). Accordingly, the causative viruses of acute rhinitis (see p. 49) are etiologically important in addition to the common bacterial organisms Haemophilus influenzae and Streptococcus pneumoniae.

Although rhinitis has a very marked tendency to involve the contiguous sinus mucosae, acute rhinitis does not invariably lead to symptomatic sinusitis. The extent of the inflammation in the sinus system and the associated symptoms depend on various factors:
• Individual functional anatomy (see Chapter 1, p. 2)
• Individual immune status
• Specific virulence of the causative organism

Besides rhinogenic sinusitis, there are also rare instances of dentogenic sinusitis arising from a dental root infection, an apical granuloma, or a maxillary sinus fistula following a tooth extraction [3.14].

Symptoms: The clinical picture is marked by the features of acute rhinitis combined with a variable degree of headache, which is exacerbated by bending over. Generally the pain is most intense over the affected sinuses (see also Fig. 1.6, p. 6). Thus, the pain of maxillary sinusitis is greatest over the maxillary sinus and the adjacent midface and temple. Ethmoid sinusitis is most painful over the bridge of the nose and the medial canthus of the eye, and frontal sinusitis over the anterior wall and floor of the frontal sinus, with pain radiating toward the medial canthus. The pain of sphenoid sinusitis is fairly nonspecific, marked by a dull, aching pressure located at the center of the skull and radiating to the occiput.

Diagnosis: Rhinoscopy or nasal endoscopy often reveals pus tracking along the middle meatus of the nasal cavity (Fig. 3.36), but a purulent track may not be seen if the mucosa is greatly swollen. With isolated sphenoid sinusitis, pus may be found about the ostium in the anterior wall of the sphenoid sinus or on the posterior wall of the pharynx.

Sinus radiographs (see also Fig. 2.7, p. 22) may show partial opacification of the affected sinus due to mucosal swelling (Fig. 3.37a) or may demonstrate a fluid level if the sinus contains free pus (Fig. 3.38). An alternative to radiography, especially for follow-up and in children and pregnant women, is ultrasonography (A-mode or B-mode), which avoids radiation exposure.

Treatment: Conservative treatment options should be exhausted before surgery is considered. The latter may be necessary in cases where the complaints of acute sinusitis do not respond to conservative treatment modalities and in cases with persistent sinus empyema.

Conservative therapy: Ventilation and drainage of the paranasal sinuses can be improved by the use of decongestant nose drops, nasal spray, or by inserting a cotton pack soaked with nose drops into the middle meatus. In more severe forms associated with fever and significant malaise, antibiotics (e.g., amoxicillin) should be administered. Heat therapy (electric light bath) and the inhalation of chamomile or sage are recommended as adjuncts.

3.14 Special forms of sinusitis

Other forms are nosocomial sinusitis resulting from prolonged nasal intubation, barosinusitis caused by pressure changes during flying or diving, and swimmer’s sinusitis caused by the entry of infectious micro-organisms into the sinus during swimming.
Surgical therapy: Maxillary sinusitis can be treated by maxillary sinus puncture following decongestion and topical anesthesia of the nasal mucosa. Two approaches are available: first, “sharp puncture” through the inferior meatus, passing the needle below the inferior turbinate; and second, “blunt puncture” via the natural maxillary sinus ostium in the middle meatus. In the sharp puncture technique, there is a significant risk of complications due to air embolism if air is inadvertently injected into the sinus after a medication has been instilled. Another potential danger is perforation of the lateral sinus wall, resulting in a buccal abscess or perforation of the sinus roof causing infection of the orbital contents (Fig. 3.39).

A frontal sinus empyema can be surgically drained through a “Beck puncture” (Fig. 3.15). There should be little hesitation in using this procedure, since the frontal sinus directly borders the cranial cavity, posing a risk of meningencephalitis or frontal brain abscess.
Chronic Sinusitis

**Etiopathogenesis:** Besides intranasal anatomic changes such as septal deviation and septal spurs, a variety of other diseases of a chronic inflammatory, allergic, traumatic or neoplastic nature can lead to chronic sinusitis. The common pathogenic mechanism is impaired ventilation of the ostiomeatal unit (see 1.3, p.7) due to stenosis or obstruction of this region. This hampers drainage of the dependent sinus systems, particularly the adjacent maxillary sinus and anterior ethmoid cells. As the mucosa becomes swollen, especially in the narrow anatomical passages of the ostiomeatal unit, a vicious cycle becomes established that initially leads to recurrent bouts of acute inflammation and eventually culminates in a persistent, chronic sinusitis.

Chronic sinusitis frequently affects the maxillary sinus and ethmoid cells, while the frontal and sphenoid sinuses are less commonly involved.

**Symptoms:** The character of the pain is variable and can range from a feeling of pressure to persistent or recurrent headaches. Many patients also complain of nasopharyngeal drainage (postnasal drip), and some complain of obstructed nasal breathing.

**Diagnosis:**

**Rhinoscopy, endoscopy:** The nasal cavity is inspected by rhinoscopy or endoscopy, giving particular attention to changes in the nasal septum, the condition of the turbinates (turbinate hyperplasia, pneumatized middle turbinate, concha bullosa), and the appearance of the ostiomeatal unit (mucosal swelling, polyps, tumors, etc., see also 1.3, p.7). Other causes of impaired ventilation and drainage in the nasal cavity itself (e.g., tumors) should also be excluded.

**Imaging studies:** Today, computed tomography is considered the only acceptable modality for imaging the paranasal sinuses if chronic sinusitis is suspected. Conventional sinus radiographs are of very limited value in diagnosing chronic sinusitis due to artifacts from superimposed structures. Also, only CT scans can accurately define the key anatomic structures that are important for accurate preoperative planning (Fig. 3.40).

**Treatment:** Conservative treatment options include decongestant nose drops (for no more than one week), heat therapy (electric light cabinet, microwaves, infrared), and broad-band antibiotics (e.g., amoxicillin) for acute exacerbations of sinusitis with fever and malaise. Mucolytics can also be administered for supportive therapy. With an allergic etiology, appropriate antiallergic therapy should also be provided (see p.52). All of these conservative therapies are of symptomatic benefit and cannot eliminate the cause of chronic sinusitis. The only definitive treatment is sinus surgery (3.16).

Fig. 3.40 Chronic sinusitis

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**3.15 Frontal sinus irrigation**

For the Beck puncture, the skin and subcutaneous soft tissues are divided at the medial border of the eyebrow, and the anterior wall of the frontal sinus is opened with a drill. Secretions and pus are aspirated from the frontal sinus, and the sinus is irrigated with decongestant nose drops and an antibiotic solution.

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Nasal Polyposis

**Pathogenesis and morphology:** Nasal polyposis is a very complex condition that develops in response to a variety of noxious stimuli, appearing morphologically as edematous, polypoid hyperplasia of the sinus mucosa (usually in the anterior ethmoid cells and maxillary sinus) and projecting into the nasal cavity in the form of polyps (Fig. 3.41).

**Etiology:** Besides genetic causes, nasal polyps are attributed mainly to chronic irritation of the mucosa, like that occurring in chronic rhinitis or sinusitis. They can also form in response to allergic rhinitis and acetylsalicylic acid (ASA) intolerance (ASA pseudoallergy). The functional anatomy of the ostiomeatal unit, with its slitlike passages, appears to have causal significance in nasal polyposis, as in chronic sinusitis, because it controls the ventilation and drainage of the frontal and maxillary sinuses. The opposing mucosal surfaces in these areas are often separated by a distance of less than 1 mm. If they come into contact, this will impair the mucociliary clearance mechanism and hamper the normal transport of harmful substances toward the nasopharynx. Most polyps form at these narrow passages.

Whether, when, and to what extent these pathologic changes lead to symptomatic polyposis varies in different individuals and may depend partly on the timing of diagnosis and treatment.

Nasal polyps are rarely observed in children. Most occur in a setting of cystic fibrosis.

**Symptoms:** The clinical manifestations of nasal polyps depend on their extent and may consist of obstructed nasal breathing, hyposmia or anosmia (due to obstruction of the olfactory groove), headache (due to impaired ventilation and drainage in the paranasal sinuses), snoring, rhinophonia clausa, and frequent throat clearing due to associated postnasal drainage. Spread to the lower airways can lead to laryngitis with hoarseness and bronchitic symptoms.

**Diagnosis:** As in chronic sinusitis, the diagnosis is established by careful rhinoscopic or endoscopic evaluation of the nasal cavity, giving particular attention to the lateral nasal wall. The imaging modality of choice is computed tomography. Further diagnostic measures consist of allergy tests and olfactory testing.

**Treatment:** Treatment may begin with symptomatic conservative measures such as the use of corticoid-containing nasal sprays and systemic antihistamines. Systemic steroids may also be tried. A partial or even complete remission of nasal polyps can sometimes be achieved with these measures alone.

Many cases will require surgical treatment, however. Besides intranasal polypectomy, which is performed mainly in older or higher-risk patients, an important current option is intranasal sinus surgery using endoscopic or microsurgical technique (see also Treatment of Chronic Sinusitis, p. 57).

**Prognosis:** Given the complex and poorly understood etiopathogenesis of nasal polyposis, which is not considered a single entity, the prognosis is guarded even with modern surgical techniques, and even the...
most meticulous ablative sinus surgery cannot prevent a recurrence. As a result, there is often no alternative to long-term medical prophylaxis with topical steroid sprays.

**Mucoceles and Pyoceles**

A mucocele is a cystlike, mucus-containing sac that can form within a paranasal sinus. A pyocele is a mucocele that contains purulent material as a result of superinfection.

**Pathogenesis:** A mucocele may be caused by adhesions (postinflammatory, post-traumatic, or postoperative) that obstruct drainage from the paranasal sinus system. Mass lesions (polyps, tumors) can also obstruct and obliterate the drainage tracts, leading to mucocele formation. The outflow obstruction causes the mucocele to exert increasing pressure on the surrounding sinus walls, resulting in progressive thinning of the bone. In this way the mass can erode into adjacent structures such as the orbit or even the cranial cavity. The most common site of occurrence is the frontal sinus, followed by the ethmoid cells, maxillary sinus, and sphenoid sinus.

**Symptoms:** A frontal sinus mucocele usually presents as an isolated, tense swelling over the anterior wall of the frontal sinus (Fig. 3.42). It may also cause inferolateral displacement of the orbital contents, especially if it has eroded through the sinus floor. On the other hand, swelling in the cheek area with upward displacement of the orbital contents is more characteristic of a maxillary sinus mucocele (Fig. 3.43 a). Proptosis, limited ocular movements, and diplopia may also occur, depending on the location of the mass. Unlike mucoceles in the frontal sinus, ethmoid cells (Fig. 3.44) and maxillary sinus, sphenoid sinus mucoceles (Fig. 3.45) often have nonspecific clinical manifestations, with complaints similar to those of sphenoid sinusitis (headache radiating to the vertex and occiput).

**Diagnosis:** A prior surgical history is often helpful in making a diagnosis. The clinical appearance may also be suggestive (Fig. 3.42, Fig. 3.43 a), whereas inspection of the nasal cavity by rhinoscopy or endoscopy usually does not advance the diagnosis. On the other hand, modern sectional imaging modalities (computed tomography and magnetic resonance imaging, Fig. 3.43 b, Fig. 3.44, Fig. 3.45) can make it possible to delineate the mucocele from surrounding tissues and differentiate it from a malignant tumor.

**Treatment:** The treatment of choice is surgical removal of the mucocele.

**Rhinosinogenic Complications**

There are various mechanisms by which serious and even life-threatening complications can arise from inflammatory diseases of the paranasal sinuses. Complications with inflammatory involvement of the orbit (orbital complications) are distinguished clinically from bone and soft-tissue complications (osteomyelitis) and from intracranial complications.

**Orbital Complications**

Inflammatory complications with orbital involvement most commonly arise from the ethmoid cells or frontal sinus and less commonly from the sphenoid or maxillary sinus. They occur with highest frequency in children under 6 years of age. Four different clinical grades of severity are distinguished, requiring a graded therapeutic approach. Ophthalmologic consultation should be obtained.

**Orbital edema:** The initial stage is marked by a doughy swelling and erythema of the eyelids (Fig. 3.46). Ocular mobility is normal, and the globe itself is not dis-
placed. The differential diagnosis should include dacryocystitis, the most common disease of the lacrimal sac, which is characterized by tenderness, erythema, and accompanying edema at the medial canthus of the eye. The treatment of choice is conservative and relies on nose drops (which may be applied in an intranasal cotton pack; see Treatment of Acute Sinusitis, p.55) and antibiotics. Corticosteroids may also be administered if required.

Periosteitis: While lid edema persists, pain develops at the medial canthus of the eye. Conservative medical treatment is still adequate at this stage in most cases.

Subperiosteal abscess: The inflammatory process has penetrated the bony barrier between the paranasal sinus and orbit, separating the orbital periosteum from the lamina papyracea and raising the pressure within the orbit. This stage is marked by an initial limitation of ocular movement with associated proptosis. Chemosis may already be present in exceptional cases. The treatment of choice is surgical drainage of the abscess, which today is done endoscopically. Nose drops and antibiotics are also administered. Ophthalmologic consultation is advised for medicolegal reasons.

Orbital cellulitis: The final stage of orbital complications is associated with proptosis and limited eye movements, pain, chemosis, and visual deterioration or even blindness.

Orbital cellulitis is a life-threatening emergency that requires immediate surgical decompression. This should be done with antibiotic coverage, as for a subperiosteal abscess.

An "orbital apex syndrome" can develop when the cellulitis spreads to involve the anatomic structures at the orbital apex (cranial nerves II–VI, ophthalmic
artery and vein). Progressive thrombophlebitis can also lead to cavernous sinus thrombosis and other intracranial complications (see below).

Bone and Soft-Tissue Inflammations (Osteitis and Osteomyelitis)

Osteomyelitis occurs mainly as a complication of frontal sinusitis when the bacterial inflammation spreads to the bony anterior wall of the frontal sinus, the frontal bone, and the surrounding soft tissues.

The main danger of frontal osteomyelitis is that the infection may spread to other bony structures of the calvaria.

Symptoms: The patient presents clinically with a tender, doughy, erythematous swelling over the forehead (Fig. 3.47). The surrounding facial soft tissues are also involved in most cases.

Diagnosis: Cranial CT scans should always be obtained to define the extent of the inflammation.

Treatment: The treatment of choice is surgical eradication of the affected bone under antibiotic coverage. Bone affected by inflammatory changes should be generously resected to prevent the further spread of inflammation.

Intracranial Complications

Intracranial complications also arise from the frontal sinuses in most cases. The ethmoid cells and sphenoid sinus are a more frequent nidus for these complications in children due to their lack of aeration.

Epidural, subdural and intracerebral abscesses: The clinical manifestations of these lesions are frequently nonspecific. As the disease progresses, it may produce signs of increased intracranial pressure with nausea, headache, vomiting, and occasional papilledema, somnolence, or seizures. The diagnosis relies critically on computed tomography (Fig. 3.48) or magnetic resonance imaging (Fig. 3.49) due to the nonspecific clinical features. The treatment of choice for the various types of abscess is surgical drainage with high-dose antibiotic coverage.

Meningitis: The main clinical manifestations are stiff neck, headache, fever, nausea, and photophobia. Some cases show increasing somnolence and clouding of consciousness, and seizures are occasionally observed. Cranial CT scans should be obtained whenever meningitis is suspected, particularly to exclude an intracranial abscess. The neurologist confirms the diagnosis by CSF sampling. In rhinogenic meningitis and with other forms, treatment relies mainly on surgical drainage of the affected sinuses under antibiotic coverage. Neurologic therapies may also be indicated (e.g., in patients prone to seizures).
Sinus Thrombosis and Thrombophlebitis

These complications, while rare, can lead to permanent neurologic deficits if diagnosed too late. They can be fatal in extreme cases.

Cavernous sinus thrombosis may present clinically with orbital edema, signs of venous congestion in the optic fundus (ophthalmologist!), chemosis, and occasional diplopia and proptosis with limited ocular movements. Magnetic resonance imaging should be performed and will often show definite evidence of sinus thrombosis. If magnetic resonance imaging (MRI) is unrewarding in patients with suggestive clinical signs, digital subtraction angiography (DSA) should be performed. As with all other intracranial complications, treatment consists of surgical drainage under antibiotic coverage.
3.9 Tumors of the External Nose and Face

The majority of facial tumors are malignancies, consisting mainly of basal cell carcinomas and spindle cell carcinomas. Other malignant neoplasms, precancerous lesions, and benign tumors are much less common in this region.

Benign Tumors

The most important benign facial tumor is *rhinophyma* (Fig. 3.50 a), a connective-tissue and sebaceous hyperplasia with angiectatic changes occurring over the cartilaginous nose. Most patients have preexisting rosacea, and so concomitant erythema is usually present. Although rosacea is more common in women than men, usually begins after age 20 and has a peak incidence at 40–50 years of age, rhinophyma is seen almost exclusively in older men. The *differential diagnosis* should include the cutaneous manifestations of lymphatic leukemia, cutaneous T-cell lymphoma, and sarcoidosis. The *treatment* of choice is surgical ablation of the hyperplastic tissue in layers, allowing the wound area to heal by spontaneous epithelialization (Fig. 3.50 b).

Precancerous Lesions

Besides *actinic keratosis* and *Bowen’s disease*, a chronic skin inflammation caused by carcinoma in situ, precancerous lesions include *cutaneous horn* and *malignant lentigo*. The latter is attributed to chronic sun exposure, grows slowly, and may progress to malignant melanoma.

Precancerous lesions of the facial skin should be watched closely, as they may progress to a malignant tumor.

Details on these precancerous lesions, which are generally rare, can be found in textbooks of dermatology.

Malignant Tumors

The most common facial malignancies are of epithelial origin, predominantly basal cell carcinomas and squamous cell carcinomas (spindle cell carcinomas). By contrast, melanomas, sarcomas, lymphomas, and cutaneous infiltration by leukemia are relatively rare in the facial region.

![Fig. 3.50 Rhinophyma](image_url)

Typical clinical appearance of rhinophyma before and after ablation of the hyperplastic tissue areas.

**Basal Cell Carcinoma**

*Synonym: basaloma*

*Epidemiology:* Basal cell carcinoma has a peak incidence between 60 and 70 years of age but may be seen in patients as young as 40.

*Etiology:* Uncertain. In addition to a genetic predisposition, prolonged sun exposure in people with very sun-sensitive skin appears to have causal significance.

*Clinical manifestations:* The tumor is classified as a malignant neoplasm because of its local invasiveness, but it has no tendency to metastasize. Basal cell carcinomas can vary greatly in their morphologic features. Solid basalomas are particularly rare in the facial region; they show central crusting and a string-of-beads margin (Fig. 3.51).

*Diagnosis and treatment:* After the diagnosis is confirmed by biopsy, treatment consists of surgical *excision* with frozen-section control of all margins. A special form, sclerodermiform basaloma, irregularly infiltrates the surrounding skin and often has ill-defined gross margins. This can lead to problems of surgical excision, as the size of the defect is often underestimated preoperatively. Following tumor removal with a margin of healthy tissue, the surgical defect is *reconstructed* in the same sitting (see 3.4, p.36).
Spindle Cell Carcinoma

Synonym: spinalioma

**Epidemiology:** Spindle cell carcinoma (Fig. 3.52a) is the second most common malignant tumor of the external nose and also tends to occur in older individuals.

**Etiology:** The etiology is uncertain, but it is very likely that exposure to ultraviolet radiation has causal significance.

**Clinical manifestations:** Unlike basal cell carcinoma, spindle cell carcinoma is a “classic” malignant tumor in that it can metastasize to regional lymph nodes.

**Treatment:** Treatment consists of surgical tumor removal. Various plastic reconstructive techniques can be used, depending on the location and size of the defect (Fig. 3.52a–d and Figs. 3.9–3.14, p.36). Patients with regional lymph-node metastases should undergo a neck dissection in the same sitting, followed by postoperative radiotherapy.
3.10  Tumors of the Nasal Cavity and Paranasal Sinuses

Benign tumors of the nasal cavity and paranasal sinuses are relatively rare. Malignancies of this region occur mainly in older patients and, since they develop in preexisting cavities, may remain asymptomatic for years.

**Benign Tumors**

Besides epithelial and connective-tissue neoplasms, benign intranasal and sinus tumors may arise from smooth muscle, peripheral nerves, or blood vessels.

**Inverted Papilloma**

Sinonasal papillomas, especially inverted papillomas, constitute a special category of epithelial masses. The inverted papilloma has certain characteristics that still prompt discussion on its etiology and appropriate management. It is a locally aggressive tumor, and transformation to squamous cell carcinoma is periodically described. Its growth characteristics resemble those of various virus-induced cutaneous and mucosal lesions (e.g., warts, condylomas, laryngeal papillomas). But while a viral etiology has been discussed, it remains unproved.

**Symptoms and diagnosis:** The clinical manifestations of inverted papilloma are as nonspecific as those of other sinonasal tumors and include nasal airway obstruction, headache, and occasional epistaxis. The lesion often has a polyp-like appearance when inspected by nasal endoscopy. In many cases only histologic examination can establish the diagnosis. Imaging (computed tomography) is helpful in defining the tumor extent.

**Treatment:** The treatment of choice is surgical removal. The special growth characteristics of this tumor require adequate exposure to allow for complete removal.

**Osteomas**

Osteomas are benign bone tumors that may occur as isolated masses, especially in the ethmoid cells and frontal sinus, or may form extensive masses that grow along the skull base (Fig. 3.53).

**Symptoms and diagnosis:** Many of these tumors are detected incidentally on x-ray films of the skull (Fig. 3.53a). Often they do not become symptomatic until they obstruct drainage tracts to or from the paranasal sinuses, leading secondarily to headaches and recurrent bouts of sinusitis. Computed tomography (Fig. 3.53b) is the imaging modality of choice for the accurate localization of osteomas.

For medicolegal and other reasons, it is essential to define the relationship of the osteoma to the skull base and other landmarks prior to surgical treatment.

**Treatment:** As soon as an osteoma becomes symptomatic, it should be surgically removed. Otherwise there is no need for therapeutic intervention.

**Malignant Tumors**

Malignant tumors of the nasal cavity and paranasal sinuses are far more common than benign masses. Histologically, the great majority (>80%) are tumors of the epithelial series (e.g., squamous cell carcinoma, adenocarcinoma, adenoid cystic carcinoma). Neoplasms of mesenchymal origin, such as osteosarcomas and chondrosarcomas, as well as malignant lymphomas are much less common. Metastases from other malignancies are occasionally found, with the primary tumor residing in the kidney, lung, breast, testis, or thyroid gland.

The main sites of predilection are the nasal cavity and maxillary sinus, followed by the ethmoid cells, frontal sinus, and sphenoid sinus.

**Symptoms:**

Because many tumors originate in the paranasal sinuses themselves, they often do not produce clinical manifestations until they have reached an advanced stage.

Symptoms that are suspicious for malignancy include sudden onset of **obstructed nasal breathing** combined with **bloody rhinorrhea** and a **fetid nasal odor**, especially in patients over 50 years of age. A malignant tumor should also be considered in the differential diagnosis of unilateral sinusitis that is refractory to treatment. Advanced tumor stages may be marked by swelling of the buccal soft tissues, swelling at the medial canthus of the eye, headache, facial pain, and hypoesthesia or numbness of the cheek due to infraorbital nerve involvement. Orbital infiltration can lead to displacement of the orbital contents, diplopia, or proptosis.
**Diagnosis:** The clinical examination includes endoscopic inspection of the nasal cavity (Fig. 3.54) and a search for regional lymph-node metastases by bimanual palpation of the cervical soft tissues. Since sinus tumors are apt to invade the nasal cavity secondarily, endoscopy alone may provide little information on the extent of the mass. For this reason, computed tomography and/or magnetic resonance imaging should always be performed (Fig. 3.55) and should cover the cervical soft tissues to check for nodal metastases. The disease can be staged based on the results of the examination. The principles of sinonasal tumor staging are outlined in Table 3.5.

**Treatment:** Treatment is individualized according to the histology and extent of the malignant tumor, and the treatment plan should be coordinated with the radiotherapist and medical oncologist. Since the great majority of lesions are squamous cell carcinomas, however, the treatment of choice will usually consist of surgery and postoperative radiation (Fig. 3.56). The goal of radical tumor removal may require a very extensive procedure with partial or complete removal of the maxilla or partial resection of the anterior skull base. As a result of close interdisciplinary cooperation with neurosurgeons, maxillofacial surgeons and ophthalmologists, as well as modern intensive-care options, even very extensive sinonasal malignancies can now be managed by surgical treatment. Since only about 20% of sinonasal malignancies metastasize to regional lymph nodes, a neck dissection is necessary only in patients who have clinically positive cervical nodes. Many of these cases will require postoperative radiotherapy (3.18).
Coronal computed tomography with a bone window demonstrates an extensive sinonasal carcinoma with sites of bone destruction in the anterior skull base and orbit (arrows).

Coronal magnetic resonance (MR) imaging in the same plane more clearly demonstrates the soft-tissue relationship of the tumor to the brain tissue and orbital contents.

Sagittal MR image.

Axial MR image.
Fig. 3.56 Melanoma of the nasal cavity and maxillary sinus

The tumor extends into the nasal vestibule, where it is visible to external inspection.

Corresponding axial computed tomography shows tumor tissue occupying the right maxillary sinus and nasal cavity.

At surgery, the tumor is exposed and resected through a lateral rhinotomy.

3.18 Esthesioneuroblastoma

Esthesioneuroblastoma is a rare neurogenic malignancy that arises from the sensory cells of the olfactory region and generally occurs in adults. Its etiology is still uncertain, but it is believed that some cases are embryogenically induced.

Clinically, esthesioneuroblastoma remains asymptomatic for some time due to its location in the olfactory groove between the upper portions of the nasal septum and the attachment of the middle turbinate. When advanced, the tumor causes obstructed nasal breathing, recurrent epistaxis, and particularly hyposmia or anosmia. Some of these tumors become symptomatic only after invading the cranial cavity or orbit, causing headache or visual deterioration. In a few cases, cervical lymph-node metastases are the primary manifestation of the disease.

Diagnosis is based on endoscopy (see image below) and especially computed tomography or magnetic resonance imaging; only these modalities can accurately define the tumor extent.

Treatment is based on a combination of tumor resection and postoperative radiotherapy.

Table 3.5 Principles for the staging of sinonasal tumors

<table>
<thead>
<tr>
<th>Regions</th>
<th>Subregions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nasal cavity</td>
<td>Nasal floor and roof</td>
</tr>
<tr>
<td>Upper level</td>
<td>Maxilloethmoid angle, ethmoid cells, sphenoid sinus, frontal sinus</td>
</tr>
<tr>
<td>Midlevel</td>
<td>Inferior, superior and medial portions of maxillary sinus</td>
</tr>
<tr>
<td>Tumor stage</td>
<td>Tumor extent</td>
</tr>
<tr>
<td>T1</td>
<td>1 subregion</td>
</tr>
<tr>
<td>T2</td>
<td>&gt; 1 subregion or 1 region</td>
</tr>
<tr>
<td>T3</td>
<td>Invasion of adjacent region</td>
</tr>
<tr>
<td>T4</td>
<td>Tumor crosses organ boundaries</td>
</tr>
</tbody>
</table>

* Tumor involves at least one of the regions listed.
4.1 Basic Anatomy and Physiology of the Lips and Oral Cavity

The lips and the soft tissues of the cheek function as the outer boundary of the oral vestibule and oral cavity, which form the initial part of the digestive tract. The tongue is situated such that the body of the tongue is within the oral cavity, while the base (root) of the tongue is in the oropharynx, forming its anterior boundary. For learning purposes, however, the tongue as a whole is included in the chapter on the oral cavity.

Functionally, the lips and oral cavity comprise the initial part of the upper digestive tract and thus play a key role in food ingestion. Speech production additionally developed during the course of phylogenesis. Finally, a large percentage of the taste receptors are located in the oral cavity.

Anatomy

Oral Vestibule

The oral vestibule is bounded externally by the lips and cheeks and internally by the alveolar processes and teeth (Fig. 4.1). When the teeth are in occlusion, the oral vestibule communicates with the oral cavity via a space behind the last molar. The oral cavity opens into the pharynx at the faucial isthmus (see 5.1, p. 98).

Lips and Cheeks

The lips and cheeks, the morphologic framework of which is formed largely by the mimetic muscles, are lined on their mucosal side by nonkeratinized squamous epithelium.

Lips: The longer upper lip and shorter lower lip are connected to each other by the labial commissures at the corners of the mouth. The lips are separated from the cheek by the nasolabial fold, an oblique sulcus that runs laterally and inferiorly from the nasal alae. The lamina propria of the lips contains numerous seromucous salivary glands (see Fig. 6.2, p. 133), the secretions from which drain into the oral vestibule. The orbicularis oris muscle forms the muscular foundation of the lips (Fig. 4.2). The lips receive their blood supply from the superior and inferior labial arteries, which arise from the facial artery. The lips are drained primarily by the facial vein, which also communicates with the orbital veins via the angular vein above the upper lip.

With inflammatory lesions of the lip (e.g., furuncles), infectious organisms can spread into the cranial cavity via connections between the orbital veins and cavernous plexus, resulting in complications (see 3.8, p. 59).

Knowledge about the lymphatic drainage of the lips is important for understanding the lymphogenous metastasis of malignant tumors of the lips. The submandibular and submental lymph nodes receive the lymphatic drainage from the lips.

The upper lip receives its sensory innervation from the infraorbital nerve, the lower lip from the mental nerve.

Cheeks: The cheeks, which form the lateral boundaries of the oral vestibule, also contain small salivary glands in their mucosa. The buccinator forms the muscular framework of the cheek. Like the orbicularis oculi, the buccinator is a mimetic muscle (Fig. 4.2) and receives its motor innervation from branches of the facial nerve. The Bichat fat pad (buccal fat pad) is located between the buccinator muscle and the overlying masseter muscle, the fibers of which run almost perpendicular to the buccinator. This fat pad smooths the cheek contour by filling in the depression at the anterior border of the masseter muscle. The excretory duct of the parotid gland runs through the buccinator muscle and opens into the mucosa of the cheek opposite the upper second molar.

Masticatory Muscles

The masseter muscle, located in the posterior part of the cheek, covers the vertical ramus of the mandible and the mandibular angle from the outside. It is one of the masticatory muscles, along with the temporalis muscle and the medial and lateral pterygoid muscles (Fig. 4.2). These muscles form both a functional and phylogenetic unit and accordingly are all supplied by the mandibular nerve (third division of the trigeminal nerve).
Teeth

The human dentition consists of two sets of teeth that vary in their individual shapes. The deciduous teeth are replaced by the permanent teeth, eight of which occupy each half of the maxilla and mandible:

- Two incisors
- One canine
- Two premolars
- Three molars

Each tooth consists of a crown and a root, which terminates at the apex. The area between the crown and root is the neck (cervix), which protrudes from sockets (dental alveoli) in the alveolar processes of the maxilla and mandible. The crown projects freely into the oral cavity and is covered externally by enamel. Internally, each tooth has a pulp chamber that contains connective tissue, nerve fibers, and blood vessels and is connected to the alveolus via the root canal. The teeth are anchored in the alveolus by the cementum, the bony alveolar wall itself, and the gingiva. These anchoring and supporting structures are known collectively as the periodontium.

The alveolar processes in the maxilla also form the floor of the maxillary sinuses—a fact that has major practical relevance.

Normally, the roots of the second premolar and first molar are very closely related to the maxillary sinus (see also Fig. 1.5, p. 4 and Fig. 1.6, p. 6). Occasionally, the bony shell between the periodontium and maxil-
lary sinus mucosa may even be absent in this region. The arteries that supply the maxilla and mandible (inferior alveolar artery, anterior and posterior superior alveolar arteries) arise from the maxillary artery. The upper teeth receive their innervation from branches of the maxillary nerve, the lower teeth from branches of the mandibular nerve.

**Oral Cavity**

The oral cavity is bounded anteriorly and laterally by the alveolar ridge and teeth, superiorly by the hard and soft palate (Fig. 4.1, p. 71), and posteriorly by the faucial isthmus. This narrow opening between the oral cavity and pharynx is bordered by the soft palate with the uvula and by the dorsum of the tongue at its junction with the tongue base.

**Palate:** The hard palate is formed by the palatine processes of the maxilla anteriorly, the incisive bone, and the horizontal plates of the palatine bones posteriorly. The oral cavity is sealed posteriorly by the soft palate with its pendulant process, the uvula. The palatal muscles that form the framework of the soft palate are the tensor veli palatini and especially the levator veli palatini, which elevates the soft palate during swallowing to keep food from entering the nose. The muscles of the soft palate are completed by the palatoglossus, which runs in the anterior faucial pillar (palatoglossal arch), and by the palatopharyngeus muscle of the posterior faucial pillar (palatopharyngeal arch, Fig. 4.1). The uvula also has its own muscle, called the muscle of the uvula. The palatal mucosa, like the mucosa of the lips and cheeks, contains numerous salivary glands (palatine glands, see also Fig. 6.2, p. 133). The motor innervation of the soft palate is described in 4.1. The palatal mucosa receives its sensory innervation from the greater and lesser palatine nerves (which arise from the second trigeminal nerve division; Fig. 4.1). The blood supply to the palate is derived from the ascending palatine branch of the facial artery.

**Tongue and oral floor:** The tongue is composed of various muscular systems (Fig. 4.3), occupies much of the oral cavity, and is continuous anteriorly and laterally with the floor of the mouth. The muscular foundation of the oral floor is formed by the mylohyoid muscle, which stretches between the anterior portions of the mandible. When the tip of the tongue is raised to expose the undersurface, the sublingual folds and sublingual papillae can be identified on both sides of the frenulum in the anterior part of the oral floor (Fig. 6.3, p. 133).

**4.1 Motor innervation of the soft palate**

Developmentally, the individual muscular components of the soft palate are derived from different structures and are therefore supplied by different cranial nerves (mainly by cranial nerves IX and X, see Fig. 16.3 and Fig. 16.4, p. 315 ff., and to a small degree by cranial nerve V). Cranial nerve deficits, especially those involving the lower cranial nerves IX and X, tend to restrict the mobility of the soft palate, causing difficulties in swallowing. During phonation (“ah”), the uvula and faucial pillars deviate toward the unaffected side.
The main anatomical subdivisions of the tongue are the apex, the body, and the base or root. The body of the tongue is separated from the base by a V-shaped groove called the terminal sulcus. The tip of this groove is directed toward the tongue base and is formed by the foramen cecum (Fig. 4.4).

The mucosa of the tongue differs from the rest of the intraoral mucosa chiefly by the presence of numerous papillae, which project from the surface of the tongue and give it its characteristic roughness. Four types of papillae are distinguished: filiform, fungiform, vallate, and foliate (Fig. 4.5). The latter three types are most important for taste perception. The microscopic taste buds are responsible for specific taste reception. They are most numerous on the vallate and foliate papillae and less so on the fungiform papillae. Small numbers of taste buds also occur in other regions of the oral cavity and pharynx (e.g., the soft palate, the anterior pillar, and the posterior wall of the oropharynx). Each taste bud consists of 30–80 elongated cells that extend superficially to the gustatory pore. This channel is located between the squamous epithelial cells and communicates with the oral cavity.

The lingual tonsil is part of the collection of lymphoepithelial tissue known as Waldeyer’s ring (see 5.1, p.101). The tongue and oral floor derive their blood supply from the lingual and sublingual arteries, which branch from the external carotid artery. Homonymous accompanying veins provide for drainage via the facial vein to the internal jugular vein.

Lymphatic drainage is handled by the ipsilateral and contralateral submandibular and submental lymph nodes, which drain to the lymph nodes at the junction of the facial and internal jugular veins (upper jugular lymph nodes, see Fig. 16.2, p.314).

The potential for contralateral lymphogenous spread should always be considered in patients with malignant tumors.

Developmentally, the tongue is derived from structures of the first through fourth branchial arches (see 16.5, p.317). This accounts for the complex innervation of the tongue, which is supplied in varying degrees by cranial nerves V, VII, IX, X, and XII (see also Figs. 16.3–16.6, p.315–317). The tongue derives its motor innervation from the hypoglossal nerve. The terminal sulcus receives its sensory supply from the lingual nerve, which branches from the third division of the trigeminal nerve (see Fig. 6.3, p.133). Sensation to the tongue base region is supplied by the glossopharyngeal and superior laryngeal nerves (from cranial nerve X). The taste buds (sensory innervation) are supplied by the chorda tympani (from cranial nerve VII) in the anterior two-thirds of the tongue and by the glossopharyngeal nerve in the posterior third. The central gustatory pathways are described in 4.2.
Physiology

Importance for Food Intake

The lips are the gateway to the digestive tract, sealing the oral cavity during chewing and swallowing to prevent the spillage of food. The orbicularis oculi muscle is chiefly responsible for this task. If the function of this muscle is impaired (e.g., due to facial nerve palsy), the resulting deficiency of lip closure can cause eating difficulties as well as drooling from the corners of the mouth at rest.

Within the oral cavity itself, the tongue has major functional importance as a “multifunction organ” with both motor and sensory properties. The complex motor functions of the tongue, like its other functions, can be traced to a specialized developmental history, which accounts for the sophisticated nerve supply derived from various cranial nerves (see p. 73 and Figs. 16.3–16.6, pp. 315–317) and for the specialized muscular structure of the tongue (Fig. 4.3, p. 72). The musculature of the tongue consists of extrinsic and intrinsic muscles. The extrinsic muscles are attached to the mandible and to the hyoid bone or styloid process, project into the body of the tongue, and greatly affect the position and movements of the tongue. The intrinsic muscles are composed of longitudinal, transverse, and vertical fiber systems and serve mainly to alter...
4.2 Central gustatory pathways

All gustatory fibers converge centrally in the area of the ipsilateral solitary tract, which ends at the solitary tract nucleus in the medulla oblongata; there the signal is relayed to the second neuron. The further course of the gustatory fibers is not yet fully understood. According to recent discoveries, the axons initially continue on to the medial parabrachial nucleus, where they synapse with the third neuron. Reportedly the fibers then travel via the dorsal trigeminothalamic tract, some crossed but most uncrossed, to the thalamus. The cortical taste areas themselves are located in the lateral part of the postcentral gyrus and in the adjacent insular cortex.

The molars have the greatest importance in chewing, because they are located closest to the insertion of the masticatory muscles themselves. This allows very high pressure to be developed between their occlusal surfaces.

Taste

There are only four basic taste sensations: sweet, sour, salty, and bitter. The sensory experience of “taste” is a much more complex phenomenon, however, and results from a combination of olfactory, thermal, mechanical, and sensory impressions.

The precise mechanism that triggers a taste sensation or transmits a gustatory signal at the molecular level remains unknown. Various theories have been advanced ranging from taste mediation by receptor proteins to taste activation through electrostatic interactions.

Importance in Phonation and Articulation

The musculature of the lips has an essential role in phonation, while “lingual articulation” controls the production of vowels, certain consonants, and palatal sounds through changes in the shape and position of the tongue. Finally, the oral cavity joins with the pharynx, nose, and paranasal sinuses (see 1.3, p.12 and 18.1, p.386) in forming the “supraglottic vocal tract,” which plays a role in the coordination of vocal sounds.
4.2 Methods of Examining the Lips and Oral Cavity

Inspection of the lips and oral cavity is an essential part of every otolaryngologic examination. Some problems that cannot be adequately investigated by clinical examination alone (e.g., taste disturbances, tumors) may require additional diagnostic procedures (e.g., taste tests, imaging studies). Disturbances of taste are rare but can be very distressful for the patient. They are reversible in many cases, depending on the cause (see Table 4.1).

Visual Inspection

After the lips have been inspected, the oral cavity is examined with the aid of a tongue blade. The examiner holds the instrument in the right hand while using the left hand to position and steady the patient’s head.

Dentures should be removed before the examination is started.

The sequence of the examination is shown in Fig. 4.6. Tongue mobility is assessed by having the patient stick out the tongue. With hypoglossal nerve palsy, the tongue will deviate toward the affected side (Fig. 4.7). Glossopharyngeal nerve palsy, in which the uvula and palatal arches deviate toward the healthy side (“backdrop sign,” Fig. 4.7 b), is excluded by assessing the mobility of the soft palate. This is done by watching the soft palate while the patient says “ah” several times.

Palpation

If inspection reveals questionable changes, the affected region or structure should next be palpated to better assess the consistency and depth of the suspicious finding. The cervical lymph nodes should also be palpated.

Taste Testing

Abnormalities of taste are classified as hypogeusia (diminished sense of taste), hypergeusia (increased sensitivity of taste), or ageusia (absence of the sense of taste).

Subjective Taste Testing

In the subjective test known as chemogustometry, aqueous solutions of glucose, NaCl, citric acid, and quinine are applied to the tongue in various concentrations to test the threshold of taste perception for the four basic qualities of sweet, salty, sour, and bitter. This test, while easy to perform, does not provide a high degree of reliability or reproducibility. Electrogonstometry is another subjective test procedure in which sensations are evoked by applying a constant anodal current to the taste receptors of the tongue.

Fig. Physical examination of the oral cavity

a Positions of the patient and examiner.
b The lips and cheeks are retracted from the teeth and alveolar ridge with a tongue blade to inspect the mucosa and assess the condition of the parotid duct orifice opposite the second upper molar.
c The patient elevates the tongue so that the examiner can evaluate the floor of the mouth and the submandibular duct orifices.
d The tongue is retracted with the blade so that the lateral oral floor can be examined.

This test has methodologic advantages over chemogustometry, providing better quantitative assessment of side-to-side differences and more accurate localization of responses.

Objective Taste Testing

Objective taste tests based on gustatory evoked potentials, for example (analogous to objective hearing and olfactory tests), are possible in principle but are very costly. They are practiced only at large centers and are used mainly in examinations for disability assessment.

Imaging Procedures

Since the anatomical structures of the oral cavity are easily accessible, the diagnosis can often be established by clinical examination alone (inspection, palpation, biopsy or local excision of a suspected tumor). As a result, imaging procedures tend to have a limited role in diseases of the lips and oral cavity. Nevertheless, there are various clinical situations (e.g., a tumor...
or extensive inflammatory process) in which a sectional imaging procedure can advance the diagnosis.

**Ultrasound**

Basically, only B-mode instruments are useful for ultrasound examinations of the oral floor and tongue (Fig. 4.8), and real-time scanning is preferred. Transducers with an operating frequency in the 5–10 MHz range are used, depending on the desired penetration depth and resolution. Newer systems have multifrequency transducers that operate at variable frequencies.

**Computed Tomography and Magnetic Resonance Imaging**

Normal findings are illustrated in Fig. 5.9 and Fig. 5.10 (pp. 106f.).

Computed tomography and magnetic resonance imaging are not only more cost-intensive than ultrasound but are also more invasive in cases where contrast media are used.

Indications:
- Pronounced inflammatory changes (e.g., Fig. 4.17, p. 84)
- Tumors. Information on tumor extent, depth of invasion, and spread across the midline are important parameters in selecting the optimum treatment modality, especially for lesions involving the tongue and floor of the mouth (see 5.4, p. 122).

Magnetic resonance imaging offers advantages over computed tomography in its superior soft-tissue discrimination.

If an imaging procedure is performed for confirmation (especially of a suspected tumor), the examination should include the soft tissues of the neck to check for regional lymph-node metastases.

**Table 4.1 Causes of taste disorders**

<table>
<thead>
<tr>
<th>Classification</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Congenital</td>
<td>Aplasia of the taste buds</td>
</tr>
<tr>
<td>Endocrine disorders</td>
<td>Diabetes mellitus, hypothyroidism, adrenal insufficiency</td>
</tr>
<tr>
<td>Drug side effects</td>
<td>E.g., D-penicillamine, various lipid-lowering drugs, ACE* inhibitors, antifungals</td>
</tr>
<tr>
<td>Peripheral nerve lesions</td>
<td>Involvement of the chorda tympani by facial nerve palsy, otitis media or previous middle ear surgery; involvement of cranial nerve IX by tumors or fractures of the skull base; very rarely after tonsillectomy</td>
</tr>
<tr>
<td>Radiotherapy</td>
<td>Radiation damage to the papillae</td>
</tr>
<tr>
<td>Exogenous chemical agents</td>
<td>Alcohol, nicotine, mouthwashes</td>
</tr>
<tr>
<td>Central taste disorders</td>
<td>E.g., head trauma, carbon monoxide poisoning</td>
</tr>
</tbody>
</table>

* ACE: angiotensin-converting enzyme.

**Fig. 4.7 Motor dysfunction of the tongue**

- a Hypoglossal nerve palsy on the left side
- b Glossopharyngeal nerve palsy on the left side

- a The tongue deviates toward the affected side when protruded.
- b The soft palate deviates toward the healthy side during phonation.

**Fig. 4.8 Ultrasound (B-mode) image of the tongue and oral floor**

Normal ultrasound anatomy of the oral floor. The transducer is placed submentally, resulting in an upside-down monitor image.
Malformations of the Lips and Oral Cavity

Malformations of the lips, cheeks, and oral cavity (especially those involving the palate and oral floor) are of epithelial origin. Although they are highly variable in their extent and clinical appearance, they are all based on a common teratogenic mechanism.

Cleft Lip and Palate

Epidemiology: Clefts of the lip, palate, and alveolar ridge can occur in various combinations. They are among the most common malformations, with an incidence of 1 in 500.

Classification: The following main groups are distinguished:
- Cleft lip and alveolar ridge
- Cleft lip, alveolar ridge, and palate and isolated cleft palate. The bifid uvula is a very mild variant of the cleft palate (Fig. 4.9).

Symptoms: Different clefts have a spectrum of clinical manifestations, depending on their morphology and extent:
- Hypernasal speech (rhinophonia aperta) due to incomplete closure of the nasopharynx
- Recurrent middle ear effusions and inflammations resulting from eustachian tube dysfunction
- Variable abnormalities of the nasal septum (septal deviation) or in the shape of the external nose

4.3 Pathogenesis of Cleft Lip and Palate

While the pathogenesis of cleft lip and palate is not yet fully understood, basically it involves a developmental anomaly of the embryonic head. In addition to genetic inheritance, clefting can result from a number of external influences, such as viral infections, placental oxygen deficiency, intrauterine bleeding, and exposure to ionizing radiation.

Fig. 4.9 Bifid uvula

The photograph shows the characteristic appearance of a median cleft in the uvula.
Diagnosis: Particularly with submucous clefts, the examination should include palpation of the hard palate in order to detect the bony discontinuity in that region.

Treatment: The adequate treatment of cleft lip and palate requires close interdisciplinary teamwork among the pediatrician, otolaryngologist, maxillofacial surgeon, orthognathic surgeon, and phoniatrist (see Fig. 4.10).
4.4 Inflammations of the Lips and Oral Cavity

Inflammatory diseases of the oral cavity and lips are often on a continuum and can have a variety of causes. An underlying systemic disorder is frequently present. Viral agents, bacteria, fungi, contact allergens, and various autoimmune diseases can incite inflammatory changes in this region, which predominantly affect the mucous membranes.

Viral Infections

Herpes Simplex Virus

**Epidemiology and pathogenesis:** Herpes simplex virus (HSV) infections of the oral mucosa are most often caused by HSV type 1 (cutaneous and oral-mucosa strain), while infections with HSV type 2 more commonly affect the genital region. The virus may be transmitted by contact or droplet infection or occasionally through superficial skin injuries. From 85% to 90% of the adult population are seropositive for HSV, particularly in urban areas.

**Symptoms:** Primary infection with HSV is usually acquired in early childhood and predominantly affects the oral mucosa as herpetic gingivostomatitis (aphthous stomatitis). The appearance of local lesions (bulla) on the oral mucosa is preceded by fever and lethargy consistent with a flulike infection. This is often accompanied by regional lymphadenitis. In rare cases, the nasal mucosa is also involved (herpetic rhinitis). Special clinical forms are reviewed in 4.5.

**Reactivation** of the HSV can occur in response to physical exertion, ultraviolet radiation, a febrile infection, emotional stress, or pregnancy (see also 4.6). Reactivation is most commonly manifested as herpes labialis. The site of predilection is the perioral region, especially the mucocutaneous junction of the lips (Fig. 4.11), but lesions sometimes occur in the mouth and nasal vestibule or on the cheeks, earlobes, or eyelids.

**Diagnosis:** The diagnosis is generally based on the history and clinical examination. Ordinarily there is no need for viral culturing or costly methods of viral identification (electron microscopy, immunofluorescence microscopy, polymerase chain reaction). A simple method is to demonstrate classic giant cells by the cytologic examination of a Tzanck smear.

**Complications:** A feared complication is secondary bacterial superinfection by *Staphylococcus aureus* or streptococci. Also known as herpes impetiginatus, this infection frequently heals by scarring, in contrast to non-superinfected cases (Fig. 4.12a). An occasional complication is postherpetic exudative erythema multiforme, characterized by skin lesions as well as typical

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**Figs. 4.11 Herpes simplex labialis**

Typical clinical appearance with vesicles about the upper and lower lip.

4.5 Severe forms of herpes simplex virus (HSV) infection

A particularly severe form of HSV infection, known as *Pospischill–Feyrter aphthoid*, can occur in immunocompromised children or as a sequel to measles, rubella, or chickenpox. A comparable form of this disease occurs much less frequently in immunocompromised adults and especially in HIV-infected patients.

A dreaded complication of primary HSV infection in children is herpetic meningoencephalitis.

4.6 Theories on the reactivation of herpes simplex virus (HSV) infection

The precise etiology of HSV reactivation is uncertain. Besides reinfection due to an exogenous cause, theories have focused mainly on an endogenous reactivation of the virus. The “precipitating factor” may be the integrated viral DNA in the host cells, which often is not detectable during latent periods but is able to induce the production of an “active” virus.

Another theory holds that the herpesviruses persist asymptomatically in the spinal cord, where they may be activated by any of a number of provocative mechanisms and then travel along sensory nerve fibers to corresponding sites on the skin or mucosa.
Ulcerative eruptions on the mucous membranes of the mouth, lips, and genitals (Fig. 4.12b).

**Treatment:** The treatment of herpes simplex labialis should include the use of topical antiseptics to prevent superinfection. Acyclovir, administered as a topical ointment or systemically, is available for severe forms of the disease. Therapy is generally continued for 5–7 days, but some cases (especially immunosuppressed patients) may require a more prolonged course of treatment.

**Varicella-Zoster Virus**

Pathogenesis: Chickenpox (varicella) and zoster are different clinical manifestations of infection with the varicella-zoster virus (VZV). **Chickenpox** occurs predominantly in children and results from primary infection with the varicella-zoster virus. After the cutaneous lesions have healed, the virus persists in the ganglion cells of sensory nerves. **Zoster** occurs as a reinfection or results from reactivation of the virus in response to various provocative mechanisms—ultraviolet radiation exposure, infectious diseases, or weakened immune defenses due, for example, to immunosuppressant therapy or human immuno-deficiency virus (HIV) infection; hence it requires previous contact with the virus.

Symptoms: **Chickenpox** presents with a characteristic skin rash consisting of erythematous papules and thin-walled vesicles with watery contents, covering the body but especially pronounced on the head and trunk. Aphtha-like vesicles also consistently appear on the oral mucosa and especially on the hard palate, buccal mucosa, and gingiva. **Zoster** presents clinically as a segmental disease, with cutaneous and mucosal lesions distributed along a sensory nerve segment and often accompanied by systemic signs such as lethargy, fatigue, and occasional neuralgiform pain in the distribution of the affected nerve. With involvement of the second and third divisions of the trigeminal nerve, aphthae or scalloped ulcerations can be found on the buccal mucosa, palate, and body of the tongue.

**Treatment:** Zoster, like HSV infection, should be treated with a 5–7-day course of acyclovir or famciclovir. Analgesics and anti-inflammatory drugs (especially carbamazepine) can also be beneficial, and antibiotics may be indicated in elderly or immuno-compromised patients to prevent superinfection. The efficacy of adjuvant cortisone therapy for zoster is a controversial issue.

**Herpangina**

Synonyms: vesicular pharyngitis, ulcerative pharyngitis

**Epidemiology and pathogenesis:** Herpangina is caused mainly by the group A coxsackievirus (CV), less commonly by the group B CV, and occasionally by retroviruses or echoviruses. The disease predominantly af-
ffects young children but also occurs in adults and is often manifested in the spring and fall.

**Symptoms:** Besides systemic symptoms such as fever, malaise, headache, and muscle pain, bullous eruptions surrounded by a red halo appear on the oral mucosa, particularly affecting the anterior faucial pillars (Fig. 4.13), uvula, and palatine tonsils. As a rule, the vesicles rupture in a few days, leaving behind shallow ulcerations.

**Differential diagnosis:** The gingivostomatitis caused by HSV (see above) is considerably more painful and runs a longer course.

**Treatment:** Treatment is purely symptomatic—anti-inflammatory agents or mouth rinses with chamomile. The disease generally resolves in 14 days without complications.

**Inflammatory Mucosal Lesions in HIV Infection**

Inflammatory lesions of the lips and oral cavity, while commonly observed in symptomatic HIV-infected patients, are not caused by the HIV itself but occur secondarily as a result of weakened host defenses.

**Candidiasis.** This disease, caused by *Candida albicans*, is the most common infection seen in HIV-positive patients (Fig. 4.14).

### 4.7 Hand-foot-mouth disease

Hand-foot-mouth disease is also caused by coxsackieviruses and predominantly affects small children from 6 months to 5 years of age. Clinically, small bullae typically appear simultaneously on the palate, tongue, and gingiva as well as on the palms of the hands, fingers, toes, and soles of the feet. As in herpangina, treatment is symptomatic. The disease generally resolves in 1–2 weeks without complications.
plex infection may have zoster-like manifestations) and to the course of the disease, which is markedly protracted in this subset of patients and is associated with more severe complaints (Fig. 4.15).

**Treatment**: Acyclovir is beneficial in HSV and VZV infections, but CMV is much less sensitive to this agent. CMV infections are treated with ganciclovir.

**Special form**: Another disease that has been linked to HIV infection since its initial description and is believed to have a viral cause is oral hairy leukoplakia (OHL). The presence of OHL is basically considered pathognomonic for an HIV infection. Today it is believed to be caused by the Epstein–Barr virus (EBV). The clinical presentation is marked by patchy, whitish, slightly raised lesions occurring predominantly on the border of the tongue (Fig. 4.16). Less commonly, the mucosal lesions are found in other regions of the oral cavity (buccal or lip mucosa, oral floor, soft palate). OHL typically runs a painless course, and dysphagia occurs only in cases with Candida superinfection. Despite their resemblance to leukoplakia (Fig. 4.21), the lesions have not been known to undergo malignant transformation.

The differential diagnosis should also include the various viral diseases of the oral mucosa such as herpes simplex (see p. 80), herpangina (see p. 81), and hand–foot–mouth disease (see 4.7, p. 72). The treatment of recurrent aphthous stomatitis is symptomatic and may include the frequent topical application of astrin- gents (tincture of myrrh) or mouth rinses with special pain-relieving electrolyte solutions (e.g., Hanks’ solution). Deficits can be corrected by means of iron, folic-acid and/or vitamin B₁₂ replacement. Chronic recurrent lesions may also benefit from the topical application of corticosteroid gel alternating with antiseptic mouth rinses.
Treatment consists of local measures such as the topical application of vitamin A acid and/or podophyllin. Virostatics should not be used due to the high incidence of side effects and the likelihood that OHL will recur within a few days after the drugs are discontinued.

HIV-infected patients are also predisposed to bacterial infections of the oral and pharyngeal mucosa. They have an increased incidence of acute and subacute tonsillitis as well as specific bacterial inflammations such as tuberculosis (see p.115), atypical mycobacterial infections, and syphilis (50% of HIV-infected patients test seropositive for syphilis; see 4.10).

Bacterial and Fungal Infections

Oral Floor Abscess

Synonym: Ludwig’s angina

Epidemiology: Oral floor abscess is a rare disease that can become potentially life-threatening if the inflammatory process spreads to the deep cervical soft tissues and mediastinum.

Pathogenesis: In many cases, the inflammation originates from the lower molars. Less commonly, the disease develops from mucosal injuries in the oral floor, leading to abscess formation in the tongue muscles or connective-tissue spaces of the oral floor. The disease can also develop as a sign of impaired host resistance, as in the case of diabetic or immunosuppressed patients (especially children).

Symptoms and diagnosis: An oral floor abscess is manifested clinically by edematous expansion with a firm, erythematous swelling in the submental to submandibular areas (Fig. 4.17 a). Patients complain of difficulty swallowing and speaking (“muffled speech”). High fever is also present. The downward spread of infection can lead to dyspnea with acute respiratory distress, and descending infection through the fascial compartments of the neck can incite a life-threatening mediastinitis. Imaging is necessary to define the extent of the oral floor abscess (often this cannot be done by physical examination alone due to local pain and induration). The principal options are ultrasonography and computed tomography (Fig. 4.17 b, c).

Fig. 4.16 Oral hairy leukoplakia

Typical whitish patches on the border of the tongue.

Fig. 4.17 Oral floor abscess (Ludwig’s angina)

a Typical clinical presentation, with erythematous swelling about the oral floor.
b, c Sequential axial computed tomography scans show the abscess, which originates from the oral floor (b) and extends downward (c).

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**Differential diagnosis:** See 4.9.

**Treatment:** The treatment of choice is incision and drainage of the abscess via the intraoral and transcutaneous route. Concomitant antibiotic therapy should be appropriate for a mixed spectrum of aerobic and anaerobic organisms.

**Lingual Abscess**

Overt or covert mucosal injuries to the tongue can become infected, resulting in a lingual abscess (Fig. 4.18). The diagnosis is established unequivocally by the clinical appearance of the tongue. **Treatment** is surgical and consists of incision and drainage of the abscess with concomitant antibiotic therapy.

**Candidiasis**

Candidiasis of the oral mucosa (synonym: thrush) occurs in persons with weakened host resistance due to radiation or cytostatic therapy, diabetes mellitus, long-term antibiotic use, corticosteroid inhalations, leukosis or leukopenia due to a different cause, or HIV infection.

**Symptoms:** Clinical examination of the oral cavity reveals whitish, firmly adherent plaques that can be scraped from the mucosa, leaving an erythematous, bleeding surface (Fig. 4.14, p. 82).

**Treatment:** Oral candidiasis is treated with topical antifungal agents such as nystatin solution or amphotericin-B lozenges. Every case should be treated, for otherwise the infection could spread to deeper portions of the alimentary tract (candida esophagitis) causing severe dysphagia, decreased food intake, and rapid weight loss.

**Differential diagnosis of oral floor abscess**

The differential diagnosis of a suspected oral floor abscess should include an abscess of the submandibular or sublingual glands as well as actinomycosis, in which subcutaneous infection by the bacterium Actinomyces israelii can cause an indurating infiltration of the fatty tissue in the oral floor region. The complaints differ from the symptoms of an oral floor abscess in that actinomycosis is less painful and tends to form an external fistula.

**Syphilis**

**Clinical features:** Although lesions of the oral and oropharyngeal mucosa can occur in all stages of syphilis, the manifestations in the primary and secondary stages are most important for the differential diagnosis of mucosal lesions in these regions. Three weeks pass until the primary lesion appears. The sites of predilection for extragenital primary lesions, after the perianal region, are the oral cavity and oropharynx. Besides the lips and buccal mucosa, the tonsil (unilateral) and tongue are most commonly affected. The primary chancre is painless and appears as an initially papular lesion that gives way to an erosive or ulcerative eruption. Concomitant regional lymphadenopathy (bubo) is frequently present and is also painless.

The secondary stage begins about 6 weeks after the primary lesion appears, as the disease becomes generalized due to hematogenous spread of the micro-organisms. The most commonly affected sites are the skin and mucous membranes. The mucosal syphilids (mucous plaques) are a dangerous source of infection, as they are teeming with infectious organisms.

The syphilitic enanthema in the secondary stage typically consists of patchy, reddish lesions on the hard and soft palate and buccal mucosa. An even more common finding is specific angina. In contrast to the unilateral tonsillar changes in the primary stage, both palatine tonsils are inflamed and covered with grayish-white coatings. A sweetly fetid breath odor is also present. A particularly severe form of secondary syphilis is malignant syphilis, which occurs predominantly in immunosuppressed and especially HIV-infected patients.

The tertiary stage may develop within a period of 3–5 years. Lesions of the oral and oropharyngeal mucosas are less common at this stage, but gummata (syphilitic granulomas) are occasionally found on the soft palate and uvula and also on the tonsil (unilateral). The tonsillar gummata appears as a sharply circumscribed ulcer with a greasy coating on its base and may initially be mistaken for a primary lesion, but the latter is almost always associated with painless regional lymphadenopathy. Interstitial glossitis is another, rare intraoral manifestation of tertiary syphilis.

**Differential diagnosis:** If specific angina is suspected, the differential diagnosis should include diphtheria, in which the mucosal lesions spread to involve the soft palate and uvula. It is important to differentiate interstitial glossitis from the innocuous fissured tongue (see Fig. 4.19, p. 87).

**Diagnosis:** Identification of the causative organism from the primary lesion can be accomplished with dark-field microscopy (the only reliable test in the early primary stage). Serologic tests such as the Treponema pallidum hemagglutination (TPHA) test or fluorescent treponemal antibody absorption (FTA-ABS) test are not positive until 3 weeks after the infection is acquired. A good follow-up test is the cardiolipin complement binding reaction (C8R), a modification of the Wassermann reaction whose titers correlate with the patient’s response to therapy.

**Treatment:** Penicillin G (600,000IU daily for 14 days) is the drug of choice for all stages of the disease. It is replaced by erythromycin in patients allergic to penicillin. Syphilis serology should be retested at the conclusion of treatment.
Superficial Tongue Lesions

Lesions in the surface of the tongue may reflect systemic diseases and thus provide important clues to the patient’s general state of health. Besides harmless morphologic variants and localized inflammatory lesions, therefore, medical disorders should always be considered in the differential diagnosis of these changes.

Hunter’s Glossitis

Synonyms: atrophic glossitis
Hunter’s glossitis is an atrophic inflammatory condition of the tongue base. It is an accompanying feature of pernicious anemia. Common symptoms are burning of the tongue, dry mouth, and altered sense of taste. Clinically, the tongue presents a typical smooth, shiny appearance with partial atrophy of the filiform papillae.

Geographic Tongue

Geographic tongue (synonym: benign migratory glossitis) is marked by areas of desquamation of the filiform papillae on the dorsal surface of the tongue. The affected areas are irregularly shaped but are clearly demarcated relative to surrounding areas. The disease is harmless, and histologic examination shows signs of inflammation. Generally, the only symptom is an occasional burning sensation. Treatment is unnecessary.

Black hairy tongue

Features: Black hairy tongue (synonym: lingua villosa nigra) is based on a hyperkeratosis of the filiform papillae, imparting a furry appearance to the tongue (Fig.).
Pathogenesis: The papillary elongation may result from failure of desquamation of the cornified layers or an excessive formation of keratin.
Etiology: Various precipitating mechanisms have been discussed. Besides antibiotic and corticosteroid use, they mainly include chronic mucosal irritation from oral hygiene procedures or nicotine abuse as well as metabolic disorders (e.g., diabetes mellitus), vitamin deficiency, and wasting diseases.
Treatment: Many cases are adequately managed by eliminating the causal factors or underlying disease.
Fissured Tongue

Fissured tongue, characterized by the presence of numerous furrows on the dorsal surface of the tongue, is a harmless hereditary condition that affects approximately 10–15% of the population. Lingual fissures may also be a sign of Melkersson–Rosenthal syndrome (see p. 294).

Clinically, the tongue presents a characteristic appearance (Fig. 4.19). The differential diagnosis should include interstitial glossitis in tertiary syphilis (see 4.10, p. 85).

Angioedema

Definition: Angioedema denotes a transient, frequently pronounced vascular reaction which, in the head and neck region, can lead to swelling of the face, lips, tongue, and larynx.

Pathogenesis: This disease occurs as one feature of an anaphylactic or anaphylactoid reaction. Drugs such as acetylsalicylic acid and angiotensin-converting enzyme (ACE) inhibitors are known to precipitate an attack. With ACE inhibitors, bradykinin appears to have a major pathogenic role. By contrast, the forms caused by a C1-esterase inhibitor (C1-INH) deficiency are much less common. Angioedema due to C1-INH deficiency may be hereditary or acquired.

Clinical manifestations: The disease is characterized by sometimes massive facial swelling that is most pronounced in the periorbital region but also affects the lips, tongue, tongue base, and laryngeal area. Massive tongue swelling in particular can cause acute obstruction of the upper airways. The hereditary form is additionally characterized by swelling of the extremities and episodes of abdominal pain.

Treatment: The cause of the angioedema is a key factor in selecting the appropriate treatment. The treatment of choice for angioedema not induced by a C1-INH deficiency is symptomatic treatment with corticosteroids or epinephrine (especially in the form of the disease induced by ACE inhibitors).

In the form that is induced by a C1-INH deficiency, direct replacement with a C1-inhibitor concentrate should be provided in acutely life-threatening cases with swelling of the tongue and larynx.

Antihistamines and cortisone preparations are of little or no benefit in this form of angioedema.

Immunologic Diseases

Systemic lupus erythematosus, see 4.13.
Pemphigus vulgaris, see 4.14.
Behçet’s disease, see 4.15.
Erythema multiforme, see 4.16.
Lichen planus, see 4.17.

Fixed Drug Eruption

A fixed drug eruption is a delayed (type IV) allergic reaction that occurs at the same cutaneous or mucosal sites (e.g., the extremities, soles of the feet, palms of the hands, external genitalia, oral mucosa) following repeated drug use. Particularly with mucosal involvement, the disease is characterized by superficial erosions that may resemble an HSV infection due to their scalloped margins. The eruption may be induced by analgesics, anti-inflammatory agents (e.g., pyrazolone,
4.14 Pemphigus vulgaris

Pathophysiology: Pemphigus vulgaris is an autoimmune disease characterized by the formation of antibodies directed against adhesion proteins in the epidermis. These antibodies can be detected in affected mucocutaneous areas and in the serum.

The etiology is uncertain. Besides a genetic disposition, the disease has been attributed to ultraviolet radiation exposure and various medications (e.g., phenylbutazone, indomethacin, ibuprofen, tuberculostatic drugs). The disease may also occur spontaneously, however, and it occasionally coexists with other autoimmune diseases (e.g., myasthenia gravis). Clinically, 50% of patients with pemphigus vulgaris show involvement of the oral mucosa with bullous eruptions or saliva-macerated bullae that can make eating extremely difficult.

Diagnosis relies mainly on the immunologic detection of pemphigus antibodies; elevated titers correlate with an exacerbation of symptoms. The diagnosis is most quickly established by the histologic or immunohistologic analysis of affected mucosal areas (see textbooks of dermatology).

Treatment: Initial treatment consists of systemic corticosteroids, whose use has led to a significant decline in the mortality of the disease. The addition of immunosuppressants is frequently indicated and can reduce the necessary corticosteroid dose. The most widely recommended local measures are oral rinses with anti-inflammatory or anesthetic solutions.

Prognosis: Untreated, the disease may lead to death within a period of months or years, generally due to secondary complications such as sepsis or bronchopneumonia.

4.15 Behçet’s disease

Behçet’s disease occurs predominantly in eastern Mediterranean countries (especially Turkey) and in Japan but is becoming more prevalent in Western urban areas as a result of immigration.

Etiology: A viral etiology has been suggested in addition to an autoimmune mechanism. The major features of the disease are oral aphthae, aphthous genital ulcers, and hypopyoniritis. The minor features are polyarthritis, gastrointestinal symptoms, and vascular lesions.

The diagnostic criteria for Behçet’s disease are either the presence of the three major features or the presence of two major and two minor features.

Treatment depends on disease severity and includes the use of corticosteroids. Cytostatics and immunosuppressants may also be required. Oral mucosal lesions can be treated locally with mouth rinses (chamomile) or pyoktanin (2%).

4.16 Erythema multiforme (EM)

Etiology and pathogenesis: Erythema multiforme has a multifactorial etiology. Known precipitating causes are viral infections such as HSV infections, hepatitis B, mumps, and measles. The main bacterial agents are streptococci infecting the upper respiratory tract. The disease may also occur in the setting of diphtheria or syphilis. Other causes are drug side effects (sulfonamides, pyrazolone derivatives, barbiturates, penicillins, phenothiazines). Systemic diseases such as polyarteritis nodosa, Wegener granulomatosis, systemic lupus erythematosus, and various malignancies (lymphoma, carcinoma) also have causal significance.

Signs and symptoms: A minor form of EM, marked initially by bulla formation on the oral mucosa and lips and later by erosive lesions (see Fig. 4.12b, p. 81), is distinguished clinically from a major form (Stevens–Johnson syndrome). Major EM runs a very severe course and may develop following a herpes infection or in response to certain drugs. Besides erythematous areas on the extremities and buttock, conspicuous mucosal erosions appear predominantly on the lips, oral mucosa, and pharynx. These lesions are very painful and interfere with eating. There may also be ocular involvement in the form of conjunctivitis, keratitis, iritis, or uveitis.

Systemic signs consist of generalized weakness, headache, and high fever. Severe cases may develop renal and cardiopulmonary disorders ranging to renal failure or toxic circulatory collapse.

Treatment: Due to the painful involvement of the oral mucosa, adequate food and fluid intake should be stressed (and may necessitate parenteral nutrition). Milder forms of the disease may respond satisfactorily to local treatments (rinsing with chamomile solution, local anesthetics if required). Corticosteroids are the agents of choice for the severe forms, accompanied by the administration of a broad-spectrum antibiotic to prevent superinfection.

Patients with frequent bouts of severe, postherpetic EM may benefit from long-term prophylaxis with acyclovir.
phenylbutazone, phenazone), antibiotics (penicillin, tetracyclines, erythromycin), chemotherapeutic agents, sulfonamides, and by certain hypnotics (e.g., barbiturates) and laxatives (phenolphthalein). Treatment consists of avoiding the suspicious substances.

4.17 Lichen planus (LP)

The etiology of lichen planus is unknown. Besides an immune pathogenesis in the setting of viral diseases (e.g., hepatitis B and C), there appears to be an association with certain medications (e.g., antimalarial drugs, organ arsenic compounds, gold salts). A psychosomatic mechanism is also likely, since lesions often appear following severe emotional trauma or other stressful situations.

Signs and symptoms: The disease often affects the skin and mucosae concomitantly, with mucosal involvement occurring in 25–70% of patients. Lesions are particularly common on the oral mucosa (fig.) and the vermilion border of the lips. Diagnosis can be difficult due to the variable clinical manifestations of the disease.

Oral lesions most typically appear as reticular white markings on the mucosa of the cheek (fig.) and tongue (Wickham’s phenomenon—requires histologic differentiation from leukoplakia). But painful ulcerations may also occur (erosive LP), requiring differentiation from the lesions of pemphigus vulgaris, systemic lupus erythematosus, and stage II syphilis.

Treatment: A variety of treatments have been recommended for LP ranging from antibiotics and tuberculostatics to antimalarial drugs. Most therapies tried to date have been unconvincing because they were not applied in a controlled-study framework. Corticosteroids are of symptomatic benefit, and aromatic retinoid and isotretinoin combined with corticosteroids are recommended for the treatment of mucosal lesions. Oral rinses with anti-inflammatory and local anesthetic solutions are recommended mainly for very painful, erosive intraoral lesions.

Prognosis of LP is guarded in terms of a complete recovery. It is common for oral mucosal lesions to persist for years, especially in the erosive form of the disease. Since it is now believed that LP is a potentially premalignant disease, regular follow-ups are also essential.

4.18 Burning mouth syndrome (BMS)

Burning mouth syndrome (BMS) is a symptom complex characterized by a burning sensation and other soreness in the oral cavity, often with an absence of objective mucosal findings. Although the tongue is most commonly affected (“burning tongue”), involvement of the hard palate, alveolar ridge (especially in denture wearers), and other regions of the oral cavity (buccal mucosa, oral floor, mucosal surfaces of the lips) has also been described. Concomitant xerostomia and dysgeusia are occasionally reported. BMS is most prevalent in postmenopausal women.

Causes:

- Local:
  - Dentures
  - Candidiasis
  - Geographic tongue
  - Allergic mucosal reactions (e.g., to sorbic acid, zimtaldehyde, nicotinic acid)
  - Toxic mucosal reactions (e.g., to nickel sulfate or mercury)
  - Radiotherapy
- Systemic:
  - Iron-deficiency anemia
  - Vitamin B12 deficiency
  - Vitamin B6, B2 and B6 deficiency
  - Folic acid deficiency
  - Sjögren disease
  - Menopause
  - Diabetes mellitus
  - Human immunodeficiency virus infection
  - Drug side effects (angiotensin-converting enzyme inhibitors)
- Psychogenic:
  - Depression
  - Cancerophobia
  - Emotional stress

Phenylbutazone, phenazone, antibiotics (penicillin, tetracyclines, erythromycin), chemotherapeutic agents, sulfonamides, and by certain hypnotics (e.g., barbiturates) and laxatives (phenolphthalein). Treatment consists of avoiding the suspicious substances.
4.5 Tumors of the Lips and Oral Cavity

Benign tumors of the lips and oral cavity, while rare, can occur as neoplasms of the various epithelial and mesenchymal tissues in this region. Among the precancerous lesions, leukoplakia is particularly important because of its morphologic similarity to carcinoma in situ. Malignant tumors of the lips and particularly the oral cavity have become more prevalent in past decades as a result of alcohol and nicotine abuse and today are counted among the most frequent head and neck malignancies along with pharyngeal and laryngeal cancers (see 5.3–5.5, pp. 108–111 and 17.7, pp. 368–378).

Benign Tumors

Benign tumors of the lips and oral cavity can arise from all epithelial and mesenchymal tissues in the head and neck region but are relatively rare. Besides papillomas (Fig. 4.20a) and pleomorphic adenomas (Fig. 4.20b), various mesenchymal tumors can occur such as fibromas, lipomas, rhabdomyomas, leiomyomas, and chondromas. There are also hemangiomas and lymphangiomas, which are congenital in most cases.

**Treatment:** Treatment is generally surgical and is indicated for patients who describe symptoms and in cases in which it is necessary to exclude a malignant tumor.

Hemangiomas and lymphangiomas are a special case. Due to the high rate of spontaneous remission during the first years of life, conventional surgical treatment or laser surgery is advised only if the tumor persists beyond that period, provided the patient does not have serious symptoms such as dyspnea or dysphagia that would necessitate earlier surgical intervention. Radiotherapy is no longer advocated for these tumors due to the potential for adverse sequelae (malignant transformation, growth disturbance).

Precancerous Lesions

Leukoplakia is the most common precancerous lesion of the lips and oral cavity (Fig. 4.21). Many of these lesions are asymptomatic and are detected incidentally. Exogenous irritants such as denture pressure or alcohol/nicotine abuse have been most strongly implicated as causal factors. Given their morphologic resemblance to carcinoma in situ and invasive carcinoma and their potential for malignant degeneration, leukoplakic lesions should always be investigated by biopsy. The treatment of choice is complete surgical removal of the neoplasm.

Bowen’s disease of the oral mucosa, a chronic inflammatory disease caused by an intraepidermal carcinoma, is rare by comparison. Its morphologic features are similar to those of leukoplakia.

Malignant Tumors

Malignant Tumors of the Lips

Malignant tumors of the lips (T categories Table 4.2, p. 93) are almost invariably squamous cell carcinomas and most commonly affect the lower lip (approximately 90% of cases). They occur predominantly in pipe smokers. Prolonged, intense sun exposure is considered a cofactor.
Symptoms and diagnosis: Early tumors often appear clinically as “intractable” ulcerations in the vermilion border of the lip (Figs. 4.22 a, 4.23 a) but may also consist of large, exophytic lesions (Fig. 4.22 b). Whenever a tumor is suspected, a biopsy should be taken to confirm the diagnosis.

Differential diagnosis: Differentiation is mainly required from keratoacanthoma and a primary syphilis chancre (see 4.10, p.85). Basal cell carcinoma (see Fig. 3.48, p.61) involves the vermilion border of the lip only by secondary spread.

Treatment: The treatment of choice is almost always surgical excision followed by a local primary closure or plastic repair of the defect using various reconstructive techniques (Fig. 4.22 c–g and Fig. 4.23 b–d). As a rule, even extensive tissue defects can be repaired using regional flap techniques. Carcinomas of the lip have an inherently low rate of metastasis to regional lymph nodes, but a neck dissection should be performed in patients with category 2 or higher tumors (see 16.5, p.332).

Malignant Tumors of the Oral Cavity

Squamous cell carcinomas also predominate in the oral mucosa and are variable in their clinical appearance (Fig. 4.24). Approximately 90% of patients have a long history of nicotine and alcohol abuse, and nearly 75% of malignant tumors form in the drainage area of the oral cavity—i.e., the trough between the base of the alveolar ridge and the border of the tongue (Figs. 4.24 a, 4.25 a).

Symptoms: Symptoms vary with the location and extent of the tumor and may consist of painful swallowing, blood-tinged saliva, and a fetid breath odor. Some tumors are completely asymptomatic, however.

Diagnosis: Visual inspection can raise the suspicion of a malignant neoplasm. This should be followed by bimanual palpation, since many tumors infiltrate deeper tissues and the visual impression of superficial findings can be misleading. The clinical examination also includes palpation of the regional cervical lymph nodes to exclude metastases. Imaging procedures (ultrasound, computed tomography, magnetic resonance imaging) are generally necessary only for extensive masses, as many tumors can be adequately evaluated clinically owing to their exposed location. But with more advanced lesions, imaging is valuable for defining the depth of tumor infiltration and assessing the involvement of adjacent structures (bone). It is also an important tool for excluding regional cervical lymph-node metastases.

Treatment: The treatment of choice in most cases is surgical removal of the primary tumor. The resulting defect is either closed primarily or reconstructed using pedicled flaps (see Fig. 4.25) or microvascular free transfers (e.g., a radial forearm flap). A unilateral or bilateral neck dissection (see p.334) may be necessary, depending on the location and T category of the primary tumor (see Table 4.2, p.93). Radiation to the tumor site and lymph areas is frequently indicated following surgery. Primary radiotherapy or combined radiochemotherapy may be considered as alternatives for T3 and T4 tumors.

Prognosis: The prognosis of oral malignancies depends on the location and stage of the disease. The five-year survival rate varies accordingly, ranging from 0% to 80%.
Fig. 4.22 Malignant tumors of the lower lip
Fig. 4.23 Malignant tumor of the upper lip

The clinical appearance of T1 (a) and T2 squamous cell carcinomas (b), which present different morphologic features. c The diagram illustrates a technique for reconstructing a postresection defect in the lower lip. d The flap is mobilized from the upper lip and transposed into the lower lip defect. e Appearance after flap inset and closure of the donor defect in the upper lip. f Good flap healing is seen 6 weeks after the operation. A “commissure plasty” can be performed at this stage to extend the obliterated left oral commissure. g Appearance 3 months after the first operation.

Table 4.2 T classification of malignant tumors of the lip, oral cavity, and oropharynx

<table>
<thead>
<tr>
<th>Largest tumor dimension</th>
<th>T1</th>
<th>T2</th>
<th>T3</th>
<th>T4</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>≤ 2 cm</td>
<td>&gt; 2 cm and &lt; 4 cm</td>
<td>≥ 4 cm but still superficial</td>
<td>Tumor of any size that invades deeper structures (e.g., bone)</td>
</tr>
</tbody>
</table>

**Table 4.2 T classification of malignant tumors of the lip, oral cavity, and oropharynx**

- a Ulcerated carcinoma of the upper lip, extending to the nasal base.
- b, c The resection defect is closed primarily by mobilizing a flap from the nasolabial fold.
- d One year later, the operative site is healed and free of irritation.
4.19 Malignant tumors of the oral cavity in patients infected with human immunodeficiency virus (HIV)

HIV-positive patients have a disproportionately high incidence of malignant tumors because of their weakened immune status. Most of these lesions in the oral cavity are **Kaposi sarcomas**. A smaller percentage are various types of **B-cell lymphoma**.

Kaposi sarcomas were first seen in association with HIV infection in the early 1980s. They are present in approximately 20% of affected homosexual and bisexual men but occur in less than 5% of HIV-infected individuals from other risk groups. The tumor has a variable appearance, depending on its location in the oral cavity. The hard palate is considered a site of predilection in this region (Fig. a). Fig. b illustrates a Kaposi sarcoma of the tongue.

The typical clinical appearance of squamous cell carcinoma of the oral floor (a), buccal mucosa (b), and soft palate (c).
The typical location of an oral floor carcinoma in the drainage area of the oral cavity between the alveolar ridge and border of the tongue. After resection of the tumor, which has infiltrated the tongue (b), a pedicled myofascial flap (here a myofascial pectoralis major flap; see also Fig. 3.12, p. 38) is outlined beneath the skin of the chest, mobilized, swung into the tissue defect, and sutured into place (c–e). 

One month later, granulation tissue is still present in the previous tumor defect that was reconstructed with the myofascial flap. 

The texture of the mucosa appears almost normal 2½ months after the operation.
Pharynx and Esophagus

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5.1 Anatomy, Physiology and Immunology of the Pharynx and Esophagus

The pharynx is a tubular, fibromuscular space extending from the skull base to the inlet of the esophagus (upper esophageal sphincter). Anatomically and clinically, the pharynx consists of a nasal part (nasopharynx), an oral part (oropharynx), and a laryngeal part (hypopharynx). The entire pharynx is bounded externally by several muscle systems, which perform diverse functions and are continuous distally with the muscles of the esophageal wall.

The primary function of the pharynx and esophagus is to coordinate the act of swallowing, which is regulated by a complex interaction of various cranial nerves and peripheral muscular and connective-tissue structures located in the oral cavity, pharynx, and esophagus. The pharynx also contains the tonsillar ring, a series of lymphoepithelial organs that are important in the immune response to infection. Finally, portions of the pharynx function as a variable resonance chamber for modulating vocal sounds.

Nasopharynx, Oropharynx, and Hypopharynx

Anatomical Extent

Nasopharynx: This highest part of the pharynx extends from the bony skull base to an imaginary horizontal line at the level of the velum (Fig. 5.1). It communicates with the nasal cavity via the choanae and with the middle ear via the orifice of the eustachian tube. The nasopharynx is bounded superiorly by the floor of the sphenoid sinus and pharyngeal roof. Also in this region is the pharyngeal tonsil, which forms part of the tonsillar ring (see below). Medial to the eustachian tube orifice, the tubal cartilage forms a projecting lip called the torus tubarius. The concavity behind it is termed the pharyngeal recess (Rosenmüller fossa) (see Fig. 5.8b, p.105). The nasopharynx is bounded posteriorly by the curve of the first cervical vertebra, with its overlying prevertebral cervical fascia and prevertebral musculature.

Oropharynx: The oral cavity communicates via the fauces isthmus (Fig. 4.1, p.71) with the oropharynx, which extends inferiorly from the lower boundary of the nasopharynx to the upper margin of the epiglottis (see Fig. 5.1). It is bounded anteriorly by the tongue base and lingual tonsil (Fig. 4.4, p.73) and posteriorly by the second and third cervical vertebrae with their prevertebral fascia. It is bounded laterally by the faucial pillars (tonsillar pillars, see 4.1, pp.70–75), which flank the palate tonsils.

Hypopharynx: The lowest pharyngeal segment is the hypopharynx, which extends from the superior border of the epiglottis to the inferior border of the cricoid cartilage plate of the larynx (Fig. 5.1), where it joins with the esophagus. Lying posterior to the hypopharynx are the third through sixth cervical vertebrae. Its anterior wall is formed by the back of the larynx, which protrudes into the hypopharynx and forms two lateral mucosal pouches (piriform sinuses), which rejoin at the level of the esophageal inlet.

Mucosal Lining

The mucosa that lines the nasopharynx consists of several rows of ciliated epithelium. At the oropharynx this gives way to a stratified, nonkeratinized squamous epithelium, which also lines the hypopharynx.

Pharyngeal Musculature

The muscular boundaries of the pharynx are formed by the constrictor pharyngis muscle group. The highest of these muscles, the constrictor pharyngis superior, begins at the level of the nasopharynx just below the tough, fibrous pharyngobasilar fascia, which in turn is suspended from the bony skull base. Just below the superior constrictor muscle are the overlapping constrictor pharyngis medius and inferior muscles, the latter of which joins distally with the esophageal musculature (Fig. 5.2).

While most of the constrictor pharyngis muscle fibers run obliquely, the lowest portions of the constrictor

5.1 Weak points in the wall of the hypopharynx

Three muscular weak points exist in the lower posterior wall of the hypopharynx. The first is the Killian triangle, located between the constrictor pharyngis inferior and the uppermost fibers of the cricopharyngeus muscle. The second area of weakness is the Killian–Jamieson region between the oblique and transverse fibers of the constrictor pharyngis. The third is the Laimer triangle, which is bounded above by the cricopharyngeus and below by the uppermost fibers of the esophageal musculature (see also Fig. 5.2). The Killian triangle is a particularly common site for the formation of hypopharyngeal diverticula.
pharyngis inferior (cricopharyngeal part) run directly horizontally, creating anatomical weak spots in the pharyngeal wall (Laimer and Killian triangles, 5.1). These weak spots are sites of predilection for the development of pulsion (Zenker) diverticula in the hypopharynx (see Fig. 5.28, p. 127).

Three additional pairs of external muscles are distributed to the pharyngeal wall and assist in controlling vertical movements of the pharynx: the stylopharyngeus, the salpingopharyngeus, and the palatopharyngeus (Fig. 5.2).

**Neurovascular Supply**

The pharynx receives its blood supply from the territory of the external carotid artery (branches of the facial artery, maxillary artery, ascending pharyngeal artery, lingual artery, and superior thyroid artery). The veins of the pharynx drain into the internal jugular vein. The lymphatic drainage of the upper portions of the pharynx is through the retropharyngeal lymph nodes, while the lower portions drain to the parapharyngeal or deep cervical nodes.
Nerve supply: The muscles and mucosa of the pharynx receive their motor and sensory innervation from the pharyngeal plexus, which in turn receives fibers from the glossopharyngeal and vagus nerves. The plexus itself is located on the outer aspect of the constrictor pharyngis medius muscle.

Parapharyngeal Space

The parapharyngeal space encompasses an anatomically well-defined region with the shape of an inverted pyramid whose base is formed by the inferior surface of the petrous bone and whose apex is at the lesser horn of the hyoid bone.

The parapharyngeal space is divided anatomically into two parts, the retropharyngeal space and the lateral pharyngeal space. The latter in turn is subdivided by the common connective-tissue sheath of the muscles arising from the stylohyoid process (stylopharyngeal aponeurosis) into a prestyloid and a retrostyloid part. The prestyloid part communicates with the parotid compartment. It contains the lateral and medial pterygoid muscles, lingual nerve, optic ganglion, and maxillary artery. Its lower part is directly adjacent to the tonsillar compartment. The retrostyloid part of the lateral pharyngeal space is traversed by neurovascular bundles made up of the internal carotid artery, internal jugular vein, and lower cranial nerves (IX–XII).

The retropharyngeal space contains smaller arterial and venous vessels and, most notably, the retropharyngeal lymph nodes that drain the nasopharynx.

Esophagus

The esophagus begins at the upper esophageal sphincter, located at the level of the C6 and C7 vertebrae (inferior border of the cricoid cartilage). The esophagus terminates at the gastric cardia in the plane of the T10 vertebra (Fig. 5.3a).

The three physiologic constrictions of the esophagus are clinically important due to the tendency for ingested foreign bodies to become lodged at those levels (Fig. 5.3b):

- **Upper constriction:** in the area of the esophageal inlet between the cricoid cartilage and the cricopharyngeal part of the constrictor pharyngis inferior muscle
- **Middle constriction:** where the aortic arch crosses over the tracheal bifurcation
- **Lower constriction:** where the esophagus pierces the diaphragm

The wall structure of the esophagus adheres to the pattern of the gastrointestinal tract as a whole, consisting of several layers:

- The mucosa, composed of stratified, nonkeratinized squamous epithelium

Fig. Anatomy of the esophagus

a The esophagus extends from C6/C7 to the gastric cardia at the level of the T10 vertebra.

b The three physiologic constrictions of the esophagus and their relationship to surrounding structures. The numerical scale shows distance in centimeters from the upper incisor teeth.
• The submucosa
• The muscularis, consisting of inner circular and outer longitudinal muscle fibers:
  – Upper fourth of the esophagus: striated fibers
  – Second fourth: mixed striated and smooth fibers
  – Lower half: smooth fibers
• The adventitia

Neurovascular Supply

Blood supply: The cervical part of the esophagus receives most of its blood supply from the inferior thyroid artery (and a lesser amount from branches of the subclavian and vertebral arteries). The thoracic esophagus is supplied by the aorta and intercostal arteries, and the abdominal esophagus by the left gastric artery and left inferior phrenic artery.

Venous blood in the neck is drained by the inferior thyroid vein. Thoracic and abdominal drainage is to the azygos and hemiazygos veins and esophageal veins. Lymphatic drainage is to the lymph nodes of the posterior mediastinum and pulmonary hilum.

Nerve supply: The upper, cervical part of the esophagus is supplied with branches from the recurrent nerve and the lower part with unnamed branches from the vagus nerve. Below the tracheal bifurcation is the esophageal plexus, formed by the two vagus nerves.

Physiology of Swallowing

Normal swallowing requires a coordinated interaction of various anatomic structures in the oral cavity, pharynx, larynx, and esophagus. From a functional standpoint, the voluntarily initiated oral phase of swallowing is distinguished from an “involuntary” pharyngeal phase and esophageal phase, which are controlled through reflex mechanisms (Fig. 5.4).

During the oral phase of swallowing, food is broken down and moistened to form a bolus that is moved toward the oropharynx. This is accomplished mainly by pressing the food against the hard palate with the tongue (Fig. 5.4, ➀).

The pharyngeal phase begins when the bolus comes into contact with receptors in the throat (especially on the tongue base), eliciting an involuntary swallowing reflex (➂). The afferent impulses for this reflex travel through the glossopharyngeal and vagus nerves, while the efferent neurons that supply the pharyngeal muscles arise from cranial nerves V3, VII, IX, X, and XII.

The extensive nerve supply highlights the complexity of swallowing as well as the potential vulnerability of this process.

While the involuntary swallowing reflex is triggered during the pharyngeal phase, the velum is elevated to close off the nasopharynx ➃. The larynx is also sealed off by elevation of the epiglottis ➄. This is accompa-
Within the lymphatic tissue, primary follicles are formed during embryonic development and differentiate into secondary follicles after birth (Fig. 5.5a). The secondary follicles mainly contain B lymphocytes at various stages of differentiation, along with scattered T lymphocytes (Fig. 5.5b–d). Besides the lymph follicles, there are also extrafollicular areas with B and T lymphocytes that enter the lymphatic tissue through the postcapillary venules.

**Functional Importance of the Tonsils in the Immune System**

The palatine tonsil in particular is considered to be an “immune organ” that plays a significant role in the defense against upper respiratory infections. By analogy with comparable lymphoepithelial tissue masses in the bronchi and intestinal tract, the lymphatic tissue in the tonsillar ring is also termed the mucosa-associated lymphatic tissue (MALT) of the upper respiratory tract. Accordingly, this tissue has the ability to mount specific immune reactions in response to various antigens. The activity of this lymphatic organ is especially pronounced during childhood, when immunologic challenges from the environment induce hyperplasia of the palatine tonsils (Fig. 5.6). Following this “active phase” of immune initiation, which lasts until about 8–10 years of age, the lymphatic tonsillar tissue becomes less important as an immune organ, and there is a corresponding decline in the density of lymphocytes in all regions of the tonsils. While the tonsils become less important immunologically with ageing, the tonsillar tissue continues to perform immune functions even at an advanced age, although this should not alter the decision to remove the tonsils if a valid indication for tonsillectomy exists (see Chronic Tonsillitis, p. 119).

While the tonsils are “learning” their immune function during childhood, extreme tonsillar hyperplasia (“kissing tonsils”) may develop, leading to functionally significant narrowing of the faucial isthmus, with eating difficulties and obstructed breathing. Especially...
when recumbent, these children may experience significant respiratory dysfunction, with periods of apnea. They also have an increased long-term risk of developing cor pulmonale. Consequently, there should be little hesitation in recommending tonsillectomy, even in small children.

**Phonation and Articulation**

Besides the oral cavity (see 4.1, Basic Anatomy and Physiology of the Lips and Oral Cavity, p. 70), the pharynx also functions as a variable resonance chamber for phonation and articulation.
5.2 Methods of Examining the Pharynx

Clinical examination of the pharynx is an essential part of any otolaryngologic examination and relies on various techniques. Besides the classic mirror examination, diagnostic endoscopic procedures have been increasingly utilized in recent years. Imaging procedures have also assumed major importance in the investigation of various pharyngeal disorders. This trend is due largely to the advent of computed tomography and magnetic resonance imaging, while conventional radiographs have become largely obsolete in the investigation of diseases of the pharynx. On the other hand, conventional radiographs are still an essential tool for the investigation of many esophageal disorders.

Mirror Examination and Endoscopy

Nasopharynx

The location of the nasopharynx can make it very difficult to access and examine, especially for beginners. Before the advent of endoscopy, the only technique available for examining the nasopharynx was posterior rhinoscopy (Fig. 5.7). The establishment of endoscopic techniques has dramatically improved the diagnostic evaluation of the nasopharynx.

Endoscopy: Nasopharyngeal endoscopy may be performed using a transoral or transnasal technique. The latter technique is described fully in 2.1 (pp.16–18) and permits the nasopharynx to be examined from the front. It can also provide detailed views of the eustachian tube region, the pharyngeal recess, and other difficult-to-reach sites (see p.98). Transoral endoscopy is basically a postrhinoscopic technique that provides the examiner with an excellent overview of the nasopharynx.

Transoral endoscopic technique: With the tongue pulled forward (to enlarge the space between the soft palate and posterior pharyngeal wall), the endoscope is introduced into the oral cavity over the left mandibular teeth and advanced past the uvula to the posterior wall of the pharynx (Fig. 5.8).

Oropharynx

Most structures of the oropharynx can also be evaluated during the examination of the oral cavity. The technique is shown in Fig. 4.6, p.76. The palatine tonsils are evaluated for their symmetry, mobility, and for the presence of any coatings or ulcerations. A laryngeal mirror or telescopic laryngoscope (see 17.2, pp.346–350) should be used to examine the tongue base and the lateral walls of the oropharynx.

Hypopharynx

Clinical examination of the hypopharynx (mirror examination, endoscopy) is performed concurrently with the examination of the larynx (see pp.346–350).

Esophagus

The esophagus can be examined by means of flexible or rigid endoscopy. Flexible esophagoscopy can be performed under local anesthesia, is generally well tolerated, and allows for concomitant examination of the stomach and duode-
num. It is the technique generally preferred by inter-
nists. Rigid esophagoscopy can also be performed under lo-
cal anesthesia in principle, but it is more comfortable 
for the patient and examiner to conduct the procedure 
under general endotracheal anesthesia. Rigid esopha-
goscopy provides a better overview, particularly when 
looking for foreign bodies, because the advancing rigid 
scope tends to flatten out the mucosal lining, making 
it easier to detect trapped foreign objects.

**Imaging Procedures**

**Conventional Radiographs**

In the area of conventional radiography, the oral con-
trast examination is the most valuable technique for 
diagnosing hypopharyngeal (see Fig. 5.28) and esopha-
geal diverticula, tumors, stenoses, and disorders of 
esophageal motility. Various contrast media can be used (e.g., barium, Gas-
trografin, Ultravist, Isovist), depending on the nature 
of the investigation and any preexisting disorders.

If there is a risk or suspicion of a perforation, bar-
um should not be used.

Another conventional radiographic technique, used 
mainly in patients with equivocal swallowing disor-
ders, is high-speed cineradiography. This technique 
can be used to evaluate the different phases of swal-
lowing with high temporal resolution (approximately 
50 images per second).

**Computed Tomography and 
Magnetic Resonance Imaging**

The modern sectional imaging modalities of comput-
ed tomography (CT) and magnetic resonance imaging 
(MRI) have significantly advanced the diagnosis of 
pharyngeal tumor masses as well as certain inflamma-
tory processes in this region. MRI (Fig. 5.9) has proven 
particularly effective for the soft-tissue discrimination 
of tumors in relation to surrounding structures, while 
CT (Fig. 5.10) is the method of choice for confirming or 
excluding osseous involvement.

---

Fig. Transoral endoscopy of the nasopharynx

a The tip of the endoscope has been positioned between the 
soft palate and the posterior wall of the pharynx in an anatom-
ical specimen.

b Normal view through a 90° endoscope, showing the posteri-
or ends of the turbinates, the posterior margin of the vomer, 
the eustachian tube orifice, and the torus tubarius.

c The same area viewed with a 120° endoscope.
Fig. Magnetic resonance imaging of the pharynx: normal findings

a) Nasopharynx: axial, without CM
b) Oropharynx: axial, with CM
c) Oropharynx: axial, with CM
d) Oropharynx: coronal, without CM
e) Oropharynx: coronal, with CM
f) Oropharynx: sagittal, with CM

Key:
- Masseter muscle
- Parotid gland
- Medial pterygoid muscle
- Torus tubarius
- Eustachian tube
- Genioglossus m.
- Longus colli m.
- Sphenoid sinus
- Nasopharynx
- Soft palate
- Oropharynx
- Mandible
- Oral floor musculature
- Masseter muscle
- Intrinsc tongue muscles
- Lateral pterygoid muscle
- Genioglossus muscles
- Geniohyoid muscle
- Tongue base
- Jugular vein
- Medial pterygoid muscle
- Sphenoid sinus
- Nasopharynx
- Soft palate
- Oral cavity and pharynx
The T1-weighted magnetic resonance images are labeled to indicate the image plane (axial, coronal, or sagittal) and whether the image was obtained with or without contrast medium (CM).
5.3 Diseases of the Nasopharynx

Diseases of the nasopharynx can have strikingly different causes at different ages. While adenoids are most common in children, tumors predominate in adults. Because lesions of the nasopharynx are difficult to examine and may produce nonspecific symptoms, malignant tumors in particular are apt to go undetected for some time. Modern endoscopic techniques using a high-intensity light source, together with the new sectional imaging modalities, have brought significant improvements, particularly in the early detection of nasopharyngeal lesions.

Adenoids

Synonyms: polyps, adenoid vegetations

“Adenoids,” the common term for hyperplasia of the pharyngeal tonsil, is a very widespread condition in children 3–6 years of age. The proliferation of lymphatic tissue in this region is so common in children that it can hardly be considered an abnormal condition, and nearly all children have some degree of adenoid hypertrophy due to the immunologic activity of that tissue. As a result, enlarged adenoids should be considered abnormal and treated accordingly only if they are causing symptoms. Not infrequently, the presence and severity of adenoidal symptoms depend on the relationship between the size of the nasopharynx and that of the adenoids.

Clinical manifestations: Common symptoms of adenoids are chronic nasal airway obstruction (“mouth breathing,” Fig. 5.11 a), nasal discharge (“runny nose”), snoring, anorexia, and a hyponasal voice (rhinophonia clausa). Also, many small patients have frequently recurring infections of the nose and paranasal sinuses with otitis media and chronic impairment of eustachian tube ventilation, caused for example by adenoid tissue obstructing the tubal orifices. Prolonged conductive hearing loss (see 9.1, pp. 198–201), especially during the first 3–4 years of life, can lead to delays in speech development. Finally, chronic mouth breathing can lead to maxillary deformity and dental malalignment. Many of these small patients also have enlarged tonsillar lymph nodes at the mandibular angle (Fig. 5.11 a).

Diagnosis: Besides posterior rhinoscopy or endoscopy (see pp. 17–18 and 104–107), the diagnostic workup includes microscopic examination of the tympanic membrane (otoscopy, see p. 166). Often this will show retraction of the tympanic membrane or a middle ear effusion resulting from chronic impairment of eustachian tube ventilation, with negative pressure in the middle ear. Additionally, hearing should be tested in adenoid patients (pure-tone audiogram, see p. 178; tympanometry, see p. 185; if necessary, otoacoustic emissions, see p. 189).

Treatment: The treatment of abnormally enlarged adenoids basically consists of surgical removal of the adenoids under general endotracheal anesthesia (adenotomy, adenoidectomy). In patients with concomitant middle ear effusion, paracentesis should be performed in the same sitting or a ventilation tube should be inserted for drainage (see p. 240).

Benign Tumors

Juvenile Angiofibroma

Epidemiology: Benign tumors of the nasopharynx are rare. The most common of these is juvenile angiofibroma, which accounts for less than 0.05% of all ear, nose, and throat (ENT) tumors and occurs exclusively in boys 10–18 years of age (Fig. 5.12).

Symptoms: Typical symptoms are obstructed nasal breathing, recurrent epistaxis, headache, impaired eustachian tube ventilation with middle ear effusion, and conductive hearing loss due to obstruction of the eustachian tube orifice.

Diagnosis: The typical endoscopic appearance is that of a well-circumscribed, vascularized mass (Fig. 5.12 a) with superficial vascular markings, situated in the nasopharynx or posterior part of the nasal cavity.

If there is clinical suspicion of an angiofibroma, a biopsy should not be performed due to the risk of heavy bleeding.

The primary workup should include MRI or CT, which can accurately define tumor extension into surrounding structures (Fig. 5.12 b). Digital subtraction angiography (DSA) is useful for identifying tumor-feeding vessels (Fig. 5.12 c).
Treatment: The treatment of choice is surgical removal of the tumor. Preoperative embolization of the feeding vessels (usually the maxillary artery) should be performed to reduce the intensity of intraoperative bleeding (Fig. 5.12d).

Malignant Tumors

Epidemiology: Carcinomas of squamous-cell origin account for the great majority of malignant nasopharyngeal tumors. A basic distinction is drawn between squamous cell carcinomas and lymphoepithelial carcinomas (Schmincke tumor). Much less common tumors of this region are adenocarcinoma, adenoid cystic carcinoma, malignant melanoma (Fig. 5.13), sarcoma, lymphoma, and plasmacytoma.

Etiology: The Epstein–Barr virus (EBV) appears to have a key role in the etiology of undifferentiated lymphoepithelial carcinoma.

Symptoms: Early symptoms of nasopharyngeal malignancies are unilateral conductive hearing loss with middle ear effusion.

Any persistent middle ear effusion of long duration in an adult patient with no prior history of middle ear disease is suspicious for a tumor and should be investigated accordingly.

Cervical lymph-node metastasis, usually involving the nodes at the mandibular angle, is another common initial finding. Features of advanced disease include nasal airway obstruction, recurrent epistaxis, headaches, and cranial nerve palsies.
Fig. Juvenile angiofibroma

**a** Endoscopy reveals a spherical, well-circumscribed tumor in the posterior portions of the right nasal cavity.

**b** Axial T1-weighted MRI after contrast administration demonstrates an enhancing mass in the nasopharynx.

**c** Digital subtraction angiography shows a well-vascularized tumor (arrows).

**d** After embolization of the feeding vessels, the tumor is no longer visible.
Diagnosis: The primary study is endoscopy of the nasopharynx (Fig. 5.14 a, b). Nasopharyngeal malignancies can have a variety of appearances ranging from a smooth, well-circumscribed tumor surface to mucosal ulcerations.

Some of these tumors are initially submucosal and are easily missed at endoscopy.

Otomicroscopy reveals unilateral tympanic membrane retraction and a middle ear effusion as a result of impaired eustachian tube ventilation. Given the EBV association of many nasopharyngeal cancers, the EBV antibody titer should be determined (this shows an elevated IgA, contrasting with the elevated IgM/ IgG that is found in infectious mononucleosis). MRI or CT is useful for defining tumor extent (Fig. 5.14 c).

Treatment: The treatment of choice for most nasopharyngeal carcinomas is primary high-voltage radiotherapy, because most of these tumors are very radiosensitive and the unfavorable tumor location and rapid invasion of the skull base preclude curative surgery in many cases.

Fig. Malignant melanoma

Right tubal orifice

Endoscopic view of a malignant melanoma in the nasopharynx.

Fig. Nasopharyngeal carcinoma

a Postrhinoscopic endoscopy demonstrates a mass that has obstructed the nasopharynx.

b In the transnasal endoscopic view, the tumor completely fills the choanae (only the right side is shown here).

c The corresponding axial T1-weighted magnetic resonance image after contrast administration shows the tumor and its extension into surrounding structures.
5.4 Diseases of the Oropharynx

The most common diseases of the oropharynx are inflammatory processes. Tumors, especially malignancies, are far less common in this region but should still be considered in the differential diagnosis, especially when certain risk factors are present (heavy smoking, alcohol abuse). Lesions of the oropharynx can also contribute to the development of sleep-related breathing disorders, particularly obstructive sleep apnea.

Injuries and Foreign Bodies

Scalds and Corrosive Injuries

Etiology: The accidental drinking of hot liquids by children can cause severe scalding of the lips, oral cavity, and oropharynx. Corrosive injuries are more common in adults due to the ingestion of caustic liquids with suicidal intent.

Symptoms: The dominant clinical symptoms are severe pain, especially on swallowing, and increased salivation.

Diagnosis: Initially the mucosa appears erythematous on mirror examination. Subsequent blistering may occur, followed by the formation of a whitish fibrin coating. Further tests are aimed at excluding injuries at lower levels of the alimentary tract and in the mediastinum. A chest radiograph should always be obtained (to check for mediastinal widening due to esophageal perforation). An early, careful endoscopic examination can be performed so that the extent of the esophageal injury can be accurately assessed.

Treatment: The initial treatment for scalds and corrosive injuries is to rinse the oral cavity with cold water. If the lips are affected, they should be treated with a corticosteroid-containing ointment. Patients with more severe injuries can additionally be treated with systemic corticosteroids, antibiotics, and analgesics.

A nasogastric feeding tube should be placed in patients with severe dysphagia who are unable to swallow.

Correct placement of the tube should be checked radiographically before the initial feeding, because the tube may perforate an esophagus that is affected by severe mucosal changes.

Foreign Bodies

Foreign bodies in the oropharynx are most commonly located in the tonsils and at the tongue base. Typical foreign objects are fish bones (Fig. 5.15) and bone fragments, usually with an obvious prior history of oral ingestion. Most patients describe well-localized pain on swallowing.

Treatment: The foreign material should be removed as soon as possible due to the risk of superinfection.

Fig. Foreign bodies in the oropharynx

a Clinical appearance of a fish bone lodged in the right tonsil (arrows).
b A fish bone lodged in the tongue base just above the vallecula (arrows).
Acute Inflammations

Acute Tonsillitis

Synonym: streptococcal angina

Definition, etiology: Acute tonsillitis is an acute bacterial inflammation of the palatine tonsils that is generally caused by group A β-hemolytic streptococci. Rare cases may be caused by staphylococci, *Haemophilus influenzae*, or pneumococci.

Symptoms: This disease is particularly common in children and adolescents and presents initially with high fever and severe pain on swallowing, which often radiates to the ear. Other symptoms are swollen tonsillar lymph nodes and muffling of speech due to oropharyngeal swelling.

Diagnosis:

**Mirror examination:** Both tonsils are swollen, bright red, and coated (Fig. 5.16).

**Inflammatory parameters:** The blood count shows leukocytosis, and the erythrocyte sedimentation rate (ESR) and C-reactive protein (CRP) are elevated.

**Bacteriologic testing:** A bacterial culture is rarely taken from throat smears because it usually takes 2–3 days to obtain a definitive result, by which time treatment should already be initiated. It is better to perform a rapid immunoassay, which can identify the causative organism as a group A streptococcus in just 10 minutes (5.2).

Fig. Acute tonsillitis

a Typical appearance of the palatine tonsils, which are bright red, swollen, and coated.

b The tonsils in this patient were so swollen that they caused respiratory distress, necessitating an immediate tonsillectomy.

5.2 Rapid streptococcal test

This rapid immunoassay makes use of colloid-labeled specific antibodies, which are placed onto reaction strips along with the pharyngeal smear. A color change in the “result window” indicates the presence of streptococcal group A antigen. The specificity and sensitivity of the various rapid tests available on the market range from 80% to 90%, making them useful tools in deciding whether to administer antibiotics. Note that a correlation exists between the test result and clinical findings—i.e., asymptomatic patients with a positive rapid test should not be placed on antibiotics. Conversely, a culture should be taken in cases where there is clinical suspicion of streptococcal tonsillitis but the rapid test is negative.
Treatment: The standard treatment for streptococcal tonsillitis is a 10–14-day course of penicillin V. This regimen should be continued for at least 7 days to avoid late complications (see below). Macrolides or oral cephalosporins can be used in patients allergic to penicillin. Analgesics are also administered for pain relief.

Complications: See 5.3 and Tonsillogenic Complications on p. 118.

Scarlet Fever

The tonsillitis in scarlet fever is also caused by infection with group A β-hemolytic streptococci. These are highly virulent bacterial strains that produce the scarlet fever exotoxin. Clinically, patients present with a rash that begins on the trunk. The area around the mouth is spared (“peri-oral pallor”). A pathognomonic feature is a bright red tongue with a glistening surface and hyperplastic papillae (“raspberry tongue,” Fig. 5.17). The tonsils are greatly swollen with a deep red color. Occasionally there is an enanthema of the soft palate with hemorrhagic areas. The diagnosis is established by the overall clinical picture combined with a positive rapid streptococcal test (see 5.2).

Medical therapy relies on penicillin, as in acute tonsillitis. Additionally, the oral cavity should be rinsed with mild antiseptic solutions, and analgesics should be given for pain.

5.3 Complications and sequelae of streptococcal tonsillitis

Lingual tonsillitis

In rare cases, the lingual tonsils may become inflamed and greatly swollen, and there may be concomitant edema involving the tongue base and laryngeal introitus. Endoscopic findings (Fig.) include marked hyperplasia of the lingual tonsils, which appear cylindrical with a stippled surface.

These patients may experience brief periods of progressive respiratory distress, requiring intubation. Patients with lingual tonsillitis should be hospitalized for observation and should receive high doses of antibiotics.

Streptococcal gingivostomatitis

In rare cases, tonsillitis may be followed or accompanied by streptococcal gingivostomatitis, characterized by diffuse inflammation and redness of the gingival mucosa and the formation of gingival abscesses. These lesions may also be seen at other sites on the oral mucosa and on the lips.

Sequelae of streptococcal tonsillitis

Rarely, a delayed-type antigen-antibody reaction can give rise to poststreptococcal diseases involving the kidneys (acute glomerulonephritis), major joints (acutereumatic fever), or heart (rheumatic endocarditis). Besides appropriate medical therapy, the treatment of choice is tonsillectomy under antibiotic coverage.

5.4 Complications of scarlet fever

A feared complication is necrotizing scarlet fever tonsillitis, which will cause extensive necrotic areas in the pharynx and oral cavity unless adequately treated. Septic complications can also arise, manifested by extensive soft-tissue infections and a toxic-shock-like syndrome. As in all infections with β-hemolytic streptococci, late sequelae can develop after an initial period of apparent recovery (rheumatic fever, diffuse hemorrhagic glomerulonephritis, and rheumatoid arthritis) (see 5.3).

Fig. Raspberry tongue

Typical clinical appearance of the tongue in scarlet fever. The bright red coloration and prominent papillae create a raspberry-like appearance.
Plaut–Vincent Angina

This inflammatory disease is caused by fusiform rods and spirochetes and presents clinically with unilateral dysphagia and a fetid breath odor with very little malaise. Mirror examination reveals a unilateral, fibrin-coated ulcer on the palatine tonsil.

Differential diagnosis: tonsillar carcinoma.

The causative organisms can be detected by the direct microscopic examination of a gram-stained smear.

Treatment: Local measures (cautery with 10% AgNO₃ or 5% chromic acid) are usually satisfactory, but should be supplemented by antibiotics (penicillin) in patients with more severe complaints.

Diphtheria

Epidemiology: Diphtheria was controlled for a time by active immunization, but lately its incidence has been rising due to low vaccination numbers, especially in immigrants from Eastern Europe, and secular fluctuations in the virulence of the toxin.

All instances of the disease must be reported to health officials.

Causative organism: The causative organism is Corynebacterium diphtheriae, which is transmitted by droplet inhalation or skin-to-skin contact. The incubation period is 1–5 days.

Pathogenesis: The bacterium produces a special endotoxin that causes epithelial cell necrosis and ulcerations.

Clinical manifestations: Two main forms are distinguished based on their clinical presentation:
- Local, benign pharyngeal diphtheria
- Primary toxic, malignant diphtheria

The disease begins with moderate fever and mild swallowing difficulties. The clinical picture becomes fully developed in approximately 24 hours, characterized by severe malaise, headache, and nausea.

Diagnosis: Mirror examination of the pharynx reveals typical grayish-yellow pseudomembranes that are firmly adherent to the tonsils and may spread to the palate and pharynx. The underlying tissue bleeds when the coatings are removed. A slightly sweet breath smell is also characteristic. The diagnosis is confirmed by the overall clinical impression, combined with smear findings.

Treatment: First, the patient should be isolated. Whenever diphtheria is suspected, even before it is confirmed by smear results, diphtheria antitoxin (200–1000 IU/kg body weight) should be administered by intravenous or intramuscular injection.

Allergy to the antitoxin should be excluded (with a skin test) before it is administered.

Penicillin G should also be administered.

Discharge from the hospital is contingent upon test results: three smears taken at 1-week intervals must all be negative. Two percent of patients continue to carry the bacterium and should undergo tonsillectomy.

Complications: Dangerous complications, which occur mainly in association with the primary toxic malignant form, are toxic myocarditis (which may terminate fatally in 10–14 days) and interstitial nephritis. The more severe the diphtheria, the earlier these complications may arise. Electrocardiography and urinalysis follow-ups should be continued for at least 6 weeks after the onset of the disease.

Tuberculosis

Epidemiology: Oral or oropharyngeal manifestations of tuberculosis most commonly occur in the setting of advanced organ tuberculosis. Although these lesions are very rare (0.2% of patients with organ tuberculosis), they should be considered in the differential diagnosis since the incidence of tuberculosis has been on the rise. It is even less common to see oropharyngeal involvement by a primary complex or in the setting of miliary tuberculosis.

Clinical manifestations:
Primary complex: A primary tuberculous complex in the tonsillar and cervical lymph-node region is most common in children who have become infected by drinking cow’s milk contaminated with tubercle bacilli. The primary complex in these cases consists of a typical ulcerative lesion of the oral mucosa and tonsil, associated with regional cervical lymphadenopathy. The swelling in the neck leads most patients to seek medical attention.

Organ tuberculosis with ulcerative mucocutaneous lesions occurs mainly in regions that may come into contact with secretions containing infectious organisms, resulting in the formation of ulcerative mucosal lesions that are sometimes necrotic. (Other forms of organ tuberculosis can affect the lung, bowel, etc.) Morphologically, the lesions may appear as mucosal ulcerations on the lips and dorsum of the tongue or as slightly raised, nodular eruptions on the palate. Skeletal involvement is also occasionally seen due to hematogenous spread. In this case “cold abscesses” may form about the cervical spine (Fig. 5.18), causing the posterior wall of the pharynx to bulge forward and
Miliary tuberculosis: Involvement of the oral mucosa can result from hematogenous spread, appearing as multiple pinhead-size papules, some hemorrhagic, that form on the oral mucosa.

Diagnosis: The diagnosis is established by the detection of acid-fast rods in smears, sputum, bronchial secretions, gastric juice, or biopsy material. The diagnostic workup should include biplane chest radiographs to check for pulmonary involvement. The tuberculin skin test is also performed to assess the reactivity of the organism to tubercle bacilli. Calcifications detected by ultrasound in enlarged cervical lymph nodes are pathognomonic for tuberculosis. If the result is equivocal, a cervical lymph-node biopsy should be taken for a histologic and bacteriologic tissue analysis.

Treatment: Inpatient antituberculous polychemotherapy is required, consisting either of a triple regimen (isoniazid, ethambutol, rifampicin) or a quadruple regimen with pyrazinamide added.

Acute Viral Pharyngitis

Etiology, symptoms: Acute viral pharyngitis, which is often caused by influenza or parainfluenza viruses, typically presents clinically with sudden onset of fever, sore throat, and headache. There may also be coughing and catarrhal symptoms (e.g., rhinitis, sinusitis). Concomitant cervical adenopathy may also be present.

Diagnosis: The pharyngeal mucosa appears red and coated on mirror examination. If a bacterial etiology is suspected, a rapid streptococcal test can be performed (see 5.2, p.113).

Treatment is supportive and consists mainly of analgesic agents. Cold compresses to the neck can also help to relieve pain. The patient should drink copious amounts of warm liquid to ease complaints.

Infectious Mononucleosis

Synonyms: Pfeiffer’s glandular fever, kissing disease

Causative organism: Infectious mononucleosis is caused by infection with the Epstein–Barr virus (EBV). It predominantly affects adolescents and young adults. The incubation period is 7–9 days.

Clinical manifestations: Although infectious mononucleosis is a systemic illness, it is common to encounter tonsillitis as the initial or cardinal symptom. Besides systemic symptoms such as fatigue, anorexia, and moderate temperature elevation (38–39°C), patients complain of severe pain on swallowing, headache, and limb pains.

Diagnosis:

Clinical examination: The tonsillar and nuchal lymph nodes, axillary nodes, and inguinal nodes are palpably enlarged. Often there is concomitant enlargement of the spleen and liver.
On mirror examination, the tonsils are found to be bright red, swollen, and covered with a grayish fibrin coating (Fig. 5.20).

Laboratory tests: The blood count initially shows leukopenia, followed later by leukocytosis (20,000/μL) with 80–90% atypical lymphocytes (lymphomonocytoid cells, Pfeiffer cells). EBV serology (especially IgM and IgG) is another important test. The enzyme-linked immunosorbent assay (ELISA) can confirm infectious mononucleosis by quantitatively detecting antibodies against the various EBV antigens (virus capsid antigen, early antigen, Epstein–Barr nuclear antigen). Rapid mononucleosis tests are also available but are less sensitive and specific than ELISA.

The serum hepatic enzymes should be determined to exclude concomitant involvement of the liver or spleen.

Upper abdominal ultrasound and an electrocardiogram are also recommended.

Treatment: Treatment centers on the symptomatic relief of pain and fever. The agents of choice for pain relief are acetaminophen or ibuprofen. Aspirin products should not be used, as they could cause bleeding problems if tonsillectomy is required. Antibiotics (penicillin V) should be given only if signs of bacterial superinfection are present.

Ampicillin and amoxicillin should be avoided because they frequently induce a pseudoallergic rash (Fig. 5.21).

In cases of infectious mononucleosis that run a severe course with persistent fever, respiratory distress or stridor, a tonsillectomy can expedite recovery by eliminating the focus of greatest viral proliferation.
Complications: Complications are rare and consist mainly of myocarditis, hemorrhage, nephritis, hepatitis, meningitis, or encephalitis.

Tonsillogenic Complications

Peritonsillar abscess: Peritonsillar abscess is a unilateral inflammatory process that involves not only the tonsillar parenchyma but also the peritonsillar tissue—i.e., the abscess spreads past the tonsil to involve the connective tissue between the parenchyma and pharyngeal musculature. The clinical features are pronounced unilateral redness and swelling of the soft palate (Fig. 5.19), muffled speech, and possible trismus. This is frequently accompanied by uvular edema, but the swelling may also spread to the tongue base and lateral pharyngeal wall, causing respiratory complications.

The treatment of choice is removal or incision of the affected tonsil under antibiotic coverage, bearing in mind that most patients harbor a mixed spectrum of aerobic and anaerobic organisms. Other complications: see 5.5.

[Fig. Pseudoallergic rash in infectious mononucleosis]

A pseudoallergic rash developed in this patient following treatment with ampicillin.
Chronic Inflammations

Chronic Pharyngitis

**Etiology:** Chronic pharyngitis is often a result of long-term exposure to various noxious agents (nicotine, alcohol, chemicals, gaseous irritants). It can also occur as a result of chronic mouth breathing due to nasal airway obstruction (e.g., deviated septum) or as an accompanying feature of chronic sinusitis.

**Symptoms:** The main clinical manifestations are a dry-throat sensation with frequent throat clearing and the drainage of a viscous mucus. Some patients have a dry cough and a foreign-body sensation in the pharynx.

**Diagnosis:** The history will often direct attention to possible noxious agents. On mirror examination, the pharyngeal mucosa appears red and “grainy” due to the hyperplasia of lymphatic tissue on the posterior pharyngeal wall (*hypertrophic form*: Fig. 5.22). The pharyngeal mucosa may also have a smooth, shiny appearance in some cases (*atrophic form*).

A thorough nasal examination should be performed to exclude nasal airway obstruction as the cause of chronic pharyngitis, giving particular attention to possible septal deviation or turbinate hyperplasia. The middle meatus should also be examined endoscopically (see 1.3, Anatomy of the Ostiomeatal Unit, p. 7).

**Treatment:** Any agents causing the pharyngitis should be avoided. Also, an herbal product such as sage or chamomile can be used in a steam inhalation to moisten the airways. In patients with nasal airway obstruction due to septal deviation or turbinate hyperplasia, a surgical procedure can be performed to improve complaints.

Chronic Tonsillitis

**Pathogenesis:** Like infections confined to the tonsillar crypts, recurrent inflammations of the tonsils and peritonsillary tissue can lead to permanent structural changes with scarring. Bacteria that grow on cellular debris in poorly drained crypts can perpetuate a smoldering inflammation, chronic tonsillitis. In this condition the palatine tonsils provide a “focus” that can sustain a variety of diseases in other parts of the body (rheumatic fever, glomerulonephritis, iritis, psoriasis, inflammatory heart disease, pustulosis palmaris and plantaris, erythema nodosum).

**Symptoms:** Chronic tonsillitis may cause recurrent episodes of pain or may run an asymptomatic course. The most frequent complaints are lethargy, poor appetite, a bad taste in the mouth, and a fetid breath odor.

**Diagnosis:** Mirror examination often reveals small, firm, immobile tonsils with associated peritonsillar redness. Occasionally a purulent liquid can be expressed from the crypts. *Tonsillar smears* are found to contain group A β-hemolytic streptococci. Palpation: The tonsillar lymph nodes at the mandibular angle may be enlarged. **Laboratory tests:** An elevated ESR and CRP and a left shift in the differential blood count are present as signs of the inflammatory process. An *antistreptolysin titer* of approximately 400 IU/mL or higher is considered pathologic.

**Treatment:** The treatment of choice is *tonsillectomy*, which is performed under general endotracheal anesthesia with the head hyperextended. The tonsil is exposed by incision of the anterior faucial pillar, shelled out along the connective-tissue plane between the parenchyma and pharyngeal muscle, and detached at its inferior pole.

Heavy postoperative bleeding may occur on the day of the tonsillectomy, during the first week after the operation, or even later in rare instances.
Peripheral Obstructive Sleep Apnea Syndrome (OSAS)

Obstructive sleep apnea syndrome (OSAS), like snoring, is a type of sleep-related breathing disorder that can have serious health effects and social consequences. The history can provide important clues to the presence of OSAS (Table 5.1).

**Etiology and pathogenesis** (Fig. 5.23): There is a tendency for the velum, oropharynx, and/or hypopharynx to collapse during sleep, narrowing the pathway for airflow and causing periods of apnea or hypopnea that can last up to 2 minutes. This leads to frequent arousals from sleep and gasping for air, preventing a normal sleep pattern. Besides disturbing the sleep–wake rhythm, OSAS can have longer-term effects due to a reduction in blood oxygen levels, with a potential for significant damage to the cardiopulmonary system. Factors that narrow the pharyngeal airway or lead to decreased muscle tone (Table 5.2) can promote or intensify the disease process.

**Symptoms**: Typical symptoms of OSAS are morning lethargy and daytime fatigue, with a tendency to fall asleep during the day. Witnesses additionally report irregular snoring with periods of apnea followed by “gasping” and loud snoring. Obesity is usually present as an accompanying condition.

**Diagnosis**: Mirror examination may demonstrate an elongated uvula, a narrow velopharyngeal passage, and a bulky soft palate with a small oropharyngeal lumen. It is also common to find a hyperplastic tongue base and hyperplasia of the palatine tonsils. The nasal

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**Table 5.1 Signs in the patient’s history that are suggestive of (obstructive) sleep apnea**

- Loud, irregular snoring
- Periods of apnea during sleep (witnessed)
- Unusual daytime sleepiness or fatigue
- Restless sleep
- Intellectual deterioration (poor concentration and impaired memory)
- Personality changes
- Loss of libido, impotence
- Nycturia, enuresis

Source: adapted from Günther, see p. 430.

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Flexible transnasal endoscopy can be used to perform a functional test (Müller maneuver) that is helpful in assessing the degree of oropharyngeal obstruction at the level of the soft palate.

Technique: The patient sits in an upright or 45° reclined position. With the endoscope in place, the examiner compresses the patient’s nostrils and instructs the patient to inhale the residual intraoral air while keeping the mouth closed.

Result: The negative inspiratory pressure during the Müller maneuver produces various collapse effects in the pharynx when OSAS is present, and these effects can be observed endoscopically. When a healthy patient performs this maneuver, it causes very little decrease in the cross-sectional area of the pharyngeal lumen.

Objective measuring techniques:

Screening: Reports given by the patient and witnesses can be objectified by recording O₂ saturation, respiratory sounds, and heart rate on an outpatient basis during sleep. However, these screening devices alone (e.g., the Madaus Electronic Sleep Apnea Monitor, MESAM) cannot provide an accurate sleep evaluation, since they do not at present include an electroencephalography (EEG) channel.

Confirming the diagnosis: The current gold standard for confirming OSAS and differentiating it from other sleep-related breathing disorders is polysomnography. Conducted as an inpatient procedure in a sleep laboratory, it measures thoracic and abdominal respiratory excursions, transcutaneous Po₂, and records an EEG in addition to the usual screening parameters.

Treatment: General treatment measures consist of weight reduction, abstinence from alcohol and nicotine, and avoiding big meals, especially at night. It is also important to establish a regular sleep–wake cycle and avoid the use of sedatives.

One nonsurgical treatment option is the Esmarch splint, an occlusive splint that advances the lower jaw. By moving the tongue base and adjacent pharynx forward, this device widens the airway in the unstable portion of the oropharynx.

In patients with severe grades of OSAS or an unsuccessful trial with an occlusive splint, the unstable portions of the airway can be “pneumatically splinted” by means of transnasal continuous positive pressure ventilation; this keeps the tissues from collapsing during sleep and obstructing airflow. A nasal continuous positive airway pressure (CPAP) mask for this purpose is custom-fitted at the sleep laboratory.

Surgical treatment options are tailored to the specific pathology causing the apnea.

Surgical treatment requires very careful patient selection, because many patients will derive little or no benefit from the operation.

The result of the Müller maneuver can be helpful in selecting patients for a surgical procedure on the soft palate.

An established procedure is the uvulopalatoplasty (UPPP) with tonsillectomy, in which redundant mucosa is resected from the posterior pillars and the remaining mucosa is tightened by suturing it to the anterior pillars. At least part of the uvula is also resected in most cases. The operation may employ conventional instruments or a laser technique (laser-assisted uvulopalatoplasty, LAUP). Some patients will also require adjuvant intranasal surgery (septoplasty, septorhinoplasty, turbinate reduction).

Differential diagnosis: The differential diagnosis should include various disorders that are associated with snoring or with the hypersomnia that is typical of obstructive sleep apnea (Table 5.3).
Tumors

Benign Tumors and Precancerous Lesions

The occurrence, incidence, clinical features, and treatment of these lesions are covered in 4.5, Benign Tumors and Precancerous Lesions of the Oral Cavity (p.90).

Malignant Tumors

The overwhelming majority of malignant tumors of the oropharynx are squamous cell carcinomas. Approximately 80% are located in the palatine tonsils or tongue base. Less common sites are the soft palate and posterior wall of the pharynx.

Etiology: In most patients, chronic nicotine and alcohol abuse have a major etiologic role in the development of oropharyngeal cancers.

Table 5.3 Differential diagnosis of sleep apnea

- Occasional or habitual nonobstructive snoring
- Upper airway resistance syndrome
- Narcolepsy
- Underlying heart disease with Cheyne–Stokes respiration (e.g., heart failure)
- Nocturnal bronchial asthma
- Periodic hypersomnia, hypersonia form of endogenous depression
- Insomnia
- Chronic alcohol and drug abuse

Source: adapted from Günther, see p. 430.

Symptoms:

Cancers at some sites in the oropharynx may remain clinically silent for some time.

Otherwise the symptoms depend on the location and extent of the tumor. Besides dysphagia and odynophagia, the symptoms may include blood-tinged saliva and a fetid breath odor. Advanced stages (Fig. 5.24a) often produce trismus, signifying that the tumor has invaded the surrounding musculature (pterygoid muscles).

Diagnosis:

Inspection: Tonsillar carcinomas may appear as exophytic lesions (Fig. 5.24a) or may show an ulcerating, infiltrating type of growth. Occasionally they are not grossly visible (microcarcinomas of the tonsils), and the first presenting symptom of the disease is cervical lymph-node metastasis.

CT and MRI (Fig. 5.24b–d) are useful for defining the extent of tumor growth and detecting the invasion of surrounding structures.

Treatment: The treatment of choice for most cases is surgical tumor removal. The resulting tissue defect may be closed primarily with local pedicled flaps or by using microvascular free tissue transfers, depending on the size and location of the defect (see Fig. 4.25, p. 95). A neck dissection (see p. 334) may be necessary on one or both sides, depending on the location and stage of the primary tumor (see Table 4.2, p. 93).

Postoperatively, radiation should usually be delivered to the tumor site and lymphatic pathways.

Alternatives for the treatment of advanced tumors (T3, T4) are primary radiotherapy or combined radiation and chemotherapy.
Visual inspection reveals an exophytic mass arising from the left tonsil. T1-weighted axial MR images before (b) and after contrast administration (c) show that the tumor (•) has deeply infiltrated the lingual muscles. The corresponding fat-suppression sequence more clearly delineates the tumor from its surroundings.
The diseases of the hypopharynx that have the greatest clinical importance are foreign bodies, hypopharyngeal diverticula, and especially malignant tumors, which frequently do not produce symptoms until they have reached an advanced stage. The ENT physician is not often confronted with diseases of the esophagus.

Injuries and Foreign Bodies

Caustic Ingestion

Etiopathogenesis: Caustic ingestion in children is almost always accidental, caused by drinking an alkaline or acidic liquid that has not been properly stored. Most such injuries in adults result from attempted suicide.

While acids cause a coagulation necrosis with the denaturation of proteins, alkalis cause a colliquative necrosis with liquefaction of the necrotic tissue. Strictures caused by scarring are common sequelae of this type of injury.

There is a long-term risk of cancer developing in esophageal strictures caused by caustic ingestion.

Symptoms:

Acute cases present with severe pain in the mouth and pharynx and possibly in the retrosternal and epigastic areas. Drooling is also present. Patients with an esophageal perforation may also present with subcutaneous emphysema in the neck or a pneumomediastinum (see Fig. 5.27, p. 127), and mediastinitis may supervene. Symptoms such as high fever and retrosternal or interscapular pain in these cases are accompanied by typical shock symptoms with an elevated pulse rate, a fall in blood pressure, cold sweats, and pallor. Generalized symptoms of intoxication such as renal and liver failure, electrolyte imbalance, and hemolysis generally do not appear until 1–2 days after the caustic injury.

In the long term, patients may develop an esophageal stricture with progressive dysphagia.

Diagnosis:

Acute evaluation begins with a mirror examination of the oral cavity, oropharynx, hypopharynx, and larynx. The mucosa initially appears erythematous and edematous and later may show epithelial defects and a whitish fibrin coating. It is also important to obtain radiographs of the chest and abdomen to exclude a perforation of the esophagus or stomach. If the diagnosis cannot be confirmed by imaging studies, esophagoscopy should be performed using careful technique.

The extent of the injuries to the oral and pharyngeal mucosa does not necessarily reflect the severity of corrosive damage to the esophagus, which may be very severe despite a normal appearance of the oral and pharyngeal mucosae.

Follow-up: Caustic injuries to the esophagus require long-term follow-up with imaging studies and periodic esophagoscopy.

Treatment:

Acute: The first priority is to treat the patient for shock. It is important to stabilize the airway, replace fluids, correct electrolyte imbalances, relieve pain, and provide sedation. Treatment should also include high doses of corticosteroids as well as antibiotics to prevent superinfection.

An esophageal stricture is treated with a dilator passed over a guide wire.
The therapeutic value of “neutralizing” agents (e.g., magnesium oxide for acid ingestion, citric or acetic acid for alkali ingestion) is questionable because the tissue damage occurs on immediate contact with the corrosive substance, and some time passes before the neutralizing agent can be administered.

**Long-term:** Some esophageal strictures can be treated by dilation. In cases with severe caustic injury, early dilation should be started just one week after the injury. The dilator should never be introduced blindly but should always be passed over a guide wire (Fig. 5.25). If a stricture cannot be expanded by dilation, the stenotic segment should be resected. It may then be necessary to perform a gastric pull-up or interpose a free segment of jejunum with microvascular anastomosis, depending on the length of the esophageal segment that has been removed.

**Foreign Bodies**

**Etiology and pathogenesis:** Foreign bodies typically become lodged in the hypopharynx or in the upper constriction of the esophagus (see Fig. 5.3, p.100). Most patients are small children who have swallowed coins, nuts, or toy parts, but in many cases they are older patients who have decreased sensation in the hard palate (e.g., due to a maxillary denture). Objects typically swallowed by adults are fish bones or larger bone fragments (Fig. 5.26a), pieces of meat (Fig. 5.26b), and denture parts.

**Symptoms:** Typical symptoms are a feeling of pressure, a “pricking” sensation, or pain in the hypopharynx or retrosternal area. Dysphagia may also be present, depending on the size and location of the foreign body.

**Diagnosis:** Inspection and palpation will disclose any cutaneous emphysema caused by perforation of the hypopharynx or esophagus (sharp object!). This is followed by indirect **mirror examination** of the hypopharynx. If this fails to locate the foreign body, diagnostic imaging should also be performed. The imaging procedure of choice depends on the nature of the foreign body suggested by the patient’s history. If a radiopaque foreign body is believed to be lodged in the hypopharynx or upper esophageal constriction, the soft tissues of the neck should be imaged with a lateral radiograph (Fig. 5.26a). Otherwise, an oral contrast examination (with a water-soluble medium) should be performed (Fig. 5.26b). An abdominal plain film can also show evidence of a foreign body in some cases (Fig. 5.26c).

Since there is always a danger of perforation, barium should never be used in oral contrast examinations (risk of foreign-body reaction or pneumonia).

**Treatment:**

The sun should never rise and set on a foreign body. Whenever an ingested foreign body is suspected, rigid esophagoscopy should be performed without delay to retrieve the foreign object. With a foreign body that has skewered and cannot be removed endoscopically, it may be necessary to expose the object through an external approach. This may require a transcervical incision or thoracotomy, depending on the location of the foreign body.

**Rupture of the Esophagus**

**Synonym:** Boerhaave syndrome

Boerhaave syndrome refers to a spontaneous rupture in the left posterolateral portion of the terminal esophagus just above the esophageal hiatus caused by forceful vomiting or retching. The condition is most common in patients with habitual vomiting and in alcoholics.

**Symptoms:** The classic symptoms occur immediately after the rupture and consist of very severe retrosternal or epigastric pain that may be accompanied by the features of an acute abdomen. Patients may also exhibit hematemesis, dyspnea, and progressive shock symptoms.

**Diagnosis:** First, an anteroposterior chest radiograph should be obtained in the standing or left lateral decubitus position. This will disclose a pneumomediastinum or possible air crescent below the diaphragm caused by air leakage from the ruptured esophagus. Neither of these signs is always evident in the chest radiograph, however, and so CT should be performed if there is lingering suspicion of a rupture (Fig. 5.27). Another option is oral contrast radiography of the esophagus, making certain to use only a water-soluble contrast medium (Gastrografin). If imaging procedures do not furnish a definitive diagnosis, the patient should undergo endoscopy.

The **treatment** of choice is immediate surgical intervention by thoracotomy with primary closure of the defect and pleural repair, leaving a drain in the pleura or mediastinum. The surgery must be performed under antibiotic coverage.
Diverticula

Two types are distinguished: pulsion diverticula, in which the mucosa herniates through a weak point in the muscular coat due to a rise of intraluminal pressure; and traction diverticula, which usually form at parabronchial sites due to scar traction following hilar lymphadenitis and involve all layers of the esophageal wall.

The treatment of esophageal diverticula is described in standard textbooks of surgery.

Hypopharyngeal Diverticulum

Synonym: Zenker diverticulum

Epidemiology: The hypopharyngeal (Zenker) diverticulum is the most common diverticulum of the esopha-
geal inlet. Most patients are middle-aged or older, with a 3:1 preponderance of males over females.

**Pathogenesis:** The herniation of esophageal mucosa (pulsion diverticulum) classically occurs at “weak points” in the posteroinferior hypopharyngeal wall located above the cricopharyngeal part of the constrictor pharyngis inferior muscle (see 5.1, p.98, and Fig. 5.2, p.104). Many patients present with a long history of reflux esophagitis.

In rare cases, a carcinoma may be the cause of a hypopharyngeal diverticulum.

**Symptoms:** Classic symptoms are dysphagia and the regurgitation of undigested food, especially in the morning and while lying down. Patients also complain of pronounced halitosis caused by food residues trapped in the diverticulum. Smaller diverticula are sometimes manifested only by a foreign-body sensation or may be completely asymptomatic.

**Diagnosis:** Mirror examination or indirect laryngoscopy (see 17.2, pp.346–349) will occasionally show the pooling of saliva in the piriform sinus, but only an imaging procedure can establish the diagnosis. An oral contrast examination of the esophagus is best for defining the diverticulum (Fig. 5.28). If reflux esophagitis is suspected (belching, heartburn, possible epigastric pain and dysphagia), the imaging study should be supplemented by ambulatory 24-hour pH-metry and esophageal manometry (5.6).

**Treatment:** The treatment of choice is surgery, using either an endoscopic or external approach.

**Endoscopic approach:** The endoscope is advanced through the mouth toward the esophageal introitus. The muscular septum formed by the cricopharyngeus is transected using either conventional technique or a CO₂ laser (Fig. 5.29), thereby reintegrating the diverticular pouch into the hypopharynx or esophagus. The endoscopic technique is particularly suitable for older patients with a high surgical risk, as the procedure is well tolerated and of relatively short duration.

**External approach:** An alternative is to resect the diverticulum through an external transcervical approach.
At surgery, the muscular septum is identified endoscopically and (in this case) divided with a CO₂ laser. A postoperative radiograph should always be taken to assess the integrity of the esophagus (Fig. 5.31).

**Tumors of the Hypopharynx**

### Benign Tumors

Benign tumors of the hypopharynx are considered a rarity. They may present clinically with dysphagia, regurgitation, or retrosternal pain. The diagnosis is established with an incisional biopsy taken endoscopically under general endotracheal anesthesia. Treatment consists of surgical removal, depending on the tumor size.

### Malignant Tumors

Histologically, almost all of these tumors are squamous cell carcinomas. As with oral and oropharyngeal carcinomas, there is an etiologic link to chronic alcohol and nicotine abuse.

**Symptoms:** Most malignant tumors of the hypopharynx are diagnosed at an advanced stage because earlier lesions do not produce symptoms. Initial complaints tend to be nonspecific, depending on tumor size and location (Table 5.4), and consist of dysphagia and a fetid breath odor. Later there may be...
pain radiating to the ear. Hoarseness and possible dyspnea signify tumor extension to the larynx. In many cases, cervical lymph-node metastasis is noted as the earliest sign of disease.

**Diagnosis:** Besides the mirror examination or indirect laryngoscopy (see 17.2, p. 346), the diagnostic workup should include endoscopic examination under general endotracheal anesthesia, as this is the best way to evaluate tumor extent. A biopsy can also be taken in the same sitting for histologic confirmation. Additionally, sectional imaging modalities can help to define the tumor size and check for involvement of adjacent structures while also evaluating the cervical lymph-node status (Fig. 5.32a).

**Treatment:** Treatment depends on tumor size but usually consists of local surgical excision with a concomitant neck dissection (see p. 334). Many malignant tumors of the hypopharynx have already spread to the larynx, making it necessary to perform a laryngectomy in the same sitting (see 17.7, pp. 368–377). The tissue defect is closed primarily whenever possible. This cannot be done with extensive hypopharyngeal resections due to the high risk of stricture formation, and larger defects should be reconstructed by means of a free jejunal transfer with microvascular anastomosis. Surgery should be followed by radiation to the tumor site and lymphatics.

*Alternative treatments* for advanced hypopharyngeal cancers are primary radiotherapy and combined radiation and chemotherapy.
6.1 Clinical Anatomy of the Salivary Glands

As an introduction, we will review the clinically relevant anatomy of the major and minor salivary glands. A knowledge of salivary gland anatomy is essential for understanding and diagnosing diseases of the salivary glands and providing appropriate treatment.

Classification

There are three pairs of major salivary glands (Fig. 6.1) and several hundred solitary, minor salivary glands distributed throughout the upper aerodigestive tract:
- **Parotid gland**: The largest salivary gland, it mainly produces a serous secretion.
- **Submandibular gland**: produces a mucouser secretion.
- **Sublingual gland**: also produces a mucouser secretion.
- **Minor salivary glands** (Fig. 6.2): These consist of labial glands in the mucosa of the lips, palatine glands in the mucosa of the palate, lingual glands in the tongue, and pharyngeal glands in the pharyngeal mucosa. They secrete a saliva that is predominantly mucous.

**Parotid Gland**

**Location**: The parotid gland descends into the retromandibular fossa between the vertical ramus of the mandible and the mastoid. Embedded in a subcutaneous pseudocapsule, it lies in contact with the sternocleidomastoid muscle and the posterior belly of the digastric muscle.

**Fibrous capsule**: The subcutaneous pseudocapsule, composed of dense fibrous tissue, is not a discrete tissue layer but blends with the skin and with the parotid gland itself. The capsule is poorly distensible and can cause severe pain when there is acute swelling of the gland. It is less dense inferiorly and medially, facilitating the spread of inflammations and tumors toward the pterygopalatine fossa and parapharyngeal space (“iceberg tumor,” see Fig. 6.5, p. 136).

**Parotid duct (Stensen duct)**: This excretory duct, approximately 6 cm long, leaves the parotid gland in its anterior superior third and passes forward over the masseter muscle (Fig. 6.1). It winds around the anterior border of the muscle and pierces the buccinator muscle and buccal mucosa. It opens opposite the second upper molar, forming an orifice with slightly raised edges.

After entering the parotid gland, the **facial nerve** branches into a plexus at the pes anserinus, subdividing the gland into a lateral and medial portion. This anatomical subdivision provides an important landmark during surgery of the parotid gland. The facial nerve is identified at its trunk and isolated. Surgical removal of the portion of the gland lateral to the pes anserinus is called a lateral parotidectomy. **Medial** to the pes are branches of the external carotid artery (superficial temporal artery, transverse facial artery) and venous vessels that drain into the internal jugular vein. **Lymphatic drainage** of the parotid gland is through several intraglandular and periglandular lymph nodes to the submandibular and deep jugular nodal chains.

**Submandibular Gland**

**Location**: The submandibular gland lies in the submandibular trigone between the two deep parts of the digastric muscle and the mandible. The gland makes a U-shaped bend around the posterior border of the mylohyoid muscle. The outer part of the gland extends past the trigone to a variable degree under cover of the outer cervical fascia. Because of this arrangement, the glandular compartment connects the sublingual and posterior portions of the oral floor, cre...
6.1 Embryology, malformations, and anomalies

Embryology

The major salivary glands arise from ectodermal tissues in the foregut between the fourth and eighty weeks of embryonic development. The surrounding mesenchyma segregates the glandular tissue and may include lymph-node primordia. The excretory ducts become patent by the 22nd week of development.

Malformations

Aplasia, hypoplasia, duct atresia: Aplasia of all the salivary glands is extremely rare. Malformations affecting individual salivary glands are also rare. Duct atresia most commonly affects the submandibular glands, and cysts may form.

Dystopias, accessory and aberrant salivary glands: Dystopia refers to the abnormal location of an otherwise normally developed salivary gland, such as a parotid gland located anterior to the masseter muscle. Accessory salivary glands are appendages of the major salivary glands that communicate with the duct system and are fully functional. They are most commonly found appended to the parotid gland (see Fig. 6.1). Aberrant salivary glands are heterotopic salivary gland primordia that do not have a duct system and are non-functional. They occur most frequently in the lateral neck or gingiva and rarely in the middle ear. Salivary gland tumors may develop from these aberrant glands.

Dysgenetic cysts and ectasias: These congenital malformations of the excretory ducts require differentiation from similar, acquired abnormalities. They can predispose to recurrent inflammations (see 6.2, Salivary gland cysts, p.143).

ating a potential route for the spread of infection (abscess or cellulitis of the oral floor).

Submandibular duct (Wharton duct): The excretory duct, approximately 5 cm long, of the submandibular gland passes with the sublingual process of the gland to the sublingual plane of the oral floor and runs forward beneath the mucosa. It crosses over the lingual nerve (Fig. 6.3) and opens at the sublingual caruncle.

Nerves and vessels: The lingual nerve is not only closely apposed to the excretory duct but also forms a genu that runs just above the gland body, where it distributes branches to the submandibular ganglion. The hypoglossal nerve runs inferomedially to the gland. The facial artery and vein loop around the posterior part of the gland.

The lymphatic drainage of the submandibular gland is to lymph nodes in the lateral and posteriorinferior portion of the gland, which also receive drainage from the face and oral cavity. Because of this, it is usually necessary to remove the submandibular gland when a neck dissection is performed (see p. 334).

Sublingual Gland

The sublingual gland lies in the anterior, submucous part of the oral floor on the mylohyoid muscle, lateral to the submandibular duct and distributed along the medial surface of the mandible (Fig. 6.3). Excretory ducts may drain into the submandibular duct or may open directly into the oral mucosa as small ducts.
6.2 Functional Morphology and Physiology of the Salivary Glands

To understand disorders of salivary gland function, it is also important to know the common histologic structure of the glands, the physiologic functions of the saliva, and the clinical terms for secretory dysfunction.

**Histologic Structure**

All salivary glands are based on a common structural principle: glandular acini connected to a system of salivary ducts (Fig. 6.4). The acini and the duct system are embedded in a glandular mesenchyma that contains connective tissue, blood and lymphatic vessels, lymphatic tissue, and nerve fibers.

**Glandular Acini**

The acini produce the primary saliva, which contains enzymes (including amylase) and salivary mucins. Several histologic types are distinguished based on the relative amounts of enzymes and mucins that are formed:

- Serous glands, which mainly produce enzymes (e.g., the parotid gland)
- Mucous glands, which mainly produce mucin (e.g., the palatine glands)
- Mixed glands (e.g., the submandibular and sublingual glands)

The acini include myoepithelial cells, which form a weblike structure that surrounds the acinus and can extrude its contents through a contractile action.

**Salivary Duct System**

The salivary duct system is not a passive transport system but actively modifies the contents and consistency of the primary saliva. The short intercalated ducts secrete mucins and regulate the electrolyte concentration. Next come the striated ducts, which can quickly and actively secrete fluid, followed by the interlobular duct system, which mainly transports the saliva while only slightly altering its properties.

**Composition of the Saliva**

The salivary glands constantly produce a certain amount of saliva. In the absence of external stimuli, this production, called the *resting secretion*, probably relates to the basic activity of the salivary nucleus. External and internal factors, particularly eating and the associated chewing movements, smells, etc., can significantly increase the rate of salivary secretion. In the parotid gland, this *stimulated secretion* is approximately 4–5 times greater than the resting secretion (Table 6.1).

<table>
<thead>
<tr>
<th>Table 6.1 Resting secretion versus stimulated secretion</th>
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<tbody>
<tr>
<td><strong>Resting secretion</strong></td>
</tr>
<tr>
<td>Origin</td>
</tr>
<tr>
<td>Consistency</td>
</tr>
<tr>
<td>Contents</td>
</tr>
</tbody>
</table>
The total daily volume of saliva secreted by all the salivary glands is from 500 to 1000 mL, subject to influence by numerous factors such as climate, fluid intake, nutrition, age, and gender (Table 6.2). The production of saliva is controlled mainly by parasympathetic-cholinergic stimuli. But sympathetic stimuli also exercise a control function via α- and β-adrenergic receptors, especially in the regulation of α-amylase.

**Physiologic Functions of Saliva**

**Nutrition and Digestion**

The saliva emulsifies and dissolves food constituents, aiding in the perception of taste. The glycoproteins in the saliva lubricate the bolus to facilitate swallowing. The actual digestive enzyme is α-amylase, a starch-splitting enzyme that is produced mainly by the parotid gland.

**Protective Functions**

The quantity and composition of saliva have a major influence on the microbiological and inorganic milieu of the oral cavity. Mechanical cleansing (irrigation) plays a role, as does the secretion of enzymes (lysozyme, muramidases, peroxidases) and immunoglobulins (mainly IgA).

The production of saliva is also important in the prevention of gum disease and dental caries.

**Excretion**

Both endogenous and exogenous substances can be excreted in the saliva. Of clinical importance is the excretion of certain ions (iodine, fluoride) and of viruses that can be transmitted via the saliva—poliomyelitis, hepatitis B, Epstein–Barr, cytomegalovirus, coxsackievirus, rabies, human immunodeficiency virus (HIV). The excretion of genetically determined glycoproteins, whose antigenic properties are similar to the ABO system but independent of it, can be of forensic importance.

**Secretory Disorders**

Disorders of salivary gland secretion, transport, and consumption can lead to changes in the quality and quantity of the saliva. The general term for these disorders is *dyschylia*. Several terms are applied to increased salivary flow and are not always clearly differentiated from one another (Table 6.3). In *sialorrhea*, the quantity of the saliva is not necessarily increased. *Sialorrhea* is particularly common in children with cerebral palsy and may become a significant nursing problem.

**Table 6.2 Factors that influence saliva production**

<table>
<thead>
<tr>
<th>Hypersalivation</th>
<th>Hyposalivation</th>
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<tbody>
<tr>
<td>Stimulation</td>
<td>Depression</td>
</tr>
<tr>
<td>Foods, acids, poisons (mercury, arsenic, lead), nausea, pregnancy</td>
<td>Inflammation of the glands, dehydration, marasmus, radiation exposure</td>
</tr>
<tr>
<td>Parasympathomimetic agents (e.g., pilocarpine, muscarine, nicotine), sympathomimetic agents (e.g., isoproterenol), iodine, bromide, fluoride, curare, theophylline, caffeine</td>
<td>Anticholinergic agents (e.g., atropine, scopolamine), alpha-blockers (e.g., phentolamine), beta blockers (e.g., propranolol), antihistamines, antihypertensive agents (e.g., clonidine, reserpine), psychoactive drugs (e.g., antidepressants), topical anesthetics</td>
</tr>
</tbody>
</table>

**Table 6.3 Terms for disorders of salivary secretion**

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Symptoms</th>
</tr>
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<tbody>
<tr>
<td>Dyschylia</td>
<td>General disturbances of salivary secretion or production</td>
</tr>
<tr>
<td>Hypersalivation</td>
<td>Increased flow of saliva</td>
</tr>
<tr>
<td>Synonym: <em>ptyalism</em></td>
<td>Causes: see Table 6.2</td>
</tr>
<tr>
<td>Sialorrhea</td>
<td>Excessive flow of saliva from the mouth (drooling, slobbering)</td>
</tr>
<tr>
<td></td>
<td>Caused by swallowing difficulties due to:</td>
</tr>
<tr>
<td></td>
<td>• Neurologic diseases (Parkinson’s disease, bulbar paralysis, myasthenia gravis, cerebral palsy)</td>
</tr>
<tr>
<td></td>
<td>• Mechanical obstruction (pharyngeal tumor, esophageal obstruction)</td>
</tr>
<tr>
<td>Hyposalivation</td>
<td>Decreased flow of saliva</td>
</tr>
<tr>
<td>Synonym: <em>sialopenia</em></td>
<td>Causes: see Table 6.2</td>
</tr>
<tr>
<td>Asalia</td>
<td>Absence of salivary secretion</td>
</tr>
<tr>
<td>Xerostomia</td>
<td>Dryness of the oral mucosa</td>
</tr>
<tr>
<td>Sicca syndrome</td>
<td>Dryness of the oral mucosa and other mucous membranes (conjunctiva, genital mucosae)</td>
</tr>
</tbody>
</table>

The condition resulting from pronounced hyposalivation may be termed *xerostomia* (dry mouth) or *sicca syndrome* (additionally involving the conjunctiva and other mucous membranes). It is frequently associated with very troublesome complaints.
6.3 Clinical Examination, Imaging Studies, and Biopsy of the Salivary Glands

In many cases, a salivary gland disease can be classified as inflammatory or neoplastic based on a detailed clinical examination. The first-line modality for further investigation is ultrasonography; computed tomography (CT) and magnetic resonance imaging (MRI) are available as second-line studies. The histologic diagnosis is particularly important, as it provides the basis for treatment.

Clinical Examination

History

Particular attention should be given to:

- Systemic diseases, especially metabolic disorders (e.g., diabetes mellitus), which can have a systemic effect on the salivary glands
- Medications (antihypertensive drugs, psychotherapeutic drugs; see Table 6.2, p. 135)
- Prior illnesses, surgical procedures, or therapies (radiotherapy) involving the glands and oral cavity
- Hypersalivation and hyposalivation, sialorrhea, or sicca syndrome (see Table 6.3, p. 135)

Inspection

The glands are inspected externally (periauricular and submandibular region) along with the oral cavity and tonsillar region. Facial nerve motor function should also be tested.

External: Normally, visual inspection reveals only the flat contour of the submandibular gland in the submandibular trigone if the overlying skin is thin. A normal parotid gland is not visible. Masseter hyperplasia can mimic enlargement of the parotid gland but is easily differentiated by having the patient squeeze the jaws together. This activates the masseter muscle, making it easily accessible to inspection and palpation.

Oral cavity: The orifices of the excretory ducts (Wharton and Stensen ducts) are evaluated for redness and swelling. The flow of saliva, either spontaneous or in response to glandular massage, is an important parameter for differentiating between obstruction, inflammation, and normal findings (saliva clear or absent, flocculent, purulent, blood-tinged).

Tonsillar region (Fig. 6.5): The parapharyngeal or tonsillar region may appear prominent due to swelling of the deep portions of the parotid gland.

With changes in the parotid gland, especially neoplastic changes, facial nerve function should always be tested and documented in a side-to-side comparison.

A tumor involving the deep portions of the parotid gland may produce a bulge in the tonsillar region (“iceberg tumor”).

Bimanual palpation of the submandibular gland.
Palpation

A normal parotid gland is barely palpable. The submandibular glands, a swelling of the sublingual glands, and the excretory ducts are palpated bimanually (Fig. 6.6), noting the size of any abnormalities in centimeters and assessing their consistency, surface contours, tenderness, and mobility relative to the skin and underlying tissues. A normal sublingual gland cannot be palpated. The intraglandular, periglandular, and cervical lymph nodes should also be examined.

Imaging Studies

Ultrasound examination: Ultrasound has proved to be a well-tolerated and rewarding study for investigating the salivary glands. It can differentiate among normal glandular parenchyma, inflammatory processes (with or without liquefaction), tumors, lymph nodes, and calculi (Fig. 6.7). Generally the ultrasound examination is the first imaging procedure used. It can be combined with fine-needle aspiration biopsy.

Radiographs: Plain radiographs of the oral floor, submandibular gland, and parotid gland are seldom rewarding because of superimposed structures. To be visible on radiographs, calculi must have a sufficient calcium content and measure at least 2–3 mm in size. For this reason, plain radiographs are of diagnostic value only in patients with a suspected salivary stone in the Wharton duct (oral floor view).

Sialography: Radiographic contrast examination of the excretory ducts after catheterization gives the most detailed view of the duct systems of the parotid and submandibular glands. Sialography is rarely performed today, however, due to potential complications (infection, abscess formation, extravasation) and the availability of ultrasound, sialoendoscopy, and MRI. Its possible indications include the detection of small stones in the excretory ducts, anomalies of the excretory ducts, sialadenosis, and chronic inflammation.

Sialography is contraindicated in the presence of acute inflammation.
CT and MRI: Most lesions of the salivary glands that cannot be adequately diagnosed from the history, palpation, and ultrasound findings must be investigated by CT or MRI. This particularly applies to tumors and masses that transcend the gland boundaries or involve the deep portions of the parotid gland (Fig. 6.8). MRI is superior to CT in the diagnosis of salivary gland tumors (Fig. 6.9a–c).

Sialoendoscopy

The major excretory ducts (Stensen and Wharton ducts) can also be inspected with special endoscopes, which are introduced under local anesthesia after dilation of the papillae. The duct is irrigated during endoscopy, which can demonstrate mucous plugs, calculi, and stenoses. Interventional procedures can also be performed such as dilation or the removal of a stone with a loop or by laser lithotripsy.

Biopsy

Fine-needle aspiration biopsy (FNAB) has established itself as a low-risk, well-tolerated procedure in the preoperative investigation of salivary gland swelling. It can also furnish material for bacteriologic analysis. More deeply situated lesions can be aspirated under ultrasound guidance. The main complication of FNAB is secondary infection of the puncture site, which is why the procedure requires meticulous aseptic technique. Core biopsy, once widely practiced, should no longer be used today.

Facial nerve injury, bleeding, and tumor cell dissemination along the needle track are potential problems in core biopsies but not in FNAB.

Incisional biopsy and intraoperative frozen section: In rare cases, an incisional biopsy of the salivary glands may be warranted for the investigation of a chronic inflammatory process. An incisional biopsy is relatively safe for ulcerative lesions and in patients with facial nerve palsy. Otherwise, an incisional biopsy should be avoided, especially in the parotid gland, due to the ever-present risk of facial nerve injury.

Incisional biopsy is contraindicated for benign salivary gland tumors.

If the tumor is most likely benign (p. 140), the tumor should be completely removed along with its capsule. This involves surgical removal of the gland combined with an intraoperative frozen section. With generalized salivary gland disease (e.g., myoepithelial sialadenitis), the biopsy of a minor salivary gland in the lower lip may yield a histologic diagnosis. This procedure is safe and easy to perform under local anesthesia, but its sensitivity is low.
Magnetic resonance images of a lymphangioma of the left parotid gland, with correlative drawings. Axial T1-weighted (a) and T2-weighted (b) postcontrast images at the level of the external auditory canal, and a coronal image (c) at the level of the sphenoid sinus.
6.4 Overview: Diagnosis and Management of Salivary Gland Swelling

Diseases of the salivary glands are often manifested by unilateral or bilateral swelling of the glands. This swelling leads to differential diagnostic considerations in which the history, clinical examination, imaging studies, and biopsy have an important role (see 6.3, pp. 136–139). This shows how the various diagnostic options can be effectively coordinated in practice (see also 6.7, pp. 148–151).

The swelling of one or more salivary glands may be a result of duct obstruction, inflammation, or neoplasia. A detailed history and physical examination will usually furnish a differential diagnosis as a basis for further testing. The clinical suspicion of a malignant tumor is particularly important in this regard. The clinical steps involved in making a differential diagnosis are outlined in Fig. 6.10. Standard laboratory tests are also helpful in distinguishing between inflammation and a tumor. An elevated white blood cell count, erythrocyte sedimentation rate (ESR) or C-reactive protein (CRP) is suggestive of inflammation. Tumors of the salivary glands generally produce no changes in routine blood tests. A somewhat rare exception to this rule is the situation where an obstructive tumor has caused salivary stasis, leading to an infection.

Generally, an ultrasound examination is the next diagnostic procedure to follow the clinical examination. It can demonstrate changes in the duct system and more accurately characterize the swelling, differentiating between a cyst and solid lesion, for example. This information will usually yield a diagnosis or indicate the need for additional, more specific studies as shown in Fig. 6.11.

The definitive diagnosis of a swollen salivary gland is occasionally made at the time of operation, particularly in the case of a tumor. Tumors are manifested clinically by a more or less painless, unilateral swelling with a palpable nodule. It is important to look for signs that are helpful in distinguishing benign and malignant tumors.

Signs suggesting a benign tumor:
- Slow growth (months to years)
- Painless, soft or tense nodule that is freely movable
- No signs of tumor infiltrating the surrounding tissue
- No additional symptoms

Signs suggesting a malignant tumor:
- Rapid growth (weeks to months)
- Painful, fixed nodule
- Evidence of tumor infiltrating muscle, skin, or nerves (facial nerve palsy, Fig. 6.12)
- Lymph-node enlargement

Facial nerve palsy associated with a tumor in the parotid gland almost always signifies a malignant tumor.

The location of a salivary gland tumor can also furnish clues to the differential diagnosis of the tumor.

Eighty percent of salivary gland tumors occur in the parotid gland, approximately 10% in the submandibular gland, and 10% in other salivary glands.

A localized swelling of the parotid gland very often signifies a tumor. This is less often the case with the submandibular gland. Only about 20% of tumors in the parotid gland are malignant, compared with approximately half of tumors in the other salivary glands.
Fig. Flowchart for the investigation of salivary gland diseases

1. History, inspection, palpation
   - Calculus, ductectasia
     - MRI, endoscopy
     - CT or MRI
     - FNAB
     - Biopsy of a minor salivary gland
     - MRI
     - Incisional biopsy
2. Ultrasound examination
   - Diffuse swelling
   - Circumscribed swelling
     - Suspected malignancy or large tumor
     - FNAB and MRI (if necessary, CT)
     - Surgery with frozen-tissue histology
     - Suspected benign tumor
     - FNAB if necessary

Fig. Facial nerve palsy due to a salivary gland tumor

Mucoepidermoid carcinoma of the right parotid gland with facial nerve palsy, signifying invasion of the nerve by the tumor.

Rule of thumb for salivary gland tumors: the smaller the salivary gland, the greater the likelihood that the tumor is malignant.

FNAB will yield an accurate cytologic diagnosis in approximately 80% of tumors. The positive detection of malignant cells has a high correlation with the actual presence of a malignant tumor, and appropriate tests should be initiated at once. It does not justify a radical procedure such as facial nerve resection, however. At the same time, the absence of malignant cells on FNAB does not confidently exclude a malignancy.
6.5 Noninflammatory Diseases and Injuries to the Salivary Glands

This deals with salivary gland diseases other than primary inflammations and tumors. The most important are sialolithiasis, sialadenosis, and injuries.

Sialolithiasis

Synonym: salivary stone disease

**Definition:** Stone formation in the excretory duct system of a salivary gland.

**Epidemiology:** Adults in the third and fourth decades are most commonly affected, with males predominating by a ratio of 2:1.

**Location:** From 60% to 70% of salivary stones are located in the main duct. Generalized lithiasis (urinary stones, gallstones, and salivary stone) is present in approximately 6–10% of all cases.

From 70% to 80% of salivary stones occur in the submandibular gland, and approximately 20% in the parotid gland. A smaller percentage occur in the minor salivary glands or sublingual gland.

**Etiopathogenesis:** Salivary stones result from the secondary calcification of “plugs” that form from enriched organic salivary contents (mucins). A microcalcification can lead to increased salivary stasis and to the precipitation of inorganic compounds.

**Symptoms:** Eating and other gustatory stimuli incite a swelling of the affected gland, often accompanied by severe pain (“salivary stone colic”). There may also be a stasis-induced “salivary tumor.” The stasis can lead to infection of the excretory duct and gland, which may present as a primary or secondary symptom (see Acute Sialadenitis, p. 145).

**Diagnosis:** Stones are often palpable in the duct system of the submandibular gland. Ultrasound reveals dilation of the duct system with typical acoustic shadowing. Approximately 70% of the stones are radiopaque, particularly those occurring in the submandibular gland. An oral floor radiograph can demonstrate stones in the distal part of the Wharton duct, but this is necessary only in the rare cases with equivocal ultrasound findings. Calculi in the parotid gland are particularly difficult to define on radiographs because they are poorly calcified and are obscured by superimposed structures.

**Differential diagnosis:** External obstruction of the excretory duct, as by a denture or tumor, is the most frequent differential diagnosis. Phleboliths are rare, and calcified lymph-node tuberculomas are very rare.

**Complications:** Infection and abscess, particularly of the oral floor.

**Treatment:** Treatment is generally surgical.

- With distal stones, it is sufficient to incise the excretory duct, extract the stone, and suture the duct epithelium to the mucosa (marsupialization).
- Intraglandular stones of the submandibular gland and a chronically damaged gland should be excised.
- Salivary stones located in the duct or near the gland can be removed by endoscopic fragmentation (using a laser beam or ultrasound shock waves) or extracorporeal lithotripsy (ultrasound shock waves).

Stone in the parotid gland are more difficult to treat. Conservative therapy is tried first (increased fluid intake, anti-inflammatory agents). A parotidectomy is rarely indicated.

Sialadenosis

**Definition:** Sialadenosis refers to a noninflammatory, symmetrical swelling of the major salivary glands caused by a systemic, frequently unknown cause. The parotid gland is most commonly affected.

**Pathogenesis:** Sialadenosis can occur in association with:

- Chronic alcoholism
- Vitamin deficiencies
- Diabetes mellitus
- Protein deficiency
- Anorexia nervosa and other eating disorders

Sialadenosis is a secretory disorder characterized by enlarged acinar cells. The most likely cause is disordered autonomic innervation of the affected salivary glands.
6.2 Salivary gland cysts

Cysts of the salivary glands require differentiation from tumors and chronic inflammations. Various forms may be encountered:

**Dysgenetic cysts:** These are primary cysts resulting from a developmental abnormality and must be distinguished from secondary, acquired cysts. Dysgenetic cysts occur most commonly in the sublingual gland and its excretory ducts. A cyst at this location is called a ranula (“little frog,” after its resemblance to the expanded vocal air sac of a frog, Fig.). A ranula of sufficient size can restrict tongue mobility and cause difficulties with speech and swallowing. It may also become infected. Treatment consists of surgical removal. Dysgenetic cysts of the parotid gland may become inflamed, particularly in children.

**Salivary duct cysts** occur predominantly in the parotid gland. They may be complicated by infection.

**Mucoceles and retention cysts of the minor salivary glands:** Injuries to minor salivary glands may allow saliva to escape into the tissue, forming a pseudocyst. Duct obstructions give rise to true cysts (“retention cysts”) with an epithelial lining. Both forms occur predominantly in the mucosa of the lower lip. The differential diagnosis should also include a possible tumor of the minor salivary glands, especially when the mass is located on the palate.

**Lymphoepithelial cysts** probably originate from lymph follicles and therefore occur mainly in the parotid gland. Histologic examination reveals lymphatic tissue in the cyst wall. This type of cyst is particularly common in human immunodeficiency virus (HIV) infections; they tend to be bilateral and occur predominantly in younger patients.

**Symptoms:** The disease causes a painless, usually symmetrical swelling that is unrelated to eating.

**Diagnosis:** A bilateral, symmetrical, painless swelling of the salivary glands is typically noted on clinical examination. CT, MRI, or fine-needle aspiration biopsy may be indicated in cases with equivocal clinical findings. The diagnosis can be histologically confirmed by glandular biopsy, but this is rarely necessary.

**Differential diagnosis:** All forms of chronic sialadenitis must be excluded. Less frequent possibilities are masseter hyperplasia and obesity with fatty hypertrophy of the gland.

**Treatment:** Treatment is directed toward the underlying cause. No specific therapy for sialadenosis is required.

**Injuries**

**Penetrating or Blunt Trauma**

Direct sharp or blunt injuries most commonly affect the parotid gland, which occupies a less protected location than the submandibular gland. It is important to distinguish injuries to the glandular parenchyma alone from injuries involving the excretory duct system or facial nerve.

Every open salivary gland injury should be surgically explored.

**Bleeding:** Bleeding in the parotid gland is not life-threatening and can usually be managed by primary compression. Coagulation, clipping, or ligation should be avoided in an obscured field (risk of facial nerve injury).

**Duct injuries:** A duct injury can result from trauma to the anterior third of the gland. Whenever possible, a microsurgical end-to-end anastomosis should be performed over a fine plastic catheter. Another option is to suture the duct stump to the mucosa, creating a neo-ostium.

**Facial nerve injuries:** see 14.2, p. 295. Immediate or early treatment is advised.

**Pneumoparotid**

The retrograde entry of air into the Stensen duct is considered a special case of parotid trauma. It results from a high positive pressure in the oral cavity, which may occur during forcible mask ventilation or when blowing a musical instrument, blowing glass, or inflating a balloon. It is marked by transient pain in the region of the parotid gland. Cutaneous emphysema is rarely detectable by palpation. Specific treatment is unnecessary, and the air should be quickly eliminated by reabsorption or leakage from the duct orifice.
6.6 Inflammatory Diseases of the Salivary Glands (Sialadenitis)

The inflammation of a salivary gland usually leads to diffuse swelling of the entire gland. Table 6.4 reviews the various forms of sialadenitis, which may be acute or chronic. They may be caused by viral infections, bacterial infections, autoimmune diseases, or ionizing radiation exposure.

Acute Sialadenitis

Acute Viral Sialadenitis, Mumps

Various organisms have been identified as the cause of acute viral inflammations of the salivary glands. Rare causes are cytomegalovirus (6.3), coxsackievirus, influenza virus, and the human immunodeficiency virus. The most common viral pathogen is the mumps virus. It is the causative organism of mumps (see below), which occurs predominantly but not exclusively in children. Synonyms: epidemic parotitis, infectious parotitis.

The causative organism is the mumps virus, from the family of paramyxoviruses. The virus is shed in the saliva, and the infection is spread by droplet transmission. The major salivary glands are infected by the hematogenous route. The incubation period is 18 (±10) days. Fifty percent of cases run a silent course, and the infection confers lifelong immunity.

Symptoms: Cases typically present with diffuse, painful swelling of the parotid glands with a doughy edema (“hamster cheeks”). Often one parotid gland is affected initially, followed several days later by swelling of the cervical lymph nodes, the opposite parotid gland, and the submandibular glands. The duct orifices are reddened and slightly swollen, and the secretions are nonpurulent. Usually only a mild fever is present, and approximately 30% of patients are afebrile. The infection should resolve in 1–2 weeks.

Diagnosis: The diagnosis is based on the clinical presentation. Serologic testing is necessary only in doubtful cases. A four-fold rise of antibody titers 2–3 weeks after the onset of the disease establishes the diagnosis.

Differential diagnosis: Differentiation is mainly required from cervical lymphadenitis and acute suppurative parotitis. Less frequent conditions to be considered are chronic recurrent parotitis, a dentogenic infection or abscess, sialolithiasis, or a tumor.

Complications: An abnormal cerebrospinal fluid examination accompanied by serous meningitis is relatively common. Serious but less frequent complications are meningencephalitis with permanent cranial nerve deficits, orchitis, and labyrinthitis. Deafness may also occur and is unilateral in most cases. The pancreas and ovaries may also be involved.

6.3 Cytomegalovirus sialadenitis

The second most common viral sialadenitis is caused by cytomegalovirus (salivary gland inclusion disease). The sialotropic virus usually affects the salivary gland with no inflammatory signs and is shed in the saliva. Often acquired perinatally, the infection may run a silent course or may produce various symptoms separate from the salivary glands themselves, such as sensorineural hearing loss. In later stages of life the infection most commonly occurs in immunocompromised patients and causes systemic symptoms that resemble mononucleosis.

Table 6.4 Types of sialadenitis

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute sialadenitis</td>
<td>Viral:</td>
</tr>
<tr>
<td></td>
<td>• Mumps</td>
</tr>
<tr>
<td></td>
<td>• Cytomegalovirus (6.3)</td>
</tr>
<tr>
<td></td>
<td>• Coxsackievirus, echovirus, parainfluenza viruses, influenza</td>
</tr>
<tr>
<td></td>
<td>Bacterial:</td>
</tr>
<tr>
<td></td>
<td>• Acute suppurative parotitis</td>
</tr>
<tr>
<td></td>
<td>• Obstructive (electrolyte) sialadenitis</td>
</tr>
<tr>
<td>Chronic sialadenitis</td>
<td>Chronic recurrent parotitis</td>
</tr>
<tr>
<td></td>
<td>Chronic recurrent sialadenitis of the submandibular gland,</td>
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<tr>
<td></td>
<td>sclerosing sialadenitis (Kütner tumor)</td>
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<tr>
<td>Radiation sialadenitis</td>
<td>Immune sialadenitis:</td>
</tr>
<tr>
<td></td>
<td>• Myoepithelial (Sjögren syndrome)</td>
</tr>
<tr>
<td></td>
<td>• Epithelioid cell (Heerfordt syndrome; 6.4)</td>
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<tr>
<td></td>
<td>Infectious granulomatous sialadenitis (6.4):</td>
</tr>
<tr>
<td></td>
<td>• Tuberculosis</td>
</tr>
<tr>
<td></td>
<td>• Actinomycosis</td>
</tr>
<tr>
<td></td>
<td>• Syphilis</td>
</tr>
</tbody>
</table>

Prophylaxis: The best preventive measure is a mumps vaccination program (96% protection rate with proper use).
Acute Bacterial Sialadenitis

Synonym: acute suppurative sialadenitis

Etiology and pathogenesis: This suppurative infection most commonly affects the parotid gland in debilitated, dehydrated patients. When the submandibular gland is affected, the cause is usually obstructive (sialolithiasis, poorly fitting denture) or dentogenic in nature.

The main causative organism is *Staphylococcus aureus*. *Streptococci*, *Haemophilus*, and other organisms also occur. The typical pattern is an ascending bacterial infection in a patient with decreased salivary flow. Usually there is a general predisposing condition for bacterial sialadenitis such as diabetes mellitus, weakened host defenses, or poor oral and dental hygiene.

Symptoms: The patient presents with a painful, diffuse swelling of the affected gland. The skin over the gland may be reddened, and the gland may become fluctuant due to tissue liquefaction (Fig. 6.13). The excretory duct orifices are red and swollen. A turbid fluid or pus can be expressed from the gland orifice or may drain spontaneously. Trismus may be present.

Diagnosis: The diagnosis is made from the typical palpable findings and suppurative discharge in patients with a corresponding prior illness. The discharge should be tested bacteriologically to determine antibiotic sensitivity.

Differential diagnosis: A dentogenic infection can produce similar findings. A less frequent cause is furuncular otitis with periauricular abscess or lymphadenitis (especially with sialadenitis of the submandibular gland).

Treatment: Generally the disease responds well to medical management with antibiotics, analgesics (nonsteroidal anti-inflammatory agents), hydration, salivary stimulation, and good oral hygiene. If an abscess develops, the parotid gland should be incised parallel to the branches of the facial nerve.

Chronic Sialadenitis

(See also 6.4, p. 146)

Chronic Recurrent Parotitis

The pathogenesis of recurrent bacterial infections of the parotid gland, which are common in childhood but also occur in adults, is uncertain. Congenital ductasia is believed to be a predisposing factor.

Symptoms: Usually there is a unilateral or alternating (rarely bilateral) swelling of the parotid gland, which may be very painful. The saliva is milky, granular, or purulent. Trismus is frequently present. The attacks recur at varying intervals. Between attacks, the patient has no subjective symptoms but the parotid gland may be indurated.

In children, the symptoms usually resolve during puberty. Cases in adults may take a very protracted course in which obliterator scarring of the parenchyma causes saliva production to dwindle and finally cease, with an associated resolution of symptoms.

Diagnosis: The diagnosis is made from the history and clinical course. Generally a normal sonogram is obtained between attacks. Though rarely indicated, sialography can demonstrate a "leafy tree" pattern (excretory ducts with ectasia of the acini and terminal duct segments).

Differential diagnosis: Differentiation from the less common immune sialadenitis can be difficult in adults and may necessitate an incisional biopsy. Immune sialadenitis occurs predominantly in women.

Complications: Abscess formation.

Treatment: Exacerbations are treated the same as acute bacterial parotitis. A conservative approach is definitely indicated in children. Adult patients may require a parotidectomy, which is difficult in these cases and carries a significant risk of facial nerve injury.
### 6.4 Other forms of chronic sialadenitis

**Epithelioid cell sialadenitis and Heerfordt syndrome:** Sarcoidosis of the salivary glands is usually characterized by unilateral or bilateral involvement of the parotid gland, which shows a moderately firm, constant swelling. Intraglandular lymph nodes may be affected in addition to the glandular parenchyma. Involvement of the minor salivary glands may occur. Pain is relatively mild, and little or no sialolopenia is present. The diagnosis is established by biopsy. Differentiation from tuberculosis is required. The gland is treated with corticosteroids.

Simultaneous involvement of the eyes (mainly the uvea) and salivary glands is known as Heerfordt syndrome (subchronic uveoparotid fever). It is often associated with cranial nerve deficits, especially facial nerve palsy.

**Tuberculosis:** Tuberculosis of the salivary glands is rare. The intraglandular lymph nodes are predominantly affected, and less commonly the glandular parenchyma. The diagnosis is established by bacteriologic testing and/or histologic examination. Treatment relies on tuberculostatic drugs.

Other chronic forms: Actinomycosis with a hard, painless swelling and a typical violaceous skin discoloration should be included in the differential diagnosis. It may develop in proximity to the parotid or submandibular gland and may involve the gland secondarily. It is generally rare.

**Syphilis** of the salivary gland is very rare but should be excluded when a granulomatous inflammation is present.

**HIV infection:** Symmetrical enlargement of the salivary glands is relatively common in patients infected with human immunodeficiency virus (HIV salivary gland disease). It is particularly common to find bilateral lymphoepithelial cysts of the parotid glands, and patients with acquired immune deficiency syndrome (AIDS) may exhibit a Sjögren-like syndrome with marked xerostomia.

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### Chronic Sclerosing Sialadenitis

**Synonyms:** chronic recurrent sialadenitis of the submandibular gland, Küttner tumor

**Definition:** This is the most common form of chronic inflammatory sialadenitis, usually occurring in association with obstruction and sialolithiasis. It may culminate in a permanent, tumor-like swelling of the affected gland.

**Pathogenesis:** An altered composition of the saliva, usually combined with obstruction of the excretory ducts, is the main pathogenic mechanism for the inflammation.

**Symptoms:** Patients present with a firm swelling of the submandibular gland, which undergoes an acutely painful enlargement that is related to eating. The end stage, called a Küttner tumor, is a firm, constant, essentially nontender enlargement of the gland that is difficult to distinguish from a neoplasm by palpation.

**Diagnosis:** The diagnosis is made by demonstrating the duct obstruction with ultrasound. MRI is necessary only in complicated cases. Fine-needle aspiration biopsy reveals inflammatory changes. The gland can be extirpated as an "excisional biopsy."

**Differential diagnosis:** Differentiation is required from other causes of excretory duct obstruction such as extrinsic pressure due to tumors, cysts, or other intraoral lesions and from intraductal obstruction by a stone or viscous plug. The differential diagnosis should also include lymph-node metastasis, especially from squamous cell carcinoma of the oral cavity. Differentiation from a dentigerous abscess can be difficult. Actinomycosis and tuberculosis are rare differential diagnoses.

**Treatment:** An acute attack is treated with antibiotics, analgesics, and anti-inflammatory agents. Excision of the gland is often necessary (Fig. 6.14).

### Myoepithelial Sialadenitis and Sjögren Syndrome

**Synonym:** benign lymphoepithelial lesion

**Definition:** This is an autoimmune form of chronic sialadenitis marked by a gradual decline in saliva production. A sicca syndrome may develop (Fig. 6.15). Sjögren syndrome consists of myoepithelial sialadenitis accompanied by keratoconjunctivitis sicca (decreased lacrimation) and a rheumatoid-type disorder (rheumatoid arthritis, lupus erythematosus, polymyositis, scleroderma).

**Pathogenesis:** Myoepithelial sialadenitis is an autoimmune disease characterized by the formation of antibodies directed against antigens of the salivary duct epithelium. Histologically, the gland exhibits parenchymal atrophy, interstitial lymphocytic infiltration, and islands of myoepithelial cells.

**Symptoms:** The disease predominantly affects women 50–60 years of age. Both parotid glands are diffusely swollen and doughy, with very little pain or tenderness. The end stage presents with atrophy of the gland. Many patients have a Sjögren syndrome with xerostomia, keratoconjunctivitis sicca, and an accompanying rheumatic disease. Sicca syndrome is manifested by troublesome oral dryness, infections of the oral mucosa, and dental caries.

**Diagnosis:** Tests show nonspecific signs of inflammation such as an elevated ESR. Cytoplasmic antibodies against excretory-duct epithelium (parotid antibodies) can generally be detected. Sialography demonstrates the bare duct system (“leafless tree” pattern) but is rarely necessary. It is often helpful to take an in-
cisional biopsy from one of the minor salivary glands of the lip, which are involved in approximately 60–70% of cases and exhibit typical histologic changes. It may also be necessary to sample tissue from the parotid gland itself.

**Complications:** Frequent complications are chronic recurrent parotitis and the sequelae of sialopenia (mucositis, dental caries) and decreased lacrimation (ulceration, infection). There is an increased incidence of non-Hodgkin lymphomas both within and outside the gland.

**Treatment:** Immunosuppressant therapy is indicated only in the setting of a rheumatic disease. Otherwise treatment is supportive and includes the use of oral saliva substitutes and eye drops to replace the lacrimal fluid. A regimen of 3 × 5 mg/day pilocarpine can be tried to stimulate salivation.

**Radiation Sialadenitis**

**Pathogenesis:** External irradiation or radioiodine therapy (iodine is excreted in the salivary glands) incites an inflammation of the salivary glands with atrophy and transient or permanent oral dryness. At doses less than 15 GY, the injury is reversible. Higher radiation doses cause irreversible injury with a variable degree of partial recovery. The most severe damage is to the serous glandular acini, causing a quantitative decrease and qualitative change in the saliva. Sicca syndrome results in dental caries and mucosal inflammation.

**Symptoms:** The main symptoms are xerostomia and burning of the tongue, frequently combined with hypogeusia or ageusia. A full-blown sicca syndrome may develop. Some degree of recovery may occur over a period of years, but many patients are left with distressful complaints.

**Treatment:** Symptomatic measures should be tried such as stimulating the production of saliva (e.g., 3 × 5 mg/day pilocarpine), administering a saliva substitute, or frequent hydration (e.g., with sage tea).

**Prophylaxis:** An application of amifostine during combined treatment with cisplatin and external irradiation may help to protect the function of the salivary glands.
6.7 Tumors of the Salivary Glands

Approximately 70% of salivary gland tumors are benign. The benign/malignant differentiation of a tumor can generally be accomplished by the history, inspection, and palpation. This was covered in 6.4, p. 140. The present unit describes the most common tumors; less common types are reviewed in Table 6.5.

Benign Tumors

General Aspects of Diagnosis and Treatment

**Diagnosis:** The presence of a painless, soft or tense, mobile nodule in the salivary gland, unaccompanied by other symptoms, signifies a benign salivary gland tumor. Clinically benign salivary gland tumors should not be investigated by incisional biopsy due to the risk of tumor cell dissemination and facial nerve injury.

**Treatment:** Biopsy and treatment are generally accomplished in one step. The treatment of benign salivary gland tumors consists of complete removal with a margin of healthy tissue; otherwise the risk of local recurrence is markedly increased. The specimen provides material for intraoperative frozen-tissue histology and for a definitive pathologic diagnosis. Since most tumors are located in the parotid gland, a *lateral parotidectomy* is the most common procedure done for benign salivary gland tumors. First the facial nerve trunk is identified, and the glandular tissues lateral to the pes anserinus of the nerve are excised. Most benign tumors can be removed in this way while safely preserving the facial nerve.

The possibility of temporary or permanent facial nerve injury exists in any type of parotid gland surgery, and the patient must be informed of this prior to the operation.

Pleomorphic Adenoma (6.5)

Archaic name: benign mixed tumor

Despite its histologic pleomorphism, the epithelial nature of pleomorphic adenoma has been well established (Fig. 6.16). It is the most common adenoma of the salivary glands, occurring predominantly in the parotid gland. Women are affected more frequently than men. Generally the tumor is surrounded by a pseudocapsule. Multifocal tumors are rare, but it is important to appreciate multifocality from a treatment standpoint.

**Symptoms:** The clinical presentation is that of a painless, firm or nodular salivary gland tumor that is freely movable, usually confined to one side, and shows no evidence of malignancy.

**Fig. Pleomorphic adenoma**

**Diagnosis:** The history, inspection, palpation, and ultrasound examination are sufficient for the diagnosis of small and superficial tumors. FNAB (p. 138) can identify the tumor as a pleomorphic adenoma in a fairly large percentage of cases. With larger tumors, CT or preferably MRI will furnish precise information on tumor extent and location. Definitive diagnosis relies on examination of the surgical specimen.

**Treatment:** See above.

**Prognosis:** The prognosis is good following a technically sound tumor resection. Prompt surgery is recommended even for smaller tumors, as this will make it easier to remove the tumor and also preserve the facial nerve. Both can be difficult to accomplish with larger tumors. Carcinoma is known to occur in pleomorphic adenoma but is rare.

Cystadenolymphoma

**Synonyms:** Warthin’s tumor, papillary cystadenoma lymphomatosum

**Definition:** Typically located at the inferior pole of the parotid gland, cystadenolymphoma is the most common monomorphic adenoma and the second most common tumor of the parotid gland. These tumors are occasionally bilateral, and 90% occur in males.
6.5 Benign parotid tumor: a case report

**History:** A 52-year-old woman presents with a painless nodule below the right ear. The patient states that the nodule has always been “about the same size” but that it may have gradually enlarged over time. Otherwise the patient feels well.

**Findings:** At examination, a tense nodule approximately 2.5 cm in diameter, freely movable relative to the skin and underlying tissues, is palpable below the right earlobe. The nodule is most clearly visible from behind and is located over the posterior part of the vertical ramus of the mandible. No other palpable abnormalities are noted, and there is no evidence of lymphadenopathy. Otoscopic findings are normal. On intraoral examination, massage of the parotid gland elicits normal salivary flow from the Stensen duct. Facial nerve function is symmetrical.

**Further tests:** Ultrasound demonstrates a solid mass in the parotid gland with no evidence of other abnormalities or enlarged lymph nodes. The next study is FNAB, which is done easily and painlessly since the nodule is directly subcutaneous and easily palpable. The pathologist describes cells from a benign adenoma, most likely a pleomorphic adenoma.

**Recommendation and informed consent:** The patient is advised to have the lesion surgically removed along with the lateral portions of the parotid gland.

The patient is informed about the risk of facial nerve injury, the increase of risk with further tumor growth, and the rare possibility of carcinoma development. The division of cutaneous nerves (great auricular nerve) may cause some degree of numbness in the cheek and auricle. Also, sweating may occur over the parotid gland during eating several months after the operation as a result of nerve damage ( gustatory sweating, Frey syndrome, or auriculotemporal syndrome).

**Treatment:** The patient consents to the proposed surgery. She is hospitalized, and the operation is performed under general anesthesia. Facial nerve function is continuously monitored during the operation by electromyography. This makes it easier to identify the nerve trunk, over which the tumor is located. Following the trunk, the surgeon identifies the facial nerve branches that run medially to the tumor. A lateral parotidectomy is performed, and the intraglandular tumor is completely removed along with its capsule and the surrounding tissue. Frozen-section histology confirms the presumptive diagnosis of pleomorphic adenoma. A suction drain is placed at the resection site, and the wound is closed. The patient shows no abnormalities of facial nerve function after the operation. The drain is removed on the second postoperative day. The wound is healing well, and the patient is discharged on the fourth postoperative day.

Pathogenesis: It is believed that the tumor forms from inclusions of glandular parenchyma in lymph nodes. Presently uncharacterized viruses may also play a role.

Symptoms: The typical patient is a male over age 60 who presents with a relatively soft, indolent swelling of the inferior pole of the parotid gland, which may be nodular in some cases. The swelling is not painful, and there are no functional deficits. Bilateral tumors are present in approximately 10% of cases.

Diagnosis: Fine-needle aspiration biopsy generally does not contribute to the diagnosis of cystadenolymphoma. Ultrasound may demonstrate one or more typical cysts. The definitive histologic diagnosis is made from the surgical specimen.

**Differential diagnosis:** Cystadenolymphoma can be difficult to differentiate from lymphomas in the parotid gland and from lymphoepithelial cysts (in HIV). Salivary duct cysts and branchiogenic cysts may also be confused with cystadenolymphoma.

**Treatment:** Treatment consists of pericapsular excision of the tumor in the inferior parotid pole. An expectant approach may be taken in cases with a typical clinical presentation.

**Prognosis:** Good. Malignant transformation does not occur, and complaints are rare. The disease may be complicated by an infected cyst.

Malignant Tumors

**General Aspects of Diagnosis and Treatment**

The clinical signs of a malignant salivary gland tumor were discussed on p.140. The TNM classification is used to describe the clinical spread of malignant salivary gland tumors.

**Diagnosis:** The suspicion of a malignant tumor can often be confirmed by FNAB. If there is clinical suspicion of a malignant salivary gland tumor, MRI or CT should be performed to define the extent of the tumor and check for invasion of adjacent structures. Obtaining a specimen for a definitive histologic diagnosis is closely related to treatment planning and can critically influence management. The pathologist can evaluate frozen sections during the operation. Because frozen tissue histology is sometimes equivocal, a two-stage surgical procedure may be necessary, especially if the facial nerve must be resected. In this case the definitive plan of treatment is decided only after the surgical tumor biopsy has yielded a definitive histologic diagnosis.

**Treatment:** The treatment concept for malignant salivary gland tumors is based on removing the tumor as completely as possible and then irradiating the tumor region. Complete tumor removal is not always possible or desirable, however, for a variety of reasons. In the case of the parotid gland, possible resection of the facial nerve must always be considered. The case
Table 6.5 Overview of benign and malignant tumors of the salivary glands

<table>
<thead>
<tr>
<th>Name of tumor</th>
<th>Percentage of all salivary gland tumors</th>
<th>Course and prognosis</th>
<th>Other features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benign tumors of epithelial origin</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pleomorphic adenoma</td>
<td>40–50%</td>
<td>Good prognosis</td>
<td>Most common in the parotid gland</td>
</tr>
<tr>
<td>Cystadenolymphoma</td>
<td>15%</td>
<td>No malignant transformation</td>
<td>Monomorphic adenoma, most common in the parotid gland, can be bilateral, 90% in men</td>
</tr>
<tr>
<td>Other monomorphic adenomas (e.g., salivary duct adenoma)</td>
<td>5%</td>
<td>Good prognosis</td>
<td></td>
</tr>
<tr>
<td>Malignant tumors of epithelial origin</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mucoepidermoid carcinoma</td>
<td>5%</td>
<td>Depends on tumor differentiation; 5-year survival rate for low-grade tumors is 90%</td>
<td>Most common in the parotid gland or minor salivary glands of the palate</td>
</tr>
<tr>
<td>Acinar cell carcinoma</td>
<td>2–3%</td>
<td>5-year survival rate 75%</td>
<td>More common in women, usually in the parotid gland</td>
</tr>
<tr>
<td>Adenoid cystic carcinoma</td>
<td>3%</td>
<td>5-year survival rate 75%</td>
<td>Arises from minor salivary glands in 70% of cases; perivascular and perineural infiltration</td>
</tr>
<tr>
<td>Adenocarcinoma</td>
<td>3%</td>
<td>Poor prognosis</td>
<td></td>
</tr>
<tr>
<td>Carcinoma in pleomorphic adenoma</td>
<td>5%</td>
<td>Poor prognosis</td>
<td>Develops in 3–5% of untreated pleomorphic adenomas</td>
</tr>
<tr>
<td>Squamous cell carcinoma</td>
<td>2%</td>
<td>Poor prognosis</td>
<td>Requires differentiation from intraglandular lymph-node metastases</td>
</tr>
<tr>
<td>Undifferentiated carcinoma</td>
<td>3%</td>
<td>Poor prognosis</td>
<td></td>
</tr>
<tr>
<td>Benign tumors of nonepithelial origin</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lipoma</td>
<td>1–2%</td>
<td>Good prognosis</td>
<td>Usually in the parotid gland, easy to remove</td>
</tr>
<tr>
<td>Hemangioma, lymphangioma</td>
<td>2%</td>
<td>See “Features” column</td>
<td>Lymphangiomas tend to recur</td>
</tr>
<tr>
<td>Others</td>
<td>&lt;1%</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>Malignant tumors of nonepithelial origin</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sarcoma</td>
<td>&lt;1%</td>
<td>Poor prognosis</td>
<td></td>
</tr>
<tr>
<td>Lymphoma</td>
<td>1–2%</td>
<td>Like other lymphomas</td>
<td>Mostly non-Hodgkin lymphomas</td>
</tr>
<tr>
<td>Others</td>
<td>&lt;1%</td>
<td>–</td>
<td></td>
</tr>
</tbody>
</table>

may require a total parotidectomy with or without facial nerve preservation or the (subtotal) removal of the temporal bone, mandible, skin, vessels, and/or cervical lymph nodes, depending on the extent of the tumor. Reconstructive measures may be considered for the facial nerve (nerve grafting), the internal carotid artery (bypass or replacement graft), and skin (advancement flaps, see 3.4, pp. 36–39).
Mucoepidermoid Carcinoma

**Definition:** Mucoepidermoid carcinoma occurs predominantly in the parotid gland and minor salivary glands of the palate. It is the most common malignant tumor of the salivary glands and can occur even in young patients.

**Pathogenesis:** A distinction is made between well-differentiated *low-grade tumors* (approximately 75% of cases) and more poorly differentiated *high-grade tumors* (approximately 25% of cases). The grade of tumor differentiation determines the prognosis, with higher-grade tumors having a considerably poorer prognosis than lower-grade lesions. Metastasis usually occurs by the lymphogenous route; hematogenous (pulmonary) metastasis is less common.

**Symptoms:** The tumor begins as a nonpainful swelling (Fig. 6.12, p.141). Sooner or later it causes pain, facial nerve palsy, and lymph-node metastases, depending on the tumor grade.

**Diagnosis:** see p. 149.

**Differential diagnosis:** see Table 6.5.

**Treatment:** Treatment consists of primary radical excision. The surgery may include the resection and reconstruction of the facial nerve and portions of the temporal bone. If lymph-node metastases are present, a neck dissection (see p. 334) is added. Most patients undergo postoperative radiotherapy.

**Prognosis:** The prognosis depends strongly on the tumor grade. The 5-year survival rate for patients with well-differentiated, low-grade tumors is 90%.

Acinar Cell Carcinoma

**Definition:** Acinar cell carcinoma is a locally invasive tumor that grows predominantly in the parotid gland and has little tendency to metastasize. The peak incidence is between 40 and 60 years of age, and women are affected more than men.

**Pathogenesis:** As in the case of mucoepidermoid carcinoma, various tumor grades are distinguished. Most acinar cell carcinomas are relatively well differentiated, however. The tumor tissue includes acinar and ductal components. Granules positive on periodic acid-Schiff (PAS) staining and amylase can be detected.

**Symptoms:** The symptoms depend on local tumor growth and infiltration.

**Diagnosis:** see p. 149.

**Differential diagnosis:** These tumors mainly require histologic differentiation from adenocarcinoma and adenoid cystic carcinoma.

**Treatment:** See mucoepidermoid tumor.

**Prognosis:** The prognosis is good following a complete resection (5-year survival rate 70%).

Adenoid Cystic Carcinoma

**Definition:** The clinical picture of adenoid cystic carcinoma is marked by a highly variable course and by perivascular and perineural infiltration. The tumor may take a relatively benign, slow course, but lymph-node metastases are common. Other cases take a fulminating course with rapid recurrence and widespread hematogenous metastases.

**Histology:** Despite its clinical malignancy, this tumor has a relatively benign and well-differentiated histologic appearance. Solid, cribriform, and tubular types are distinguished, and these histologic categories may bear some relationship to the prognosis.

**Symptoms:** The symptoms are dependent on tumor location. Local infiltration leads to pain or nerve deficits caused by typical, early perineural infiltration by the tumor.

**Diagnosis:** See p. 149. The lungs and skeleton should be screened for additional sites of hematogenous metastasis.

**Treatment:** Treatment is surgical due to the poor radiosensitivity of the tumor. The value of an ultraradical procedure that sacrifices major structures (facial nerve, temporal bone, carotid artery) is disputed due to the slow growth rate and frequent metastasis and should be decided on a case-by-case basis. Surgery may be appropriate even when pulmonary metastases are present, depending on the frequently slow growth of the metastases.

**Prognosis:** Aside from a few fulminating cases, adenoid cystic carcinoma generally grows slowly, but cures are infrequent. The 5-year survival rate is 75%, but the 10-year survival rate is only 30%.

Other Carcinomas and Malignant Tumors

Other malignant tumors are generally rare and, like most tumors, occur predominantly in the parotid gland. The prognosis tends to be unfavorable. These tumors are summarized in Table 6.5.
### Anatomy and Physiology of the Ear

#### 7.1 Basic Anatomy and Physiology of the Ear
- Peripheral Auditory System: 154
- Central Auditory System: 158

#### 7.2 Anatomy and Function of the Cochlea
- Structure of the Cochlea: 160
- Function of the Cochlea: 161
7.1 Basic Anatomy and Physiology of the Ear

The goal of this is to describe the basic anatomic and physiologic principles of the auditory and vestibular apparatus and thus provide a basis for understanding the complex disorders that can affect this sensory organ. Given the complicated structure of the cochlea, its anatomy and function are described in a separate unit (7.2, pp. 160–163). Further details, especially pertaining to the vestibular apparatus, are presented in Chapters 10–14.

Hearing in humans plays a central role in social communication, while also serving as a warning and orientation system that functions in all spatial directions. The vestibular system is important for maintaining balance and stability and for spatial orientation. The following systems are responsible for carrying out these functions:

- The peripheral auditory and vestibular system (the "ear"). The function of the peripheral auditory system is to perceive periodic air-pressure variations and process them into neural signals. The vestibular function of the ear is described in 13.1 Clinically Relevant Anatomy and Function of the Vestibular System, pp. 272–274).

- The central auditory system, which further processes the acoustic information and is particularly instrumental in directional hearing and sound pattern recognition.

Speech is the most important sound pattern for human hearing.

- The central vestibular system, which establishes the connections between the vestibular apparatus and the effectors for spatial orientation and balance (see 13.1, pp. 272–274).

The morphologic and anatomic boundary between the peripheral and central systems is located at the site where the vestibulocochlear nerve enters the brainstem. The functional boundary is formed by the central synapse of the peripheral neuron.

Peripheral Auditory System

The peripheral auditory system is divided into three parts (Fig. 7.1):

- The external ear, consisting of the auricle and external auditory canal
- The middle ear, consisting of the tympanic membrane, tympanic cavity, auditory ossicles, intra-aural muscles, and the air cells of the temporal bone
- The inner ear, which is embedded in the petrous bone. It is subdivided into the vestibule and the semicircular canal system of the vestibular end or-

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**Fig. Peripheral auditory system**

The peripheral auditory system can be divided into three parts: the external ear (blue), middle ear (green), and inner ear (red). The vestibulocochlear nerve is shown in yellow.
gan (see 13.1, Clinically Relevant Anatomy and Function of the Vestibular System, pp. 272–274) and the cochlea, which is the auditory end organ (see 7.2, Anatomy and Function of the Cochlea, p. 160).

The acoustic systems of these three regions are coordinated in such a way that the principal frequencies of speech are transmitted with particular efficiency.

The vestibulocochlear nerve (cranial nerve VIII) is part of the peripheral auditory system. It runs in the internal auditory canal and connects the peripheral end organs to the central nervous system (CNS).

**External Ear**

**Anatomy:** The external ear consists of the auricle (pinna) and the external auditory canal (external acoustic meatus, ear canal). The formative elements of the external ear are composed of flexible cartilage and of bone, which are attached to the skin by their perichondrium and periosteum (see 10.1, Special Anatomy and Examination of the External Ear, p. 208).

**Physiology:** The function of the external ear is that of an acoustic antenna which transmits sound waves to the sensitive middle ear structures in a discriminating way. The auricle and ear canal together form an acoustic funnel that amplifies selected frequency bands, chiefly in the range from 2 to 4 kHz. This explains why noise in this particular frequency range can damage hearing. The amplification does not involve an increase in the amplitude of sound waves but is based on the law of resonance, meaning that certain wavelengths vibrate better, similar to the columns of air in a pipe organ. The resonant frequency may be altered by cerumen, insert earphones, or by the earmolds of hearing aids. Owing to the differential refraction of sound waves by the shape of the auricle, two different acoustic pathways exist: a direct route through the conchal cavi-ty and an indirect route via the helix and anthelix. This slightly longer pathway creates a brief sound delay of approximately 0.2 ms, which has an important role in acoustic analysis, especially for localizing a sound source in the vertical plane. The auricle also functions as a windbreak by creating air turbulence, thereby diminishing the constant acoustic effects of moving air.

On the whole, however, these effects are of relatively minor importance, and the loss of an auricle does not cause serious functional impairment.

**Middle Ear**

**Anatomy:** The middle ear is comprised of air-filled cavities that are subdivided into the tympanic cavity and mastoid air cells. These cavities communicate with the nasopharynx via the eustachian tube (see Fig. 7.1). The middle ear, then, may be viewed as a highly specialized paranasal sinus. Like the sinuses, it is lined by a respiratory ciliated epithelium that contains goblet cells.

Topographically, the middle ear borders on or encloses functionally important structures such as the facial nerve, the internal carotid artery, venous sinuses from inside the skull, the dura, and the inner ear. The principal middle ear space is the tympanic cavity. It is separated from the external auditory canal by the tympanic membrane, which in turn is mechanically linked to the inner ear by a chain of three auditory ossicles. The ossicles form the sound conduction apparatus of the middle ear and, together with the chorda tympani, make up the actual contents of the tympanic cavity. The auditory ossicles are also connected to the two intra-aural muscles, the stapedius and tensor tympani.

**Physiology:** The main function of the middle ear is impedance matching. Due to the different acoustic impedances of air and fluid—i.e., the different resistance that each medium offers to the propagation of sound—the direct transmission of air vibrations to the fluid-filled inner ear would cause more than 99% of the incident sound to be reflected from the fluid surface (Fig. 7.2). The task of the middle ear is to keep this transmission loss as small as possible and, at certain frequencies, transmit virtually all of the energy of the vibrating air to the inner-ear fluid. This is made possible by the approximately 20:1 size disparity between the transmitting surfaces of the tympanic membrane and the stapes footplate.

Like the external ear, the middle ear has a resonant frequency at which sound energy is transmitted most efficiently. That frequency is approximately 1 kHz.

But for the middle ear to maintain effective impedance matching at all times, even in the face of atmospheric pressure changes, it has a second important function to perform: it must equalize the static air pressure, which is in a constant state of flux due to weather and altitude changes.

The greatest atmospheric pressure variations occur in response to significant altitude changes, as in flying or riding in a cablecar. They are many times greater than the dynamic air pressure changes that occur in the acoustic range. An altitude change of 2 m is roughly equivalent to the sound pressure amplitude at 120 dB, bearing in mind that the static pressure, unlike the alternating acoustic pressure, acts in one direction only. Ambient pressure changes are counterregulated by the eustachian tube, which periodically equalizes the pressures.
The middle ear is an impedance transformer that also compensates for static pressure differences (air pressure). a Without the interposed ossicular chain of the middle ear, more than 99% of the sound energy would be reflected at the fluid surface of the inner ear. b The impedance is modified by the difference in area between the tympanic membrane and the stapes footplate. The ossicular chain and its joints are shown schematically in green. With static pressure changes that produce a negative pressure in the middle ear (black line in b), the joints of the auditory ossicles maintain the function of the sound conduction apparatus and protect the inner ear.

pressures between the environment and the tympanic cavity but cannot entirely prevent pressure changes within the middle ear. The articular connections between the auditory ossicles also help to equalize the static pressures. These connections protect the inner ear by preventing extreme displacement of the stapes footplate.

Inner Ear

Located in the petrous part of the temporal bone, the inner ear consists of multiple interconnected ducts that are collectively called the labyrinth.

Membranous labyrinth: The membranous labyrinth is filled with a potassium-rich fluid, the endolymph, and contains cilia-bearing sensory cells that are also known as hair cells. It is divided into the vestibular labyrinth and the cochlea (cochlear labyrinth), which are interconnected by the narrow ductus reuniens. The vestibular labyrinth is composed of three semicircular canals, the utricle, and the saccule. The nature and function of the sensory cells in the vestibular labyrinth are described on p. 272. The utricle and saccule are connected by another duct of the membranous labyrinth, the utriculosaccular duct. Another membranous labyrinthine structure arises from the utriculosaccular duct: the endolymphatic duct (vestibularosaccular duct), which extends to the endolymphatic sac on the posterior surface of the petrous bone (see 7.1). The function of the endolymphatic sac is not fully understood. It appears to have a secretory function and is believed to play a role in endolymph regulation and in the immune processes of the inner ear.

The membranous labyrinth of the cochlea is the cochlear duct (scala media), which makes two and one-half spiral turns. The anatomy and physiology of the cochlea are described in the next (p.160), and those of the vestibular labyrinth are covered fully in 13.1, pp.272–274.

Bony labyrinth: The membranous labyrinth is embedded in the bony labyrinth within the petrous bone. The membranous and bony labyrinth are separated by a space filled with perilymph. The composition of the perilymph, unlike the endolymph, is very similar to that of the extracellular fluid compartment. The bony labyrinth can be subdivided into three parts: the semicircular canal system, the cochlea, and the vestibule. The bony labyrinth encloses the membranous semicircular canals and reproduces their shape. The cochlear duct is attached between the inner and outer walls of the bony cochlea. This arrangement creates two separate ducts called the scala vestibuli and scala tympani, which are connected at the cochlear apex by the helicotrema.

Between the semicircular canals and cochlea, the vestibule forms a large cavity that contains the saccule, the utricle, the base of the cochlear duct, and the connecting ducts of the membranous labyrinth. The oval window that links the inner ear to the middle ear is covered by the stapes footplate and represents the

The vestibular labyrinth (blue) and cochlea (green) are interconnected by the narrow ductus reuniens (dark blue). The neural connection to the cochlear nerve is shown in yellow. For clarity, the vestibular nerve is not shown.
“acoustic entrance” to the labyrinth. It is the site where impedance-matched acoustic vibrations are transmitted from the middle ear to the perilymph. Because the entire bony labyrinth is filled with incompressible perilymph, a “pressure valve” is needed so that the vibrations can be effectively transmitted. This function is served by the round window. Located inferior to the oval window at the end of the scala tympani and sealed by a mobile membrane, the round window provides a second opening between the bony labyrinth and the tympanic cavity.

The perilymphatic space of the bony labyrinth communicates with the subarachnoid cerebrospinal fluid (CSF) space via the perilymphatic duct, known also as the cochlear aqueduct. This duct begins at the scala tympani below the round window and ends at the posterior surface of the pyramid below the internal porous acusticus. Most likely, it is consistently open only in children and is often sealed by fibrous tissue in adults.

Blood supply: The inner ear derives its blood supply from the labyrinthine artery, which usually arises from the anterior inferior cerebellar artery or the basilar artery. It runs with the vestibulocochlear nerve through the internal auditory canal, where it divides into the vestibular artery and cochlear artery. These vessels may anastomose with the middle ear vessels. Several veins drain blood from the inner ear to the superior bulb of the jugular vein and to the inferior petrosal sinus.

Vestibulocochlear Nerve

The vestibulocochlear nerve (cranial nerve VIII) leaves the brainstem as a nerve trunk that has a grossly homogeneous appearance. Functionally, however, it consists of an anterior superior part, the vestibular nerve, which is distinct from the posterior inferior cochlear nerve. This subdivision becomes anatomically apparent in the internal auditory canal, where the vestibular and cochlear nerves appear as separate structures.

In the fundus of the internal canal, the vestibular nerve forms the vestibular ganglion from which various nerve fibers are distributed to the structures of the vestibular end organ (utricleoamphullar nerve, saccular nerve, posterior ampullary nerve). The cochlear ganglion (spiral cochlear ganglion) is not located in the internal auditory canal but in the bony modiolus of the cochlea (see Fig. 7.6, p. 160).

The vestibulocochlear nerve mainly contains afferent fibers, which lead to the vestibular and cochlear nuclei in the brainstem. Efferent fibers are also present, but generally they are less well myelinated than the afferent fibers. The function of the afferent vestibulocochlear nerve can be described as passive information transfer. The information is already transformed into complex neurobiologic signals (action potentials) at the level of the synapses in a process that can be likened to digital data transmission. The precise timing of the transmission is crucial.

The facial nerve (see also p. 290) is anatomically separate from the vestibulocochlear nerve throughout its course but approaches it very closely in the internal auditory canal.
Embryology and Development of the Peripheral Auditory System

The auricle, middle ear, inner ear, and internal auditory canal undergo a more or less separate embryonic development. As a result, a malformation affecting one part of the ear is not necessarily combined with malformations of other parts. On the other hand, malformations of the external auditory canal are frequently associated with anomalies of the middle ear.

The germ layers that give rise to the various auditory structures and the timetable of inner ear development are shown in 7.1.

Central Auditory System

The central auditory system begins in the brainstem at the cochlear nucleus, where the cochlear nerve terminates (Fig. 7.5). The cochlear nucleus, unlike many other nuclei, receives its afferents entirely from one side. Past the cochlear nucleus, the auditory pathways run mainly but not exclusively via the two inferior olivary complexes, the inferior colliculus of the midbrain, and the thalamus to the contralateral areas of the auditory cortex, which are located mainly in the temporal lobe.

The following systems, while not strictly separate anatomically, are distinguishable in a functional sense:

- **Tonotopic system**: The frequency processing that occurs in the cochlea already assigns certain frequencies to specific fibers in the acoustic nerve. This "tonotopic principle" is maintained in some of the central auditory pathways as far as the cerebral cortex.

- **Non-tonotopic system**: Parallel to the tonotopic system are other modes of central processing that are largely independent of frequency analysis and rely on other parameters such as temporal information.

- **Polymodal or polysensory system**: This system establishes connections with other sensory and nonsensory centers at various levels in the CNS. An example is the stapedial reflex, which induces contraction of the stapedius muscle by the stapedius nerve (a branch of the facial nerve) in response to a sufficiently loud acoustic stimulus.

Besides these anatomically and physiologically distinct systems of the ascending auditory pathway, numerous collateral connections are present at all levels. The neurons generally become more numerous at the higher centers; this property of the ascending auditory pathway is called *diversification*.
Sound localization is based largely on the binaural auditory information pathways in the brainstem: sound reaches the ear closer to the sound source earlier and with greater intensity than the ear farther from the source. Without having to identify the sound source in terms of recognition, this system permits one or more acoustic sources to be localized at the same time. Sound pattern recognition, which involves the naming and identification of a sound source, is a cognitive cerebral function that is based on experience and learning. It relies on the preliminary neural processing of the sound information in the cochlea and brainstem. Directional hearing and its associated functions, then, form an essential foundation for sound pattern recognition. Sound identification or pattern recognition includes the separation of “desired” auditory information or sound sources from “extraneous” sources, or noise. While one picture after another is recognized in the visual system, auditory “picture recognition” is essentially dynamic and transitory. The recognition of speech sounds and their combination into syllables, words, phrases, and sentences in human communication is the most important example of recognizing rapidly changing sound patterns. Music may be considered a specialized form of sound pattern recognition. Auditory hallucinations or illusions are sound pattern recognition without a physical correlate in the outside world. Tinnitus also lacks an external physical correlate but does not involve pattern recognition; only “noise” is perceived.

While the cochlear nerve contains only a few efferent fibers, the efferent fibers outnumber the afferents in many of the connections from higher to lower centers in the CNS. They assist in controlling the flow of information.

Function of the Central Auditory System

In the peripheral system, auditory information is collected and relayed to the CNS with maximum fidelity as a complex mix of signals. The task of the CNS is to separate and recognize the auditory signals. Two basic functions are distinguished in this process: sound localization (where?) and sound pattern recognition (what? who?). These two functions are complementary and cannot always be clearly separated from each other. They are very highly evolved in humans, because a human being receives visual information only in the frontal plane and must rely on hearing for orientation in all other spatial planes.
7.2 Anatomy and Function of the Cochlea

As the actual sensory organ for hearing, the cochlea is of key physiologic and clinical importance. The main function of the cochlea is to translate auditory events into a pattern of neural impulses that precisely reflects the nature and timing of the sound stimulus. To perform this neural coding, the cochlea divides the broad frequency spectrum of the sound into narrow frequency bands that are matched to the neural processing capacity of the organ. This shows how the cochlea performs this task with the help of the cochlear amplifier and the tonotopic principle. Clinical implications are also discussed.

Structure of the Cochlea

The bony cochlear canal spirals around the axis of the cochlea (modiolus) for a length of 3–3.5 cm to the cochlear apex. It contains three separate cavities: the scala media, scala vestibuli, and scala tympani.

The basilar membrane (Figs. 7.6, 7.7), which is narrow (0.1 mm) and relatively thick at the base of the cochlea and considerably wider (0.5 mm) and thinner at the cochlear apex, stretches between the spiral lamina, a bony shelf projecting from the modiolus, and the spiral ligament of the outer cochlear wall.

Together with the Corti organ, the basilar membrane forms the floor of the scala media (cochlear duct), which is filled with endolymph. The ionic composition of the endolymph is similar to that of the extracellular space (see also Fig. 7.8).

The scala vestibuli lies above the cochlear duct, contains perilymph, and is separated from the endolymphic space of the scala media by the thin Reissner membrane. Because the composition of the perilymph is the same as in the extracellular space, a potential difference is created at the membrane (see also Fig. 7.8). The scala vestibuli begins at the base of the cochlea in the area of the oval window. At the apex of the cochlea it is connected to the scala tympani by the helicotrema.

The scala tympani lies below the basilar membrane. It is also filled with perilymph and runs downward from the helicotrema to the round window.

The mechanical properties of the basilar membrane change along the course of the cochlea. The membrane is markedly stiffer and less compliant at the base of the cochlea than at the apex. Its resonance is tuned to higher frequencies in the basal area of the cochlea and to lower frequencies in the apical area.

The formation of the passive traveling wave is based on these mechanical properties of the basilar membrane, with a frequency-dependent maximum amplitude occurring at different sites along the cochlea.

The organ of Corti (spiral organ. Fig. 7.8) lies on the inner part of the basilar membrane facing the modiolus and contains the sensory and supporting cells. The tectorial membrane, an acellular structure composed of amorphous material and fibrils, covers the sensory cell region of the Corti organ starting from the spiral lamina.

![Fig. Axial section through the cochlea](image)

The cochlea has been sectioned through the center of the modiolus. The cochlear nerve ends there and divides into separate nerve fibers that run in small bony canals and form the spiral ganglion near the bony spiral lamina. The scala vestibuli and scala tympani, which contain perilymph, are shown in blue. The endolymph-filled cochlear duct is shown in orange.

![Fig. Model of the basilar membrane](image)

Model of an uncoiled cochlea demonstrates the basilar membrane (red), organ of Corti (green), and perilymphatic spaces (blue) in which traveling waves can be produced. These waves produce maximal vibrations of the basilar membrane for high frequencies at the cochlear base and for low frequencies at the cochlear apex.
na. The reticular membrane connects the ciliated surfaces of the sensory cells with one another and creates a partition between the endolymphatic space above the membrane, which contains the cilia and tectorial membrane, and the perilymphatic space below the reticular membrane. Because of this arrangement, the reticular membrane forms a voltage boundary.

Hair cells are mechanoreceptors surmounted by a bundle of stereocilia of varying length on a specialized surface. The stereocilia are arranged in longitudinal rows and packed together in hexagonal arrays. The stereocilia themselves are tiny, stiff rods that are interconnected by transverse fibrils. Any deflection of the stereocilia bundle at its base toward the longest stereocilia generates an adequate excitatory stimulus for the sensory cell. Two types of hair cells are distinguished (Fig. 7.9):

- **Inner hair cells**: Normally the cochlea contains more than 3000 inner hair cells, which are arranged in a single row along the cochlea and are surrounded by supporting cells (see Fig. 7.8). Their stereocilia form a continuous palisade. Each inner hair cell is connected to several afferent fibers of the cochlear nerve.

  The inner hair cells are the actual “hearing cells,” which transform acoustic information into nerve impulses.

- **Outer hair cells**: There are three to four times more outer hair cells than inner hair cells (approximately 12,000) in the cochlea. The outer hair cells are cylindrical, are generally arranged in three rows along the cochlea, and are anchored only at their base and apex by a complex network of supporting cells. Except from their ciliated end, which projects into the endolymphatic space, they are surrounded by perilymph (see Fig. 7.8). Their stereocilia are firmly attached to the tectorial membrane. The outer hair cells have few afferent connections and are supplied mainly by efferent cochlear nerve fibers. We know from experiments on isolated outer hair cells that they can undergo rhythmic contractions up to a frequency of 30,000 Hz. Based on these findings as well as their anatomy and arrangement, the outer hair cells are viewed as the effector or “motor” of the cochlear amplifier.

### Function of the Cochlea

The translation of acoustic information into neural signals is basically a problem of temporal resolution, since much of the information contained in acoustic vibrations lies in their temporal structure. Vibrations in the typical range of several thousand hertz are transmitted to neural structures with a characteristic refractory period, which allows no more than several hundred impulses to be processed each second. This precludes a 1:1 temporal transduction of acoustic information into neural signals. Transduction is also hampered by noise and by very large differences in pressure levels.

The cochlea solves these complex problems with the aid of two different mechanical functions:
• **Frequency analysis:** Certain frequencies are assigned to nerve fibers at specific locations. This is the *tonotopic principle*.

• **Biomechanical amplification:** Vibrations at low amplitude are magnified with the aid of the *cochlear amplifier*.

The functions of the cochlea can be further subdivided into a *macromechanical* function within the fluid compartment and a *micromechanical* function at the cellular level. The macromechanical function is mainly concerned with frequency analysis, while the micromechanical function deals more with amplification. The two functions are closely interrelated, however. With its frequency analysis and amplifier functions, the cochlea is able to process vibrations at the limit of what is physically possible. However, the physiologic details of this process are not yet fully understood.

**Macromechanical Function: Traveling Wave**

This function of the cochlea is based on the special arrangement of the scala vestibuli and tympani and of the basilar membrane (see Fig. 7.7). Sound waves that are transmitted from the stapes footplate to the perilymph generate traveling waves in the basilar membrane. Each frequency causes a maximum deflection at a different site along the membrane (see Fig. 7.10). Described by Bekesy in 1928, this mechanism allows for:

• Passive frequency analysis and

• Tonotopicity—i.e., the representation of certain frequencies in corresponding areas of the basilar membrane, thereby assigning the frequencies to specific nerve fibers.

High frequencies cause maximum vibration of the basilar membrane near the base of the cochlea, while low frequencies induce maximum vibrations near the apex. The resolution of these passive traveling waves is much too crude for a discriminating acoustic analysis, however.

**Micromechanical Function: Cochlear Amplifier**

The micromechanical function serves to *fine-tune* the basilar membrane vibrations produced by the passive macromechanical cochlear function. This is chiefly made possible by the activity of the outer hair cells in the organ of Corti, which amplify low-amplitude vibrations. This fine tuning and amplification by the *cochlear amplifier* (Fig. 7.11) result in a sharp, detailed sound pattern at the basilar membrane. Events are as following: The pressure waves in the perilymph induce a frequency-dependent traveling wave in the basilar membrane (Fig. 7.11: ①). This traveling wave displaces the outer hair cells attached to the basilar membrane (②). Because their cilia are connected to the tectorial membrane, they are deflected radial to the cochlea, creating an excitatory stimulus. This produces an intrinsic vibration of the outer hair cells (③), resulting in a positive feedback that amplifies the vibration. The motile capacity of the outer hair cells is powered by the endocochlear potential, which works like a battery. It generates a voltage of approximately 85 mV based on the different ionic compositions of the endolymph and perilymph (see Fig. 7.8, p. 161).

The endocochlear potential is the largest extracellular potential difference in the human body.

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**Fig. Cochlear amplifier**

- **Tectorial membrane**
- **Inner hair cell**
- **Outer hair cells**
- **Impedance matching (middle ear)**
- **Cochlear amplifier**
- **Transduction**
- **Afferent cochlear nerve fibers**

Schematic representation of the cochlear amplifier, which is based on a positive feedback for low-amplitude vibrations between the basilar membrane and outer hair cells. The circled numbers refer to numbers in the text.

---

**Fig. Basilar membrane vibration at two frequencies**

- **Middle ear**
- **Round window membrane**
- **Basilar membrane**
- **Helicotrema**

Non-scale model of the passive traveling wave. The maximum deflections of the basilar membrane by the traveling wave are frequency-dependent, creating a crude representation of the sound pattern. The pressure changes induced by movements of the stapes footplate are equalized at the round window.
An even greater potential difference of approximately 155 mV exists between the cytoplasm of the outer hair cells and the endolymph.

The potential gradient is maintained by active ion-exchange processes in the stria vascularis, a specialized region of the spiral ligament that borders the endolymphatic duct. The outer hair cells are suspended in this voltage field, and any reduction in the voltage leads to a reduction or cessation of the active amplification process.

**Transduction:** The inner hair cells transform the physical stimulus of the acoustic vibration into nerve potentials. As in the case of the outer hair cells, a deflection of the cilia radial to the cochlea produces an adequate stimulus for the inner hair cells (8). It is not yet known how the deflection of the basilar membrane is transformed into this stimulus for the inner hair cells. The vibration of the basilar membrane does not stimulate the cells directly, and the stimulus is probably evoked by radial streaming of the endolymph.

**Nonlinear Function—Otoacoustic Emissions**

While the passive traveling wave grows in proportion to the sound level, the cochlear amplifier does not function in a linear way and becomes saturated at approximately 60 dB SPL.

The lower the vibrational energy, the greater the amplification factor.

The proportional amplification of weak and strong vibrations would result in an unstable and nonfunctional system.

Nonlinear amplification like that in the cochlear amplifier tends to have natural modes of vibration and is subject to distortion. Because the ear transmits vibrations not only from outside to inside (antegrade) but also in retrograde fashion from the cochlea through the middle ear to the tympanic membrane, which emits the vibrations into the ear canal like the membrane of a loudspeaker, these natural vibrations of the cochlear amplifier can be detected as faint sounds by a small, sensitive microphone placed in the ear canal. These sounds emitted by the cochlea, called spontaneous otoacoustic emissions (SOAEs, Fig. 7.12), occur at certain frequencies in many normal-hearing persons as evidence of a functioning cochlear amplifier. One theory is that they result from a slightly asymmetrical arrangement of the hair cells at certain locations. An acoustic stimulus acting on the cochlea from the outside (e.g., a click) induces evoked emissions that can also be recorded in the ear canal. Distortion products from the cochlear amplifier can also be detected (see Otoacoustic Emissions, pp. 189–191).

**Clinical Implications**

The cochlea is constantly exposed to acoustic stimuli, keeping it in a mechanically active state. The number of sensory cells in the cochlea is relatively small, and it is unlikely that they regenerate in humans. This makes it all the more remarkable that the system generally functions flawlessly and remains stable for many decades.

Mechanical overloading of the cochlear amplifier is the cause of noise-induced hearing loss (see pp. 260–265) and may also be a factor in age-related hearing loss, or presbycusis (see p. 266).

The highly specialized metabolism of the hair cells and stria vascularis is also susceptible to dysfunction. Drugs can lead to cochlear hearing loss in this way. For example, aminoglycoside antibiotics can damage the metabolism of the hair cells, and loop diuretics such as furosemide can alter the metabolism of the stria vascularis, thereby affecting the endocochlear potential.

Loss of the cochlear amplifier is at least partly responsible for the phenomenon of recruitment, or the abnormal growth of loudness. Recruitment is a clinical sign of cochlear hearing loss caused by abnormal dynamics of sound processing. With loss of the cochlear amplifier, soft sounds are not perceived whereas loud sound events are of undiminished loudness because an amplifier is not needed to perceive them.
Audiology (Auditory Testing)

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8.1 Examination of the Ear and Clinical Auditory Testing

The goal of clinical auditory testing is to acquire information on the integrity and sidedness of hearing by means of simple tests. The results of these tests can then be used to select more specific tests for the further investigation of hearing. Like any clinical examination, auditory testing consists of history taking, inspection (otoscopy), and function tests.

History

Every patient should be questioned about the three most important symptoms of an inner ear disorder:
- Diminished hearing (hypoacusis)
- Tinnitus
- Vertigo

If the patient has specific complaints involving the ear or hearing, the examiner should ask about additional symptoms such as pain or aural discharge. Descriptions of any previous ear surgery, tympanic membrane perforation, or other ear injuries are also important.

Inspection and Otoscopy

Examination of the ear and hearing should always start with a thorough inspection of the auricle and its surroundings. Attention should be given to:
- Changes in the shape of the auricle or ear canal
- Surgical scars
- Crusting in the external ear canal and discharge: cerumen, mucus, pus, blood, cerebrospinal fluid (CSF)
- Redness and swelling of the auricle or surrounding areas

Otoscopy

Before performing otoscopy, the examiner should check for tragal tenderness and pull on the auricle to check for pain.

These signs indicate otitis externa (see 10.3, p. 216 and 10.4, pp. 218–223). When present, they warrant a particularly careful otoscopic technique.

Technique (see also 10.1, p. 209): Otoscopy is performed with a hand-held otoscope (Fig. 8.1) or by an otologist using an otomicroscope. The auricle is rotated gently backward and upward for the examination, avoiding excessive traction. This maneuver straightens the external ear canal and brings the lateral cartilaginous part of the canal in line with the medial bony part. The diameter of the ear speculum should conform to anatomic constraints, keeping in mind that a broad speculum provides better exposure and illumination. The speculum is slowly introduced into the ear canal under visual guidance, inserting it past the vibrissae but without touching the bony and pain-sensitive medial portion of the ear canal. This should afford a clear view of the ear canal and tympanic membrane. Abnormalities and cleansing of the ear canal are described in Chapters 10 and 11 (pp. 210–228).

Clinical evaluation of the tympanic membrane: Not infrequently, the anterior angle of the tympanic membrane cannot be seen with the otoscope, because it is obscured by the prominence of the temporomandibular joint. The normal tympanic membrane has a grayish color and variable transparency. With a thin tympanic membrane, it is possible to identify middle ear
structures such as the long process of the incus or the chorda tympani (Fig. 8.2, see also Chapter 11, pp. 228–253). The normal tympanic membrane exhibits the following three properties:

- **It reflects light:** The tympanic membrane is covered by smooth squamous epithelium that reflects light in a typical way. A “cone of light” is often seen in the anterior inferior quadrant, but other reflections may be seen at other sites on the normal tympanic membrane, depending on the position of the membrane. When the smooth epithelium becomes swollen due to inflammation, the normal light reflexes disappear.

- **It is differentiated:** Normal anatomic structures such as the fibrocartilaginous ring and malleus handle can be distinguished (Fig. 8.2). When an acute inflammation is present, these structures can no longer be identified and the tympanic membrane has an undifferentiated appearance.

- **It is mobile:** To perform its function, the tympanic membrane must be able to vibrate. The mobility of the tympanic membrane may be restricted by effusion in the middle ear or by scars or defects in the membrane. Its mobility can be tested by having the patient perform a Valsalva maneuver or by using a Siegle pneumatic otoscope (see 11.1, Examination of the Middle Ear, p. 231). As a rule, the mobility of the tympanic membrane is most clearly appreciated in the posterior superior quadrant.

**Clinical Hearing Tests**

**Tuning Fork Tests**

The goal of tuning fork tests is to differentiate between conductive and sensorineural hearing loss. Two tests are adequate for this purpose: the Weber test (Fig. 8.3) and the Rinne test (Fig. 8.4).

**Conductive hearing loss** is caused by disease of the external auditory canal or middle ear, whereas sensorineural hearing loss has its cause in the cochlea or the neural structures of the auditory system.

Hearing loss is not detected directly with tuning fork tests.

**Technique:** A tuning fork that vibrates between about 250 and 800 Hz is used. Lower frequencies are not suitable for auditory testing due to interference from the perception of low-frequency vibrations. The resonant frequency of the middle ear is approximately 1000 Hz, and test results in this higher range are often equivocal.

The tuning fork should have a broad base with a large surface area. To test bone conduction, the base of the vibrating tuning fork must be pressed firmly against the cranial bone in order to transmit the vibrations to the bone and overcome dampening by the skin.

**Weber test:**

**Technique:** The tuning fork is placed in the midline of the skull, usually on the vertex or the forehead (Fig. 8.3a). The vibrations are transmitted by bone conduction to the cochlea.

**Interpretation:** When hearing is normal, the vibrations are perceived as equally loud on both sides, and so the sound is heard midway between the ears. In an abnormal test, the sound will be lateralized to one side or the other.

- If the patient has sensorineural hearing loss, the tuning fork is lateralized to the better-hearing ear (Fig. 8.3b).
- If the patient has a conductive hearing loss, the tuning fork is lateralized to the affected ear because the vibrational energy is more poorly transmitted from the cochlea through the middle ear and it is more difficult for ambient sounds to reach the cochlea (less masking). As a result, more vibrational energy is present in the normally functioning cochlea, and the sound is perceived as louder (Fig. 8.3c).

**Humming test:** Laterization can also be detected by having the patient hum, since a loud hum also induces vibration of the cranial bone.
The Weber test is performed by placing a vibrating tuning fork on the midline of the skull.

a When hearing is symmetrical, the sound is perceived with equal loudness in or between both ears.

b With unilateral sensorineural hearing loss, the sound is lateralized to the better ear.

c With unilateral conductive hearing loss, the sound is lateralized to the affected side.

Rinne test:

Principle: The Rinne test compares the levels of air and bone conduction in the same ear (unlike the Weber test, which compares the right and left ears).

Technique: To create standard conditions, air conduction is tested by holding the tuning fork just outside the ear canal without touching it, and bone conduction is tested by pressing the tuning fork firmly against the mastoid.

- The patient is told to compare the loudness in the first position (air conduction) with that in the second position (bone conduction).
- If the patient is unsure which is louder, air and bone conduction can be compared by testing for threshold: The tuning fork is struck and pressed to the mastoid, and the patient tells the examiner when the sound becomes inaudible. Then the tuning fork (without being struck again) is shifted to a position just outside the ear canal (see above).

Interpretation: In a normal (positive) test the tuning fork vibration is transmitted to the cochlea better by air conduction than by bone conduction.

- In a positive Rinne test, air-conducted sound is perceived as louder than bone-conducted sound and lasts at least 15 seconds longer (Fig. 8.4a).
- When conductive hearing loss is present, the sound is perceived as louder on the mastoid than outside the ear canal (Fig. 8.4b). The Rinne test is negative.
- When sensorineural hearing is better on one side than the other, it is necessary to mask the opposite ear before performing the Rinne test (see Fig. 8.6a, p. 170). Without masking, the opposite ear may perceive the sound as louder via bone conduction than the test ear via air conduction, leading to a negative Rinne test. The opposite ear is rarely masked in routine tests, however, and the hearing threshold in this situation is usually assessed by pure-tone audiometry (see pp. 178–183).

Interpreting the tuning fork tests: Fig. 8.5 shows the information that can be obtained from a combination of the Weber and Rinne tests. Occasionally, however, these tests may yield “illogical” findings that cannot be definitively interpreted.

Speech Test

The severity of hearing loss can be clinically assessed without instrumented test methods by having the patient listen to and repeat spoken numbers. This is a simple screening test used to detect a threshold difference between the right and left ears rather than make an accurate evaluation of hearing loss. The speech test and tuning fork tests supply the clinical information that is necessary for selecting appropriate audiometric test procedures.

Formerly, the degree of hearing loss was tested by determining the range at which the patient could hear spoken or whispered numbers. These measurements are imprecise and depend on many uncontrolled factors, and today the quantitative degree of hearing loss should be determined only by audiometric testing.
Air and bone conduction are compared in the same ear to determine the auditory threshold for the tuning fork and/or its loudness. 
**a** In the absence of conductive hearing loss, air conduction is perceived as being louder and/or of longer duration than bone conduction. 
**b** When conductive hearing loss is present, bone conduction is perceived as being louder and/or more prolonged than air conduction.

**Technique:** The range for hearing whispered speech and speech at a normal conversational level is tested separately for each ear. The non-test ear should be masked to preclude crossover hearing. This is done by inserting a moist cotton wad into the non-test ear canal and creating a masking noise by gently wiggling the cotton with the finger (Fig. 8.6). Now the examiner whispers two-digit numbers with his head turned away from the patient and instructs the patient to repeat the words aloud. If the patient does not understand the numbers, the examiner presents them again at progressively smaller distances from the test ear. If the numbers are still not understood when whispered just outside the ear canal, the test is repeated with the numbers spoken at a normal loudness level.

The examiner should maintain a constant loudness level during the test and provide a standardized test environment.

The range of hearing can be tested in meters if a large testing suite and an assistant are available (Fig. 8.7). The assistant masks the non-test ear and blocks the patient’s view of the examiner. The patient turns the test ear toward the examiner, who whispers numbers toward the patient from a distance of 6 meters. If the patient does not understand the numbers, the examiner moves closer and determines the range at which the words become intelligible. If necessary, the numbers may be spoken a normal conversational level. It is essential that constant test conditions be maintained.

**Interpretation:** Hearing loss can be stated in terms of the distance at which the numbers are still intelligible. In a normal test, the subject can understand two-digit numbers whispered from approximately 6 meters away. Numbers whispered with the head turned away from the patient can normally be heard at approximately 4 meters.
Fig.  Speech test

Hearing for spoken numbers is tested separately on each side as a simple screening test.

a. The examiner masks the non-test ear with one hand and shields the patient’s view with the other hand.

b. The examiner whispers numbers while turned away from the patient, or whispers them closer to the test ear when hearing loss is present.

Fig.  Hearing range test

To test hearing range, an assistant masks the non-test ear while shielding the patient’s view.

If the numbers are unintelligible or are understood only when whispered just outside the ear canal, the patient is considered to have **severe hearing loss**, at least for high speech frequencies. If the numbers are unintelligible when spoken close to the ear at a normal or even loud level, the patient is considered to have **functional deafness** for speech in that ear.

With some practice, an examiner can reliably assess hearing with this simple test and can recognize differences between the right and left sides.
8.2 Basic Principles of Audiometry

Audiometry is the measurement of auditory functions. Since it employs acoustic stimuli that are physically defined, we begin this unit by reviewing some of the basic physical principles of acoustics. The response to an acoustic stimuli may be recorded in the form of a voluntary reaction, such as pushing a button, or an involuntary physiologic response such as contraction of the stapedius muscle. Behavioral audiometry is based on voluntary responses and is described more fully in 8.3 (pp. 178–183). Involuntary responses, which are measured in objective audiometry, are discussed in 8.4 (pp. 184–191).

The goals of clinical audiometry are the:
- detection,
- localization, and
- quantification of a hearing disorder.

When sound travels from one medium to another, as from air to water, it is either reflected or absorbed. As a rule, sound is partly absorbed and partly reflected, depending on its frequency.

Basic Concepts in Acoustics

Production and Propagation of Sound

Sound is produced by mechanical vibrations (Fig. 8.8) which propagate as sound waves in an elastic medium (air, liquid, or a solid medium such as bone). The sound waves can be described in terms of the following properties:
- Frequency
- Sound pressure
- Propagation velocity

The velocity of sound ranges from 340 m/s in air to approximately 5000 m/s in solid media such as bone.

Fig. Production and propagation of sound waves

Sound is produced by a vibrating source, in this case a tuning fork. It propagates uniformly in a medium. Human hearing is specialized for the perception of sound in air.
The frequency spectra of different auditory stimuli: tone, musical sound, and noise

- **A musical sound** consists of a fundamental frequency plus harmonic overtones, which are integral multiples of the fundamental frequency. The sound is characterized by, and can be recognized by, its harmonic overtones (Fig. 8.9b). Based on the different spectra of the overtones, the listener can easily distinguish a note played on a violin from the same note played on a piano.

- **Noise** consists of sound events containing multiple frequencies that are harmonically unrelated (Fig. 8.9c)—i.e., they are not integral multiples of one another. Noise is by far the most common acoustic stimulus.

The most important sound source for humans is the voice, whose fundamental frequency of approximately 100 Hz in men and 200 Hz in women is produced by vibrations of the vocal cords (see 18.1, pp. 386–389). The spectrum of basic frequencies is individually modulated by the resonance of the upper airways, enabling the voice to serve as a means of identification.

**Sound Pressure**

Sound waves are extremely small fluctuations of atmospheric air pressure caused by the alternating condensation and rarefaction of atoms and molecules. These variations of pressure amplitude can be physically measured. The unit of measurement is the pascal (Pa):

$$1 \text{ Pa} = 1 \text{ N/m}^2 = 10 \mu\text{bar}$$

1 Pa is approximately equal to the sound pressure in a discotheque. By comparison, the atmospheric pressure is on the order of $10^5$ Pa. Human hearing can just perceive sound pressure variations as small as 20 μPa (2 × $10^{-5}$ Pa, the hearing threshold), which is 10 orders of magnitude lower than the atmospheric pressure. The pain threshold is reached at a sound pressure of approximately 20 Pa, which is approximately 1 million times greater than the normal threshold of hearing. Examples of sound pressure values associated with common events are shown in Fig. 8.10.

The range over which a person can perceive an acoustic stimulus without discomfort is called the dynamic range.

The human ear perceives the loudness levels of different sound pressures in a logarithmic fashion, rather than on a linear scale. This logarithmic relationship between stimulus and perception, called the Weber–Fechner law, holds true for all sensory modalities.

This is why a logarithmic scale is used for sound pressures in acoustics and audiology.
Sound Pressure Level (dB SPL)

**Definitions:** A logarithmic sound-pressure scale can be created by relating the measured sound pressure $p$ to a designated reference value $p_0$. The resulting sound pressure ratio is called the level. The International Organization for Standardization (ISO) has defined the reference value as $20\, \mu Pa$. This is the sound pressure at the threshold of hearing—i.e., the pressure at which a normal listener can just perceive a continuous tone between 2 and $3\, \text{kHz}$. The use of logarithms enables us to bring the physical and physiologic scales closer together and to appreciate more clearly the large range of physical values. The sound pressure level is stated in decibels (dB, named for the inventor of the telephone, Alexander Graham Bell). The suffix SPL stands for sound pressure level (see Applications, below). The formula for the sound pressure level $L_p$ is as follows:

$$L_p = 20 \times \log_{10} \frac{p}{p_0} \, [\text{dB SPL}]$$

or, according to ISO 131–1979:

$$L_p = 20 \times \log_{10} \frac{p}{20 \mu Pa} \, [\text{dB SPL}]$$

**Interpretation and calculations:** Since we are dealing with a logarithmic scale, the increase in physical values is disproportionately greater at high decibel levels than at low decibels. For example, when the sound pressure level is increased by $10\, \text{dB}$, from $0$ to $10\, \text{dB SPL}$, the sound pressure increases by $44\, \mu Pa$; but an increase from $100$ to $110\, \text{dB SPL}$ corresponds to a $4.4\, \text{Pa}$ increase, which is $100,000$ times greater in physical terms.

Decibel values cannot be added or subtracted.

When the sound pressure is doubled, the sound pressure level rises by $6\, \text{dB}!$ Additional examples and the relationship between sound pressure and sound power are shown in Table 8.1 and Fig. 8.11.

**Applications:** The sound pressure level (dB SPL) is a physical scale that is widely used in technology, including hearing aid technology. In audiology, it is used in speech audiometry but not in pure-tone audiometry.

**Explanation:** The sensitivity of hearing is frequency-dependent. It is highest in the range of approximately $1–4\, \text{kHz}$ and corresponds to a sound pressure level of about $0\, \text{dB SPL}$ in young, normal-hearing individuals. Hearing becomes less sensitive at higher and lower frequencies, and considerably greater sound pressure levels are needed to achieve a normal threshold. Because of this, a curved line is obtained when the hear-
8.1 Psychoacoustics

Psychoacoustics is a branch of psychophysics that studies the relationship between the physical properties of a stimulus and the behavioral response. When a person responds to an acoustic stimulus, we call it “hearing.” Psychoacoustics, then, studies the relationships between the physical properties of the acoustic stimulus and hearing and thus forms the scientific basis for much of audiology. Concepts such as “threshold” and “loudness” are derived from psychoacoustics. Several of these concepts are briefly described below.

Threshold

Threshold may refer to the smallest difference that can be perceived between two auditory stimuli that differ with regard to some physical property. This is called the difference threshold. In audiometry, however, the term “threshold” almost always means the absolute intensity threshold, or the minimum sound pressure level of an acoustic stimulus that can still be perceived. In clinical pure-tone audiometry, this threshold is determined for sine-wave tones presented at different frequencies. In this test the sound pressure level of the tone is varied and the subject indicates whether or not he still hears the tone.

Regardless of the method of threshold determination used in clinical audiometry, different threshold values may be determined for different test subjects even though the subjects have identical hearing from a psychoacoustic standpoint. One subject may have to detect a tone convincingly and unequivocally before signaling that it is heard, while another may respond to a much fainter signal. This will yield two different thresholds that relate less to actual hearing ability than to the internal criteria, or bias, of the persons tested.

In psychoacoustics, special methods are used to measure these criteria along with the threshold so that the actual threshold of hearing can be ascertained. These methods basically involve presenting the acoustic stimuli in random order along with equally long periods of silence. The subject responds after each period, indicating whether or not he has heard an acoustic signal. If the level of the stimulus is well above the auditory threshold, the discrimination is easily made. But if the stimulus level is well below the auditory threshold, the stimulus and the silent period will be detected or undetected with equal frequency. To make a threshold determination, it is necessary to determine the ratio of detected and undetected stimuli and silent periods at various levels between these extremes. The actual threshold can then be determined mathematically.

These methods are very time-consuming and are not practical for clinical audiometry. When ordinary clinical methods are carefully applied in a cooperative patient, the differences that are associated with different methods are small and have no diagnostic significance, particularly since the threshold in pure-tone audiometry is determined to an accuracy of only 5 dB.

Loudness level and loudness

The intensity of a stimulus is described physically by its level. But the sound pressure level is a poor criterion for describing the subjective perception of loudness, because this perception depends not only on the sound pressure level but also on other physical parameters such as the duration and spectrum of the stimulus.

The loudness level is a psychoacoustic quantity that is used to compare the subjective loudness of a sound with the loudness of a tone at 1000 Hz. When the 1000-Hz tone is perceived as having the same loudness as the sound, then the sound pressure level of the tone in dB SPL describes the loudness level.

The unit of subjective loudness is the phon. Thus, a 1000-Hz tone at 60 dB SPL has a subjective loudness of 60 phons. But a 50-Hz tone, for example, must have a sound pressure of approximately 80 dB SPL to reach a loudness level of 60 phons.

The values on the phon scale are often poor for describing the subjective perception of a gain in loudness as the sound intensity increases. This prompted the development of a separate scale for subjective loudness based on the sone. One sone is equivalent to 40 phons at a frequency of 1000 Hz. Adding 10 phons doubles the sone value, and so 50 phons are equivalent to 2 sones, 60 phons to 4 sones, etc.

The subjective psychoacoustic units of measure for loudness level and loudness are not used for ordinary technical measurements. Street noise, for example, is measured using correction factors and standard filters—usually the A filter, which is matched to the auditory threshold. The measured sound level is designated by the attribute (A), i.e., dB (A). Again, however, the measurements are not necessarily a good indicator of subjective loudness.

Masking

The perception of a sound event can be diminished by other simultaneous or near-simultaneous sound events. This process, called masking, is a general psychoacoustic phenomenon that, like loudness, cannot be accurately described using simple spectral and temporal parameters.

Masking is important in audiometry when only one ear is being tested. The non-test ear is masked with noise that prevents perception of the test stimulus.

Adaptation and auditory fatigue

Adaptation is a physiologic change in the perception of an acoustic stimulus that occurs in response to constant stimulation. Neural excitation decreases over time, usually accompanied by a rapid decline in loudness perception. If adaptation exceeds a certain measure, which depends on the physical properties of the stimulus, abnormal adaptation is said to be present. Formerly, this phenomenon was applied clinically in the diagnosis of neural injury (see 8.3, Carhart decay test, p. 182).

Auditory fatigue is different from adaptation; it denotes a gradual raising of the auditory threshold during or following an acoustic stress. Some degree of temporary threshold shift (TTS) consistently occurs when the acoustic stress exceeds a certain level.
The normal auditory threshold forms a curved line when plotted against a physical sound pressure level scale (dB SPL, red curve and red scale on the right). A scale with corresponding correction factors is used clinically (dB HL, green scale on the left), and the normal auditory threshold (green) is plotted as a horizontal line. The dB HL scale directly indicates hearing loss relative to a normal population (0 dB HL).

**Audiologic Examination**

**Goals of Audiologic Examination**

Audiologic examination is designed to test the functions of hearing. It has the following main goals:

- Detect a hearing disorder
- Classify a hearing disorder (diagnostic audiometry)
- Quantify a hearing disorder

The various methods that are used in audiometry are of varying usefulness in achieving these goals. An effort must be made to address the clinical problem as efficiently as possible using the most appropriate test methods.

**Methods of Audiologic Examination**

Audiometric tests may be used to *diagnose* or *screen for* hearing impairment (Fig. 8.13). As a rule, the patient already has a known or presumed hearing disorder in diagnostic examinations, whereas screening is designed to detect an unrecognized hearing disorder. Audiologic screening is used to detect hearing problems in newborns, for example, or for the early detection of noise-induced hearing loss in occupationally exposed persons.

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**Decibel Scale for Hearing Levels (dB HL)**

When the pure-tone threshold is measured in audiology, the physical sound pressure is of less interest than comparing the measured threshold with a normal threshold. For this reason, the values measured in dB SPL are corrected to values determined by the ISO based on the auditory thresholds of normal-hearing 18-year-olds for different frequencies (ISO 389 and 7566). The threshold at 250 Hz, for example, is approximately 25 dB SPL.

This relative decibel scale is called the dB HL (“hearing level”) scale. In this type of graph, the normal hearing threshold is represented as a horizontal line at the 0 dB hearing level, making it easier to read the audiogram. We are still dealing with a sound-pressure scale, however, that is based on the physical pressure of the acoustic stimulus.
Both voluntary and involuntary patient responses are measured in diagnostic audiology and in audiologic screening.

Methods in diagnostic audiology are classified according to the nature of the tested response:
- **Behavioral audiometry** (see 8.3, pp.178–183) is based on an active and usually voluntary response from the test subject.
- **Objective audiometry** tests hearing functions based on “objectively” measured parameters that represent an involuntary physiologic response (see 8.4, pp.184–191).

**Site-of-Lesion Determination**

The cause of a hearing disorder may lie at any level of the auditory system, from the cerumen in the external ear canal to circulatory disturbances in the auditory cortex. An important aspect of clinical and audiologic testing is site-of-lesion determination—i.e., localizing the causal pathology to a particular structure or structures. A precise clinical history and physical examination (see 8.1, pp.166–170) are important for narrowing the diagnosis. The clinical examination should be able to distinguish between:
- **Conductive hearing loss**, where the lesion is in the ear canal or middle ear; and
- **Sensorineural hearing loss**, where the lesion involves the cochlea or the neural structures of the auditory system.

**Quantitative Classification of Hearing Impairment**

Hearing disorders can be classified into various grades of severity based on audiometric findings (Table 8.2).

This type of classification is useful only for orientation purposes and says nothing about the actual degree of hearing handicap, which depends on many personal and social factors.
This flowchart shows the most important audiologic tests that are used in making a site-of-lesion diagnosis.

Table 8.2 Classification of hearing loss by severity

<table>
<thead>
<tr>
<th>Designation</th>
<th>Hearing loss in dB</th>
<th>Hearing loss in %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal hearing</td>
<td>&lt; 20</td>
<td>0–20%</td>
</tr>
<tr>
<td>Mild hearing loss</td>
<td>20–40</td>
<td>20–40%</td>
</tr>
<tr>
<td>Moderate hearing loss</td>
<td>40–60</td>
<td>40–60%</td>
</tr>
<tr>
<td>Severe hearing loss</td>
<td>60–90</td>
<td>60–80%</td>
</tr>
<tr>
<td>Profound hearing loss</td>
<td>90–110</td>
<td>80–95%</td>
</tr>
<tr>
<td>Deafness</td>
<td>&gt; 110</td>
<td>100%</td>
</tr>
</tbody>
</table>

Congenital deafness refers to the absence of hearing. Hearing develops during the first years of life and is closely related to speech and language development. With congenital deafness this development fails to occur, and the central patterns of phonation are not established. As a result, nonhearing individuals have fundamentally different speech and language concepts compared with individuals who have lost their hearing.

In addition, various computational methods can be used to determine the percentage hearing loss based on the pure-tone audiogram or speech audiometry. These methods are important for insurance matters and in rehabilitation.

Acquired deafness refers to a loss of the sense of hearing. It is difficult to rule out the vibrational perception of low frequencies, and so deafness is often defined functionally as a complete loss of speech comprehension.
8.3 Behavioral Audiometric Testing

Behavioral audiometric tests are the most commonly used test methods in diagnostic audiology. They are based on an active, usually voluntary response from the test subject. In principle, these methods can test the entire auditory system including higher cognitive functions. Clinical testing is based largely on the subjective auditory response to tones and speech signals:

- Pure-tone audiometry, which is the most common audiometric test method; and
- Speech audiometry.

Pure-Tone Audiometry

Threshold Determination

In pure-tone audiometry, the sensitivity to pure sine-wave tones is measured by determining the hearing threshold (see 8.1, p.174). The threshold is usually measured at frequencies from 125 Hz to 8 kHz, increasing by octaves or half-octaves. This is done separately for the left and right ears, and the thresholds are tested for both air conduction and bone conduction.

**Equipment:** An electronic device called an audiometer is used to generate pure tones of varying frequency and loudness and control their presentation. Frequencies below 125 Hz are difficult to distinguish from vibratory sensations; and with tones higher than 8 kHz, the sound pressure level cannot be accurately calibrated with ordinary headphones. Special audiometers are available for measuring thresholds from 8 to 16 kHz (high-tone audiometry), but these tests show greater interindividual variation than routine audiometry.

A bone vibrator requires considerably more energy to produce sound than headphones, which is why the threshold for bone conduction can be measured only to maximum values that are 40–50 dB lower than those for air conduction. More distortion occurs at higher frequencies (> 4 kHz), and the measurements become less reliable.

**Technique:** The tones are first presented to one ear only by air conduction using headphones or special insert phones. Cross-hearing is prevented by masking the non-test ear with noise (p.169). The threshold for bone conduction is measured with a vibrator pressed against the mastoid or forehead. This device sets the cranial bones and cranial contents into vibration, transmitting the test sound to the inner ear.

**Interpretation:** With proper calibration and normal sound conduction, the thresholds for air conduction and bone conduction should be equal (Fig. 8.15a). If the air conduction threshold is higher than the bone conduction threshold (i.e., if perception by air conduction requires a higher loudness level), the subject has a conductive hearing loss (Fig. 8.15b).

If sensorineural hearing loss is present, no significant difference is found between the thresholds for air and bone conduction. The hearing threshold is raised, often more at high frequencies than at low frequencies (Fig. 8.15c and Fig. 12.1a, p.258). The following audiographic signs indicate a mixed hearing loss:

- Greater air conduction loss compared with bone conduction, indicating impaired sound conduction
- An increased threshold for bone conduction (Fig. 8.15d)

The threshold in the pure-tone audiogram may show typical patterns that reflect the nature of the hearing disorder, such as a notch at 4–6 kHz signifying noise-induced hearing loss (Fig. 12.5, p.262), low-frequency hearing loss at the onset of Ménière disease (Fig. 13.14a, p. 285), or conductive loss accompanied by a bone conduction notch at intermediate frequencies (Carhart notch) in otosclerotic stapes fixation (see pp.251–253).

Suprathreshold Tests

In addition to the pure-tone threshold, measurements are also performed at suprathreshold levels. These measurements can detect recruitment (see 8.3, p.182) or abnormal adaptation (see 8.1, p.174). Examples of suprathreshold tests are the Fowler test, Lüscher test, and short increment sensitivity index (SISI) test.

Simpler and more reliable tests (usually objective) are now available for obtaining these measurements. The classic suprathreshold tests are rarely practiced any longer.

Speech Audiometry

The essential functions of human hearing include the perception and recognition of speech. Consequently, the use of speech signals has a major role in audiometric testing. Speech audiometry is particularly important in hearing rehabilitation and fitting patients with hearing aids. Speech signals display typical patterns that result from a broad frequency spectrum and rapid changes
of frequencies and levels in periods of milliseconds. These patterns form the basis for speech detection and speech recognition.

Speech audiometry generally measures the recognition (understanding) of speech rather than the threshold for the detection of speech signals.

**Principle:** The speech material is available in standardized form on compact disks and is reproducibly presented at designated levels using an audiometer. The material can be presented to one ear using headphones or to both ears simultaneously using loudspeakers in the sound-field environment. The test material may consist of syllables, words with a set numbers of syllables (e.g., monosyllabic words or words with two long or equally stressed syllables called spondees), or sentences. Unlike sustained tones or noise, the level of speech signals cannot be stated precisely but only as a statistical average. Speech audiometry uses specific level measuring functions in the frequency and time domain (filters and incrementation slopes) to determine the speech sound levels. The test subject repeats what is understood at different sound levels, and these responses are used to plot
a discrimination function or speech recognition curve (Fig. 8.16).

**Interpretation**: The speech audiogram indicates the percentage of syllables, words, or sentences that the subject has heard correctly in each test series. Thus, the result of a typical speech audiogram depends not only on hearing but also on higher cognitive functions such as language comprehension, memory, and motor speech. Individual factors such as native language and vocabulary also influence audiometric speech recognition.

**Speech Tests**

Speech tests can be differentiated according to their speech material. **Open-set tests** use single words or phrases, and the patient repeats what he or she is hearing. **Closed-set tests** give the patient a selection of words, and one of these words is presented. The patient chooses the word from the selection which he was hearing.

Routine clinical speech audiometry uses primarily open-set tests. They consist typically of several lists of words with a fixed number of syllables. Each list may have between 10 to 50 words, and the lists are usually phonetically balanced, meaning that the distribution of the various sounds in these words corresponds to that of the language in general.

The PB-50, W-22, or NU-6 are examples of commonly used open-set tests consisting each of lists of 50 monosyllabic words. The words are presented at a designated sound level, and the percentage of correct responses is plotted against the sound levels (speech loudness level in dB SPL). This yields a speech recognition curve, known also as the performance-intensity function (Fig. 8.16). This function will shift to the left and become steeper when words with more than one syllable are being tested. Typically used lists are composed of spondees, which are two-syllable words spoken with equal emphasis on each syllable (pancake, hardware, playground).

The performance-intensity function is steepest at 50% comprehension and flattens out at lower and higher levels. The speech hearing loss is most accurately read at the point where a line parallel to the abscissa intersects the curve at (almost) a right angle. This point is called speech-recognition thresholds (SRT). The scale for speech hearing loss, therefore, is usually placed at the 50% comprehension level (the zero baseline for numbers is at 18.5 dB SPL, yellow scale in Fig. 8.16). In patients with **conductive hearing loss**, the performance-intensity function shows a parallel shift toward higher sound levels. Even so, 100% comprehension is still achieved at sufficiently high levels. **Sensorineural hearing loss** leads to a flattening of the performance-intensity function for monosyllabic words. Loss of intelligibility and a decline in speech recognition at higher sound levels are signs of abnormal speech processing like that caused by damage to the cochlea or neural structures (Fig. 8.17).
8.2 Additional speech tests

**Closed set speech tests**: Words or sentences can be presented as a closed test. Many different tests exist for monosyllabic words, in which a selection of words with similar sounds is presented. From these words, the patient selects the word that he was hearing as a stimulus. Examples are the California Consonant Test (CCT), in which the stimulus consists of one of four words, such as “pin – thin – tin – kin”, or the Four Alternative Auditory Feature (FAAF) test, in which the initial or final consonant in consonant–vowel–consonant words, such as “bad – bag – bat – back” or “gab – dab – tab – cab”, are varied. Closed sets can also be applied for sentences. An example is the commonly used Synthetic Sentence Identification (SSI) test. Ten synthetic sentences are constructed so that each successive group of three words in a sentence are meaningful, but the entire sentence is not (“Go change your car color is red”). Such sentences are thought to be more difficult to guess.

**Sentence tests**: As a rule, meaning in auditory communication is conveyed in sentences rather than single words. Sentence tests are therefore used in speech audiometry for fitting hearing aids and other rehabilitative measures. Sentence tests are particularly useful for testing speech perception in noise. The noise presented in these tests has the same frequency content as speech and can therefore mask spoken sentences. While the noise is usually presented at a constant level, the speech sound levels are varied until a prescribed degree of speech recognition is achieved. The absolute sound pressure level of the sentences, which depends on the selected noise level, is of less interest in these tests than the difference between the noise level and the speech level, known as the signal-to-noise ratio (SN ratio).
8.3 Other behavioral audiometric tests

Besides the behavioral audiometric tests used in routine clinical audiometry (pure-tone and speech audiometry), there are a number of other behavioral audiometric procedures that are used for specific purposes.

Dynamic range scaling
As noted in 7.2, p.163, sensorineural hearing loss is apt to produce distortion effects at suprathreshold levels. A typical distortion effect is an abnormal increase in loudness perception, called recruitment. Although the hearing threshold is increased, with the result that tones at a relatively high sound level are either not heard or faintly perceived, the opposite may occur in the perception of suprathreshold sounds—i.e., tones that are well tolerated by normal-hearing individuals are perceived as louder or uncomfortable. As a result, the range of useful loudness levels, called the dynamic range, is significantly reduced in patients with sensorineural hearing impairment.

The reduction and distortion of the dynamic range play a major role in fitting patients with hearing aids and cochlear implants, because soft signals should be amplified as much as possible whereas loud signals should not. Consequently, an audiometric evaluation of the dynamic range should be performed in patients fitted for a hearing aid.

**Technique:**
- The simplest method is to determine the uncomfortable loudness level (ULL). This is done by determining the level of a pure tone, or preferably noise with a narrow frequency spectrum, at which the test subject perceives the signal as uncomfortably loud.
- Dynamic range measurement (dynamic range scaling) evaluates how subjects perceive increasing loudness within their dynamic range. The subject rates the subjective loudness of signals presented at various loudness levels and at various frequencies (500, 1000, 2000, 4000Hz) as “very soft,” “soft,” “medium,” “loud,” “very loud,” or “too loud.”

Carhart decay test
Abnormal adaptation (see 8.1, p.174) can be detected with the Carhart decay test.

**Technique:** Tones of approximately 0.5 s duration are used for ordinary threshold determination. After the hearing threshold has been determined, a continuous tone 5 dB above the threshold is presented for 60 s. When the tone is no longer perceived, the level is increased by 5 dB (this may be done several times if necessary).

**Interpretation:** In subjects with normal adaptation, the level must be increased by no more than 10 dB. Higher threshold decay values indicate abnormal adaptation. If the decay is 30 dB or more, retrocochlear impairment is present.

Tests for nonorganic hearing loss

**Indications:** Behavioral audiometric tests may elicit an intentionally or unintentionally false response indicating an increased hearing threshold. Malingering refers to the deliberate feigning of hearing impairment by a normal-hearing individual. Other malingerers may exaggerate the severity of an existing hearing problem. A far more common phenomenon, however, is self-deception by a patient who has psychogenic hearing loss. Objective tests (see 8.4, pp.184–191) are available that can quickly disclose the feigning of hearing loss.

There are also a number of special behavioral audiometric tests that can suggest or confirm psychogenic hearing loss or malingering, but they are not widely practiced today. An experienced examiner may suspect a “false” hearing threshold based on certain behavioral cues (reaction time, changing responses).

Feigning is also suggested by a discrepancy between the pure-tone threshold and speech recognition threshold. The loudness of speech signals is much more difficult to assess subjectively than the loudness of pure tones. As a result, subjects with nonorganic hearing loss may show less hearing impairment in speech audiometry than in pure-tone audiometry.

**Methods:**
- Special behavioral audiometric methods used in patients with unilateral nonorganic hearing loss are the crossover hearing test (Langenbeck test), which reveals the absence of a normal crossover hearing threshold at approximately 40–50 dB, and the Stenger test. The latter test is based on the observation that subjects lateralize a binaural acoustic signal to the ear in which the signal is perceived as louder and do not perceive the signal in the other ear. A tone is presented that is well above the threshold for the “good” side. The malingerer will not respond when this clearly audible tone is accompanied by stimulation of the “poor” side with a slightly louder tone because the tone is perceived only on the “poor” side and is no longer heard on the “good” side. With an organic hearing loss, the additional tone on the affected side does not affect the test because it is not perceived.

- Bilateral nonorganic hearing loss can be detected by the Lombard reading test. While the patient reads aloud, bilateral noise is presented through headphones to disrupt the patient’s perception and monitoring of his own voice. With a nonorganic hearing loss, this masking causes the malingerer to falter and speak more loudly but will not affect the patient with organic hearing loss.

A similar effect is achieved by recording the subject’s voice and playing it back to him with a slight delay. This feedback makes it impossible for a hearing subject to read aloud (Lee test).
8.3 Other behavioral audiometric tests (continuation)

Tests for central hearing disorders

The most important central auditory functions are directional hearing and auditory pattern recognition, which includes central speech recognition. As a result, speech audiometry always tests central auditory functions as well, especially when testing is done with background noise. Tests for central hearing disorders are designed to detect a central auditory processing disorder separate from peripheral hearing. This can be accomplished only to a degree, however, since it is rarely possible to distinguish precisely between peripheral and central hearing disorders. Often a combination of objective and behavioral audiometric tests are used to detect central hearing disorders. The main objective tests are the stapedius reflex and auditory evoked potentials (see 8.4, pp. 185–191).

Methods:

- Numerous behavioral audiometric tests of central auditory functions are based on binaural speech recognition under difficult conditions. Various options are available:
  - Different spectra from the same speech signal are presented to the right and left ears (binaural summation, fig. a).
  - Segments of the speech are presented alternately to the right and left ears (alternating binaural speech, fig. b).
  - Dichotic method—i.e., different, competitive speech signals are presented to the right and left ears (fig. c).

Directional hearing can be tested by determining the temporal or level difference in an acoustic stimulus between the two ears at which the sound is still perceived as originating at the center of the head. Complex sound-field setups make it possible to measure directional hearing directly (angle in degrees). Directional hearing is managed chiefly by brainstem neurons of the central auditory pathway. Many central hearing disorders do not cause significant impairment of directional hearing, and therefore this examination is of only minor importance in central auditory testing.

Tinnitus measurement

**Indications:** We must be very cautious in assessing the diagnostic value of tinnitus measurements, especially for site-of-lesion determination. The reproducibility of tinnitus measurements is poor, and so they are not very useful for follow-up. They are mainly important in disability compensation and can aid in better understanding the complaints of a tinnitus sufferer and establishing a rapport.

**Methods:** Tinnitus (see 12.1, p. 256) is a subjective sensation that cannot be tested objectively and can be evaluated only by behavioral audiometric methods.

In **tinnitus matching**, an attempt is made to find a tone or noise that most closely matches the subjective character, frequency, and loudness of the tinnitus. The matching signal may be presented ipsilaterally or to the opposite ear with unilateral tinnitus. The pitch (frequency) is determined first, then the loudness. The loudness of tinnitus usually measures only about 5–10 dB above the hearing threshold; higher values are unusual.

In **tinnitus masking**, an attempt is made to cover up the tinnitus with noise. The examiner determines the loudness of the masking noise at which the patient no longer hears the tinnitus. Every case of tinnitus cannot be masked by noise, however, and occasionally the masking will cause the tinnitus to disappear for a short time or, very rarely, for a longer period. This phenomenon is called residual inhibition and can be selectively tested.
8.4 Objective Hearing Tests

Unlike the subjective psychoacoustic methods used in behavioral audiometry, objective audiometry makes it possible to test hearing without eliciting an active response from the patient. Objective audiometry employs tests that measure hearing functions based on involuntary physiologic responses and “objective” parameters. These responses may consist of the stapedial reflex (see Imittance Measurements), the bio-
electric potentials of neural structures (see Auditory Evoked Potentials), or the acoustic vibrations of the cochlea (see Otoacoustic Emissions). Objective test findings aid in the interpretation of behavioral audiometric results. Objective audiometric tests are also useful in infants, small children (see 9.2, p. 202), and patients with mental or cognitive impairment.

**Methods:** Three main types of response are measured clinically in objective audiometry using various techniques:

- Imittance measurements: changes in the acoustic impedance of the tympanic membrane are measured with an intra-aural probe.
- Auditory evoked potentials (AEPs): acoustically evoked bioelectric responses of the cochlea, auditory nerve, auditory tract neurons, or cerebral cortex are analyzed using surface electrodes and averaging techniques (see also pp. 186–188).
- Otoacoustic emissions (OAEs): a microphone probe is used to measure sound events in the ear canal that are produced by spontaneous or acoustically evoked active biomechanical vibrations in the cochlea (see also pp. 188–191).

**Interpretation:** It cannot always be determined with complete confidence whether or not a response is present in objective audiometry. But modern, computer-assisted methods of measurement will generally establish whether a stimulus response has occurred.

The objectivity of these methods relates to the selection of the stimulus response, not to the interpretation of the test.

**Imittance Measurements**

**Definition:** The impedance of an acoustic system is a measure of the resistance that the system (e.g., the middle ear) offers to the absorption of sound waves. A system with a high acoustic impedance reflects most of the sound energy and absorbs very little. Conversely, a system with low impedance absorbs a large amount of sound energy in the form of vibrations. Sound absorption is also referred to as the compliance of the tympanic membrane. The middle ear transforms the sound waves in air in such a way that they can induce waves in the cochlear fluid with little resistance (impedance matching by the middle ear, see 7.1, p. 155).

**Principle:** The acoustic impedance of the external ear canal and tympanic membrane can be measured with an intra-aural probe. The probe emits a tone at a certain frequency, usually 220 Hz, into the ear canal. The impedance value of this probe tone depends on the overall acoustic system comprising the ear canal, tympanic membrane, middle ear, and cochlea as well as the frequency of the tone and individual factors. A freely vibrating tympanic membrane, for example, will absorb more energy and reflect less energy than a stiff tympanic membrane. The impedance value will be lower. The absolute value of impedance is of less interest in audiometry than the impedance changes that are caused by specific external manipulations. Two main types of impedance testing are performed clinically (Fig. 8.18):

![Diagram of Imittance Measurements](image_url)
• **Tympanometry**, which provides a graphic representation of the impedance changes caused by applied air pressure in the external ear canal; and

• The **stapedial reflex (SR)**, which produces an acoustically evoked change of impedance.

**Tympanometry**

Selectively raising or lowering the air pressure in the external auditory canal causes a stiffening of the middle ear, thereby increasing the acoustic impedance in the ear canal. As a result of this, more sound is reflected from the tympanic membrane. Normally the air pressures in the ear canal and middle ear are equal and correspond to the atmospheric pressure. In this condition, the tympanic membrane has the lowest impedance (resistance) and therefore absorbs sound best. The greater the positive or negative pressure in the ear canal, the greater the “acoustic stiffness” of the tympanic membrane, and the lower its sound absorption or compliance. The tympanogram is a graphic representation of compliance changes as the applied air pressure is varied over a negative-to-positive range. This requires that the intra-aural probe be hermetically sealed in the ear canal. The pressures are usually varied over a range of ±300 mmH2O or daPa. Pathologic changes in the tympanic membrane and middle ear lead to a change in compliance, which correlates clinically with various tympanogram shapes (Fig. 8.19a–c). With partial atrophy of the tympanic membrane or an ossicular discontinuity, the tympanogram may also exhibit multiple peaks.

**Stapedial Reflex (SR)**

**Physiology** (Fig. 8.20): The stapedius muscle inserts on the stapes, and its contraction has the effect of stiffening the sound conduction apparatus. This changes the impedance of the middle ear and tympanic membrane, and the impedance change can be measured with a probe placed in the ear canal. The stapedius muscle contracts as a reflex in response to acoustic stimuli of a certain intensity. With normal hearing, a tone of approximately 80–90 dB HL is sufficient to evoke the stapedial reflex. Broad-band stimuli evoke the reflex at sound levels approximately 10–20 dB lower.

**Definitions:** When sound is delivered to only one ear, the stapedius muscles on both sides contract via the acousticofacial reflex arc. The stapedial reflex on the acoustically stimulated side is called the uncrossed or ipsilateral stapedial reflex. The reflex recorded on the opposite side is called the crossed or contralateral stapedial reflex. The stapedial reflex threshold is the minimum sound pressure level needed to produce a measurable change in tympanic membrane impedance.

![Normal and abnormal tympanogram patterns](image)

- **a** Type A: normal finding
- **b** Type B: flat curve
- **c** Type C: negative pressure peak

- The normal tympanogram has a prominent, sharp peak between +100 and –100 daPa.
- The type B tympanogram is flatter or has a very low, rounded peak. This indicates immobility of the tympanic membrane, which may be due to fluid in the middle ear or tympanic atelectasis.
- The type C tympanogram has a peak in the negative pressure region below –100 daPa, consistent with impaired middle ear ventilation.

A type A or C tympanogram must be present in order to test the stapedial reflex.

**Technique:**

- For **ipsilateral testing**, the “probe ear” is stimulated with tone pulses at 500 to 4000 Hz or with broadband stimuli at incremental sound pressure levels that are 70–90 dB above the hearing threshold. The first impedance change recorded in response to the probe tone is equal to the **stapedial reflex threshold**.

- For **contralateral testing**, which follows the same principle, the “probe ear” or “response ear” is different from the “stimulus ear.”

**Interpretation:** Absence of the stapedial reflex or an increased threshold for the reflex may be caused by a lesion at various sites in the reflex pathway.
The stapedial reflex is evoked by peripheral acoustic stimulation. The reflex arc extends from the cochlear nucleus to the superior olivary complex and the facial nerve nuclei in the brainstem.

- **Ossicular chain** pathology (e.g., disruption or stiffening due to otosclerosis, see 11.4, pp. 251–253)
- Abnormal sound reception by the **cochlea** and/or **auditory nerve** in a patient with cochlear or retrocochlear impairment (e.g., vestibular schwannoma, see 15.3, p. 308)
- Lesion of the **brainstem** (e.g., multiple sclerosis, hemorrhage)
- Lesion of the **facial nerve** (e.g., idiopathic facial nerve palsy, see 14.2, p. 294)
- Disease of the **stapedius muscle** (e.g., myasthenia gravis)

**Applications of Immittance Measurements**

**Tympanometry** is used in diagnosing middle ear pathology. It is of minor value by itself, however, and should always be interpreted in conjunction with otoscopic examination of the tympanic membrane. **Measurement of the stapedial reflex** is always performed after tympanometry. It is useful for the investigation of numerous hearing disorders. The stapedial reflex threshold should be determined on both sides for both the crossed and uncrossed reflexes. The various stapedial reflex patterns are useful in differentiating middle ear hearing loss from cochlear and retrocochlear hearing loss. Moreover, the difference between the subjective hearing threshold and the stapedial reflex threshold provides a measure for evaluating recruitment. This difference is 60 dB or more in normal-hearing subjects. In patients with cochlear hearing loss and abnormal recruitment, it may be possible to evoke the stapedial reflex at only 10 dB above the auditory threshold ("objective" or Merz recruitment).

**Auditory Evoked Potentials (AEPs)**

**Principle:** The physiologic process of hearing involves a great many bioelectric potential changes that take place in the cochlea, auditory nerve, and central nervous system (CNS). These potential changes can be utilized in the objective testing of auditory function.

**Technique:** Auditory evoked potentials (AEPs) are recorded from the scalp using needle or surface electrodes. As in an electroencephalogram (EEG), the potentials from many cells are recorded simultaneously by this technique. The potential changes caused by the auditory system are not detectable in an ordinary EEG trace because their amplitude is so small relative to the total activity of the CNS. **Averaging** makes it possible to record very small potential changes that are buried in noise. It involves the iterative summation of a short EEG segment in a computer, making certain that a constant temporal relationship is maintained between the EEG segment to be analyzed and a constant, repetitive acoustic stimulus. The intermittent stimulus evokes specific, uniform potentials during this time interval, which always occur at the same time and can be amplified by the repetitive summing of the EEG segment. This summation also tends to reduce unwanted, randomly timed background potentials that do not correlate with the auditory stimulus (stimulus-independent EEG activi-
ty). Since positive and negative background potentials are added together, they cancel out after a sufficiently large number of summations. When the summed entries are averaged, the displayed potentials can be assigned temporally to the acoustic stimulus as auditory evoked potentials.

Classification and Terminology of Auditory Evoked Potentials

The properties and shapes of auditory evoked potentials (AEPs) depend partly on the time at which they occur after presentation of the acoustic stimulus, or their latency (in milliseconds). AEPs with a short latency occur very shortly after the stimulus and originate from structures that respond very quickly to the stimulus. Several types of electrical response audiometry are distinguished based on the different sites of origin and latencies of the AEPs:

- **Electrocochleography (ECochG)**: measures the potentials arising in the cochlea and auditory nerve. These potentials occur approximately 1–3 ms after the stimulus is presented.

- **Auditory brainstem response (ABR)** (also known as brainstem electrical response audiometry): measures the potentials arising in the auditory nerve and brainstem structures, with a latency up to approximately 10 ms.

- **Auditory middle latency response (AMLR)** audiometry: measures potentials with a latency of 10–100 ms that originate in the thalamus and primary auditory cortex.

- **Cortical evoked potentials (CEP)** (also known as cortical electrical response audiometry): measures potentials with a latency of 100–1000 ms.

The term “electrical response audiometry” (ERA) is often used in audiology as a synonym for AEP.

Auditory Brainstem Response (ABR)

The auditory evoked potentials that are most commonly recorded for diagnostic purposes are the brainstem potentials, usually referred to as the auditory brainstem response. The ABR occurs during about the first 10 ms after an acoustic stimulus is presented. It is usually evoked by a click stimulus that lasts only a few milliseconds and has a broad frequency spectrum. To record the ABR, the stimulus must be repeated from 1000 to 2000 times and the EEG responses are averaged. Adhesive surface electrodes placed on the vertex and over the mastoid can record a typical waveform that is virtually unchanged even during sleep and under general anesthesia (important in small children). It is characterized by the presence of five or six waves numbered from I to VI as described by Jewett. With a normal brainstem response, the individual waves can be roughly assigned to specific anatomical structures (Fig. 8.21). This cannot be done with an abnormal ABR.

**Indications:** The main clinical applications of the auditory brainstem response are in differentiating between cochlear and retrocochlear hearing loss and in the objective measurement of the hearing threshold.

- Both the shape of the curve (absence of waves, indistinct waves, etc.) and the latent period between waves I and V (normal interpeak latency is approximately 4.3 ms or less) are important in the diagnosis of retrocochlear hearing loss and the exclusion of a tumor in the cerebellopontine angle or internal auditory canal. A prolonged interpeak latency is a sign of retrocochlear hearing loss and should prompt further investigations (see 12.1, Diagnostic Imaging, p. 258).

- The ABR can also be important in the diagnosis of neurologic diseases (multiple sclerosis, ischemic brainstem lesions, etc.).

- The ABR is important for threshold testing in pediatric audiology (see 9.2, p. 202). Because potential thresholds are of interest in determining hearing thresholds in small children, it is important to have quiet examination conditions. While the ABR can usually be measured in adults without difficulty, infants and small children must be tested while sleeping or sedated and occasionally under general anesthesia.
Auditory evoked brainstem potentials are also tested intraoperatively to monitor hearing. The ABR mainly tests hearing at middle and high frequencies (>1 kHz). It is more difficult to obtain information on low-frequency hearing.

Interpretation: The most important parameters of the ABR are the time intervals between the waves and the threshold for the detection of wave V. Normally, wave V can be detected at only about 10 dB above the hearing threshold.

Otoacoustic Emissions (OAEs)

The vibrations produced by the biomechanical amplifier of the cochlea (see 7.2, pp. 161–163), either spontaneously or in response to an acoustic stimulus, are transmitted in retrograde fashion across the ossicles to the tympanic membrane, which acts like the membrane of a loudspeaker, emitting the vibrations as sound waves into the external ear canal. A sensitive microphone probe inserted into the ear canal can detect these active cochlear vibrations, which are called otoacoustic emissions (OAEs). Otoacoustic emissions are clinically important in that they can be used to test the function of the “cochlear amplifier.” The emissions reflect the functional integrity of the cochlea. The outer hair cells are a particularly important source of OAEs; the auditory nerve is not involved. The detection of OAEs in the ear canal is contingent upon normal middle ear function, for otherwise the cochlear vibrations would not be transmitted to the tympanic membrane.

Classification: Owing to the high sensitivity of the cochlear amplifier, vibrations can arise spontaneously in the cochlea without an external stimulus (spontaneous OAEs). At the same time, an acoustic stimulus of low to moderate intensity will consistently induce cochlear vibrations and emissions (evoked OAEs). Otoacoustic emissions are classified into several types based on the nature of the stimulus:
• **Spontaneous otoacoustic emissions (SOAEs):** These emissions occur without an external acoustic stimulus in approximately 50% of normal-hearing subjects and are detectable as low-level, continuous tones. They have little clinical importance.

• **Transient evoked otoacoustic emissions (TEOAEs):** These emissions are consistently detected in response to a brief stimulus (click) in subjects with normal cochlear function. They are detected using an averaging technique similar to that described for auditory evoked potentials (see p. 186). The measurement of TEOAEs is a commonly used objective audiometric test method.

• **Distortion product otoacoustic emissions (DPOAEs):** Acoustic distortions in the cochlear amplifier can be detected by stimulation with two continuous tones that have different but adjacent frequencies. The measurement of DPOAEs is another frequently used objective audiometric study.

• **Stimulus frequency otoacoustic emissions (SFOAEs):** Stimulation with a sine-wave tone evokes tonal emissions of the same frequency. These emissions are more difficult to detect than TEOAEs and DPOAEs. They have little clinical significance.

### Transient Evoked Otoacoustic Emissions (TEOAEs)

TEOAEs are recorded in response to a brief stimulus (click) and reflect the spectrum of the stimulus. A microphone probe inserted into the external ear canal can record the acoustic signals, which are averaged in a similar way as the bioelectric signals in auditory evoked potentials (see p. 186).

**Interpretation:** The click consistently evokes a cochlear acoustic response (TEOAE) in subjects with normal hearing (Fig. 8.22). Almost always, this confirms the functional integrity of the cochlea and middle ear. TEOAEs do not occur in patients with middle ear disease or a cochlear hearing loss with an approximately 30 dB threshold increase. TEOAEs recorded in normal-hearing infants usually have a greater amplitude than in adults.

### Distortion Product Otoacoustic Emissions (DPOAEs)

DPOAEs are also used in clinical testing. The test involves measuring the otoacoustic emissions that are evoked by stimulation with two continuous tones (Fig. 8.23). When the two stimulus frequencies are properly selected, distortion products occur as interference tones that bear a fixed relationship to the stimulus frequencies but are not identical to them. The use of continuous tones allows for measurements within a narrower frequency range and at higher sound levels than are possible with TEOAEs. As a result, it may still be possible to detect DPOAEs even when the function of the cochlear amplifier is impaired. Automated measuring systems can be used to quickly measure the response of the cochlear amplifier in discrete frequency ranges. On the other hand, it is easier to eliminate artifacts when TEOAEs are measured. The clinical indications for DPOAEs are the same as for TEOAEs.

### Application of Otoacoustic Emissions

The most important application of otoacoustic emissions is for screening cochlear function in newborns, infants, and small children (see 9.2, p. 202). TEOAEs and DPOAEs provide a fast and simple way to test cochlear function without sedation or general anesthesia, thus facilitating the early detection of hearing problems. The majority of hearing disorders in this age group have a cochlear etiology. The location of the hearing loss (middle ear or sensorineural) and its degree cannot be determined by analyzing otoacoustic emissions.

In the absence of OAEs, additional audiological tests such as auditory evoked potentials and behavioral audiometry should be used. OAEs can also be used to investigate nonorganic hearing loss, to objectify audiometric findings in adults, and to assess cochlear function in risk groups (e.g., patients using ototoxic medications).
Fig. Transient evoked otoacoustic emissions (TEOAEs)

a Setup for measuring TEOAEs: measuring probe with microphone and loudspeaker.

b Waveform of the stimulus, which lasts approximately 2 ms. Note that the scale of the stimulus sound pressure (Pa) is 1000 times greater than the scale for the sound pressure responses recorded in the ear canal (see c).

c The waveform recorded 2.5 ms after initiation of the stimulus reflects the time course and amplitude of the sound pressure of the TEOAEs.

d Spectrum of the evoked response (from c), indicating the frequency distribution of the TEOAEs (purple). The orange trace represents noise.
The cochlea is stimulated with two tones, $f_1$ and $f_2$, delivered by a probe. A microphone records the sound pressure in the external ear canal, detecting both the emitted sound pressure of the response and the sound pressure of both primary tones $f_1$ and $f_2$. The curve represents the frequency spectrum of the microphone signal. Besides the stimulus tones $f_1$ and $f_2$, an additional tone with a frequency of $2f_1 - f_2$ is recorded. This tone originates in the cochlea as a DPOAE.
8.5 Rehabilitation and Hearing Aids

The general goal of auditory rehabilitation is to restore or improve auditory communication. Specific rehabilitative measures are tailored to the degree of hearing loss, the needs of the patient, and other individual or social requirements. Hearing aids are frequently prescribed for moderate to severe hearing loss, while cochlear implants (inner ear implants) may be used in patients with profound hearing loss or deafness. Ancillary measures such as auditory training and learning to lip read are just as important as the accurate fitting of a hearing aid. The indications and basic principles of hearing aid care in children are reviewed in 9.3.

Indications and Possibilities of Auditory Rehabilitation

Hearing loss that is measurable by audiometry is always an “impairment”—i.e., it always compromises the physical integrity of the patient. Whether hearing loss is disabling in everyday life depends on factors that include the degree of the hearing loss as well as individual auditory requirements and demands. Thus, measures to improve auditory performance should not be based entirely on audiometric hearing loss but should also take into account the individual auditory disability and the resulting handicap.

Medical treatment is an option for hearing loss only in rare cases.

If the patient with hearing loss does not experience subjective hearing impairment and is not handicapped by the loss, rehabilitative measures will generally be unrewarding due to a lack of motivation.

On the other hand, rehabilitative measures should not be long delayed in patients who are subjectively disabled by their hearing loss, because rehabilitation at an earlier age is more successful and can prevent further disability relating to auditory deprivation.

A variety of rehabilitative options are available, depending on the nature and degree of the hearing loss and the degree of handicap that it imposes.

Surgery: A surgical operation to improve hearing can provide functional restoration of hearing in patients with conductive hearing loss (e.g., stapes surgery in otosclerosis, see p. 252).

Hearing aids: A hearing aid selectively amplifies auditory signals (Fig. 8.24). The hearing aid may be worn behind the ear (BTE aid) or inside the ear canal (ITE aid).

Active middle ear implants: An amplifier is implanted which transforms sound waves into mechanical vibrations. The vibrations are transmitted directly to the cranial bone or ossicular chain.

Cochlear implant: In patients with a complete or almost complete absence of cochlear function, this surgically implanted device can transform sound waves into electrical impulses that directly stimulate the auditory nerve with intracochlear electrodes.

Vibrotactile aids: Acoustic signals are picked up by a microphone and converted to vibrations that are transmitted to the wrist or fingers.

Other assistive devices: A variety of assistive devices can improve communication for hearing-impaired patients. These devices include optical or vibrating wake-up alarms, light flashers, telephone amplifiers, text telephones, television headphones, and digital communication devices such as faxes, text messaging, and the internet.

Training: Hearing-impaired patients can be trained in:

- Selective listening (listening tactics, auditory training)
- The proper use of hearing aids and other assistive devices
- Learning to lip read
- Improving their speech

The goal of rehabilitation is to achieve a maximum restoration of aural communication, which is crucial for social functioning and patient well-being.

Hearing Aid Fitting in Adults

The hearing aid is a special type of acoustic amplifier (Fig. 8.24). Sound from the environment is received by a microphone, amplified, and transmitted by a loudspeaker to the ear. For a hearing aid to work properly, the cochlea must be able to receive the amplified and processed signal and relay it to the auditory nerve. Microchip and digital technology allows for differentiated processing of the acoustic signal and makes it possible to tailor the hearing aid to the patient’s individual hearing loss and requirements.

The fitting of a hearing aid, which is an important means of aural rehabilitation for many hearing-impaired patients, involves a series of steps:

Determining candidacy: Before the patient is fitted with a hearing aid, candidacy must be determined on the basis of an accurate history and audiologic evaluation.
For example, the following conditions have been established in Germany for the prescription of a hearing aid:

- Hearing loss of 30 dB HL or more in the better ear by pure-tone audiometry, measured for at least one frequency in the range from 500 to 3000 Hz.
- In speech audiometry, 80% or fewer correct responses for monosyllabic word recognition at 65 dB SPL.
- The patient must be willing and able to use and maintain the hearing aid.

There should be no reasonable prospect of improving the patient’s hearing by surgery.

**Audiologic examination:** The audiologic examination should include pure-tone audiometry, speech audiometry, and a determination of the dynamic range—e.g., by dynamic range scaling (see 8.3).

**Hearing aid trial:** Based on the clinical and audiologic findings and the desires of the patient, a hearing aid is selected on a trial basis. Fitting of the hearing aid is done by a trained specialist (audiologist or hearing aid specialist). The hearing aid (usually a behind-the-ear or in-the-ear device) is physically and acoustically coupled to the patient’s auditory canal with a custom-fabricated earmold. Changes in the auricle or ear canal (e.g., due to surgery) may require the construction of a new earmold. A poorly fitting earmold may allow acoustic feedback to occur between the microphone and loudspeaker, causing a high-pitched squeal.

Patients with a largely symmetrical hearing loss should be fitted with binaural aids, as these will provide a significant gain (binaural summation), allow directional hearing, and improve speech recognition in background noise. Binaural fitting will also prevent auditory deprivation of the unaided ear.

**Final hearing aid selection:** The hearing aid specialist should offer the patient several devices to choose from. The final selection and fine-tuning of the amplifier can be done only in practical trials. **Speech audiometric testing** with and without background noise, the subjective auditory impression of the patient, and acoustic measurements in the ear canal (in situ measurements) support the final selection. After the fitting, the patient must become accustomed to the device and the auditory sensation it produces in order to derive optimum benefit.

**Follow-up care:** The follow-up regimen includes the points listed above under Training in addition to an audiologic evaluation.

Despite the sophisticated technology, a hearing aid is still a peripheral listening aid that cannot replace or substantially improve the fine frequency resolution of the cochlea or essential central auditory functions.

These limitations can be particularly noticeable in a noisy environment. As a result, hearing aids often do not provide a satisfactory gain in speech recognition when background noise is present.
Additional technical measures can dampen background noise in lecture halls where a microphone is being used. For example, the microphone signal can be directly transmitted as an electromagnetic signal using an induction coil, bypassing the microphone component of the hearing aid. Induction coils of this kind are built into many lecture halls and churches. Another option is direct radio-signal transmission from the speaker’s microphone to the hearing aid (audio input, FM transmitters).

Cochlear Implant in Adults

Even the best hearing aids reach their limits as the degree of hearing loss becomes more severe. When the cochlea contains few or no hair cells able to transform a vibration into bioelectric signals, even the strongest and most discriminating acoustic amplification will be unable to improve aural communication. The cochlear implant (CI) is an option for these cases.

**Principle:** The functional principle of the CI is shown in Fig. 8.25. An electronic receiver is implanted into the temporal bone under the skin. The receiver is connected to an electrode array inserted into the cochlea. The electrodes directly stimulate the auditory nerve and spiral ganglion, functionally bypassing the cochlear hair cells. The acoustic signal that is received by a directional microphone worn behind the ear (Fig. 8.25) is processed by an external speech processor, which is usually worn behind the ear like a hearing aid (2). This processor extracts the useful sound components that are important for speech recognition, transmits the signal through the skin to the implanted receiver, and stimulates the various intracochlear electrodes (3). After stimulating the auditory nerve (4), the signal undergoes further central processing like any acoustic signal.

Marked individual differences occur in the position of the intracochlear electrodes and in the number of residual nerve fibers and ganglion cells that are still present. This makes it necessary to adjust the electrical stimulation, and thus the processing of the speech signals by the speech processor, individually for each patient.

**Indication:** The criteria for prescribing a cochlear implant are as follows:

- **Acquired**, bilateral, predominantly cochlear deafness (postlingual deafness) with a functional auditory nerve and intact central auditory pathway
- **Congenital or early acquired** deafness in children (prelingual deafness), see 9.3, p. 204
- Lack of benefit from binaural hearing aid fitting despite optimum adjustment of the hearing aids
- A motivated patient willing to learn the operation and maintenance of the implant

The cochlear implant requires a partially functional auditory nerve that is responsive to stimulation by the intracochlear electrodes. The electrical responsiveness of the auditory nerve can be tested prior to implantation by the **promontory test**, in which the lowest cochlear turn (promontory) is directly stimulated with a transstympanic electrode. If intact auditory nerve fibers are present, this test will produce an auditory impression (**positive promontory test**). The implantation surgery is a safe, standardized procedure that has relatively few side effects. Later complications, including possible malfunction, are also rare.

**Follow-up care:** Adjustment of the speech processor begins several weeks after the operation and generally lasts for several months. The follow-up program includes:

- A technical check of CI function
- Checking and adjusting the speech processor; and
- Auditory training.

Under the guidance of speech therapists and/or hearing-impaired teachers, the patients learn to interpret the electrical impulses as speech. Speech recognition is further supported by teaching the patients to lip read.

Over 90% of patients derive definite benefit from a CI, and more than 50% can achieve open speech comprehension—i.e., they can understand CI-mediated speech without relying on visual cues (e.g., over the telephone or a public address system).

The success of a cochlear implant is partly influenced by the time of onset and duration of the deafness. In adults, patients with deafness of short duration derive the greatest benefit from the implant.
The sound is received by a directional microphone worn on the ear and fed to the speech processor as an analog signal 
\(\text{①}\). The speech processor, which is worn externally, processes the microphone signal, extracts the speech components that are necessary for comprehension, and converts them into a series of electrical impulses 
\(\text{②}\). A transmitting coil worn behind the ear transmits the impulses as radio frequencies through the skin to the implanted portion of the CI (receiving coil) 
\(\text{③}\). The necessary power supply is also transmitted; the implant itself does not require a separate power source.
Hearing Disorders in Children—Pediatric Audiology

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9.3 Treatment of Pediatric Hearing Disorders
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9.1 Causes and Effects of Pediatric Hearing Disorders

Hearing disorders in infants and small children can be difficult to recognize and often go undetected until the child is older. While hearing disorders in children can have the same causes as in adults, special perinatal causes are also encountered in infants and newborns. The frequency distribution of the causes is also different. The development of speech is closely related to hearing ability, and hearing disorders during this formative period can have specific consequences in terms of language and personality development.

**Epidemiology:** Significant hearing impairment is present at birth in approximately one in 1000 newborns. Most of these cases involve sensorineural hearing loss due to cochlear impairment.

During the first years of life, permanent hearing loss, usually sensorineural, develops in one in 1000–2000 children, and so the number of hearing-impaired children rises by 30–50%. Hearing disorders are much more prevalent in small children and preschoolers. Hearing impairment in this age group, however, is usually a temporary conductive loss secondary to otitis media; this occurs in approximately 3–4% of children.

**Causes of Pediatric Hearing Disorders**

**Classification:** The causes of early childhood hearing loss fall into several categories:

- Hereditary genetic causes
  - Congenital
  - Occurring postnatally or later
- Acquired causes
  - Acquired in utero (also congenital)
  - Acquired perinatally
  - Acquired postnatally or later

**Monosymptomatic Genetic Hearing Loss**

The majority of hearing disorders due to a genetic cause affect only hearing and are described as monosymptomatic. Except for the hearing impairment, the child is normal. Ninety percent of these hearing disorders are inherited as an autosomal-recessive trait, and so the family history is usually negative.

The hearing impairment is not obvious in these cases and, without screening, is often detected later due to abnormal speech development.

The forms of monosymptomatic genetic hearing loss are covered in 12.2 (p. 260) because they are not confined to the pediatric age group.

**Hearing Loss in Genetic Syndromes**

Rarely, multiple gene loci or chromosomes are affected resulting in hearing impairment accompanied by other symptoms. Thus, the hearing loss is one feature of a congenital syndrome. Table 9.1 reviews several of the more than 300 known syndromes that are associated with hearing loss. They are classified according to the affected organs or tissues.

The presence of hearing loss profoundly affects the further development and rehabilitation of children with these syndromes.

Aside from cases with an anomaly of the auricle or ear canal, there are no direct signs that call attention to a hearing disorder when these children are infants. Consequently, children with this syndrome should be specifically tested for hearing impairment as soon after birth as possible.

**Acquired Hearing Loss**

Hearing loss may be acquired before birth (intrauterine, perinatally), during birth (perinatally), or after birth (postnatally). Table 9.2 lists the most frequent causes of acquired hearing loss in newborns and infants.

Besides genetic syndromes, there are some congenital syndromes that can be acquired in utero or perinatally, such as:

- **Rubella syndrome** with cochlear hearing loss, pulmonary stenosis, mental retardation, and microphthalmia
- **Hyperbilirubinemia syndrome** with athetoid cerebral palsy and sensorineural hearing loss
- **Congenital syphilis** with interstitial keratitis, Hutchinson teeth, and sensorineural hearing loss

**Hearing Impairment in Newborns**

It is often difficult to distinguish between a congenital and perinatally acquired hearing loss. Because hearing loss unassociated with other disorders or malformations is not obvious in newborns, delayed speech development is unfortunately the earliest sign of hearing impairment in many unscreened children. A delay in diagnosis makes it even more difficult to distinguish between congenital and perinatally acquired disorders.
Certain structures of the auditory system are particularly susceptible to harmful influences at certain times during the neonatal period. A typical example is keri
nicterus, in which high bilirubin levels in the blood cause damage to central structures. It is also believed that the cochlea in newborns is exceptionally vulnera-
ble to hypoxia, toxic agents (antibiotics), and noise. The maturity of the newborn is an important factor in this regard, and premature infants are at particularly high risk. Overall, hearing impairment is present in approximately 4% of all newborns that require treatment in an intensive-care unit.

Table 9.1 Congenital syndromes that are associated with hearing loss

<table>
<thead>
<tr>
<th>Classification by anomalies</th>
<th>Syndrome</th>
<th>Inheritance</th>
<th>Typical features</th>
<th>Type of hearing loss</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anomalies of the external ear</td>
<td>Mandibulofacial dysostosis (Treacher–Collins syndrome)</td>
<td>Autosomal-dominant</td>
<td>Anomalies of the external and middle ear</td>
<td>X</td>
</tr>
<tr>
<td>BOR syndrome (brachio-otorenal syndrome)</td>
<td>Autosomal-dominant</td>
<td>Anomalies of the external ear</td>
<td>X or X or X</td>
<td></td>
</tr>
<tr>
<td>CHARGE syndrome: coloboma, heart defect, atresia of choanae, retarded growth, genital hypoplasia, ear anomalies</td>
<td>Sporadic</td>
<td>Anomalies of the external ear</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Retinal degeneration and ocular anomalies</td>
<td>Usher syndrome: retinitis pigmentosa and sensorineural hearing loss</td>
<td>Autosomal-recessive</td>
<td>Type I–III retinal degeneration</td>
<td>X (progressive)</td>
</tr>
<tr>
<td>Musculoskeletal disorders</td>
<td>Craniosynostosis: • Apert syndrome • Crouzon syndrome</td>
<td>Autosomal-dominant</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Osteogenesis imperfecta (various forms)</td>
<td></td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Renal function impairment</td>
<td>Alport syndrome</td>
<td>Variable</td>
<td>Chronic nephritis</td>
<td>X (progressive)</td>
</tr>
<tr>
<td>Nervous system disorders (with ataxia)</td>
<td>E.g., Cockayne, Lichtenstein–Knorr, Klippel–Durante</td>
<td>Variable</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Endocrine and metabolic dysfunctions</td>
<td>Pendred syndrome (thyroid dysfunction)</td>
<td>Autosomal-recessive</td>
<td>Patent vestibular aqueduct, possible goiter</td>
<td>X (severe)</td>
</tr>
<tr>
<td>Mucopolysaccharidosis II (Hunter syndrome)</td>
<td>Autosomal-recessive</td>
<td></td>
<td>X or X or X</td>
<td></td>
</tr>
<tr>
<td>Cutaneous and pigmented anomalies</td>
<td>Waardenburg syndrome</td>
<td>Autosomal-dominant</td>
<td>White forelock, heterochromia iridis, telescanthus</td>
<td>X</td>
</tr>
<tr>
<td>Cardiac anomalies</td>
<td>Jervell–Lange–Nielsen syndrome</td>
<td>Autosomal-recessive</td>
<td>Prolonged QT interval</td>
<td>X (severe)</td>
</tr>
<tr>
<td>Chromosome abnormalities</td>
<td>Ulrich–Turner syndrome Trisomy 21</td>
<td>Sporadic</td>
<td></td>
<td>X (frequent) or X (frequent) or X or X</td>
</tr>
</tbody>
</table>
9.1 Anomalies involving the ear

Anomalies of the middle ear
The embryology of the ear was reviewed in 7.1 (p. 158). Conspicuous anomalies of the auricle and external ear canal are frequently associated with an anomaly of the middle ear. On the other hand, isolated auricular anomalies often develop without hearing impairment, and isolated changes in the ear canal often lead to conductive hearing loss. These anomalies are described in 10.1 (p. 211). The significance of external ear anomalies in syndromic conditions is reviewed in Table 9.1. External malformations, unlike isolated middle ear anomalies, are easily diagnosed and should always prompt auditory testing.

Atresia of the ear canal necessarily causes changes in the tympanic membrane plane and malleus. The rest of the middle ear may be normally developed, but there are often concomitant anomalies of varying degree affecting other middle ear structures. Congenital aural atresia is therefore classified as a middle ear anomaly.

Middle ear anomalies with a normal external ear are not detectable by external inspection. They invariably cause hearing impairment, usually consisting of an isolated, significant conductive hearing loss. Middle ear anomalies relevant to hearing involve the ossicular chain and particularly the stapes. Other middle ear structures such as the facial nerve, vessels, or intra-aural muscles may also show anomalies of varying significance.

Middle ear anomalies are classified into various grades of severity. Usually it is sufficient to distinguish between “major” and “minor” anomalies. A more precise analysis of the malformed structures can supply prognostic information that is useful in planning surgical treatment.

Treatment: Bilateral middle ear anomalies should be treated as early as possible with special hearing aids that employ a bone-conduction vibrator. Corrective surgery is usually deferred until preschool or school age. With a unilateral anomaly, it must be determined immediately after birth whether normal hearing is present on the opposite side. If that is the case, there is no need for immediate surgery to improve hearing.

Anomalies of the inner ear
Inner ear anomalies result from an arrest or abnormality of embryonic development between the third and ninth weeks of gestation (see 7.1, p. 158). The division of the otocyst into two anatomically and functionally distinct parts (the pars utriculovestibularis and pars saccocochlearis) plays a key role in the pathogenesis of anomalies, since the division itself may be abnormal (forming a “common cavity”), or independent anomalies may develop in one or both parts after the otocyst has divided.

Case report: inner ear anomaly
A male newborn was found to be deaf shortly after birth. Computed tomography (CT, Fig. a) revealed bilateral inner ear anomalies consistent with cochleovestibular dysplasia. The vestibulocochlear nerve could be identified in the cerebellopontine angle. Cochlear implant (CI) surgery was performed on the right side at 2 years, 4 months of age. When the cochlea was opened for electrode insertion, profuse cerebrospinal fluid (CSF) leakage occurred, and the defect was sealed with connective-tissue pieces and tissue adhesive. Several months later, CSF leakage occurred through the right stapes footplate, which showed dysplastic thinning and bulging. A follow-up CT scan (Fig. b) shows the metal electrode assembly. The CI was left in place at reoperation, and the leak was repaired.

Two years later a CSF leak occurred in the left, nonoperated middle ear as the result of a trivial injury. Again, a defective footplate was found and the leak was surgically repaired. The boy made good hearing progress with his cochlear implant. He understood most ordinary instructions and began to communicate verbally.

Hearing Impairment in Infants and Small Children

A hearing loss produces very few signs in infancy, regardless of its degree of severity. Even when parents notice “something is wrong,” they are often reassured that their child’s development is merely delayed, and auditory tests are not performed. An opportunity is missed to refer the child for special services.

When the slightest evidence of hearing impairment is noted in an infant or small child, an audiologic evaluation should be performed. In many cases, this evidence comes from parents and is too often ignored. “The parents are always right” is a good rule to follow.

An acquired, temporary hearing loss due to otitis media is the most common type of hearing loss in small children and preschoolers. Both acute and chronic forms of middle ear effusion are very common in this age group, and up to 75% of children have this condition for some period of time. The clinical features of otitis media are described in 11.2 and 11.3 (pp. 234-253).

Acute otitis media itself, the subsequent fluid collection in the middle ear, and chronic middle ear effu-
sions lead to varying degrees of conductive hearing impairment. A chronic, bilateral middle ear effusion may be present for months or years without causing any symptoms other than conductive hearing loss. In cases of this kind, or in children who have frequently recurring acute middle ear infections, a prolonged hearing impairment can cause delays in speech and language development with articulation problems, vocabulary deficits, and dysgrammatism (see 19.1 and 19.2, pp. 398–408).

### Effects of Hearing Loss in Children

#### Bilateral Hearing Loss

A hearing loss in infants and small children threatens normal speech and personality development, underscoring the unique importance of hearing loss in this age group.

Hearing is crucial to the development of verbal communication. The central nervous system of the newborn and infant appears to include structures whose maturation is triggered by linguistic stimuli. It also appears that this maturation takes place only during certain critical periods. The longer the child is without auditory stimulation by speech, the more difficult it is to acquire the missed linguistic skills. The impaired speech development and altered acoustic environment have a direct influence on personality development and the relationship of the individual to society. The severity of the handicap depends on various factors. The most important, perhaps, is the degree of hearing loss (Table 9.3). The effects also depend on the time of onset of the hearing disorder. When deafness occurs during the initial months after meningitis, for example, early rehabilitation can build upon previous acoustic experience. This advantage is not available in a congenitally deaf child who has never experienced speech perception through the auditory canal. The earlier and more selectively auditory services are provided, the more effectively the natural developmental periods of the child can be utilized. This emphasizes the critical important of an early diagnosis in hearing-impaired children.

#### Unilateral Hearing Loss

A **mild to moderate** unilateral hearing loss in early childhood usually has no adverse sequelae. Often the child is unaware that he or she has a hearing problem, and the condition is first detected by audiometric testing when the child enters school or even later. With a **severe** unilateral hearing loss, however, it is possible that speech development may be hampered. Usually the child experiences little disability from the disorder. Subtle deficits and handicaps may arise and in some cases are reflected in poor academic performance and delayed speech development. Difficulties with unilateral hearing loss tend to arise in acoustic settings where there is a poor signal-to-noise ratio. Situations of this kind are common in classrooms and can lead to fragmentary perception of the academic material.

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### Table 9.2 Causes of acquired hearing loss in newborns and infants

<table>
<thead>
<tr>
<th>Timing of insult</th>
<th>Classification</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prenatal</td>
<td>Infectious</td>
<td>Rubella, toxoplasmosis, congenital syphilis</td>
</tr>
<tr>
<td></td>
<td>Drug toxicity</td>
<td>Quinine, alcohol, thalidomide</td>
</tr>
<tr>
<td>Perinatal</td>
<td>Infectious</td>
<td>Cytomegalovirus, herpes simplex virus, etc.</td>
</tr>
<tr>
<td>Metabolic</td>
<td>Kemicterus, asphyxia</td>
<td></td>
</tr>
<tr>
<td>Obstetric trauma</td>
<td>Forceps, intracerebral and intracochlear hemorrhage</td>
<td></td>
</tr>
<tr>
<td>Postnatal</td>
<td>Infectious</td>
<td>Meningitis, labyrinthitis, otitis media, mumps, measles</td>
</tr>
<tr>
<td>Drug toxicity</td>
<td>E.g., aminoglycoside antibiotics</td>
<td></td>
</tr>
<tr>
<td>Traumatic</td>
<td>Noise (susceptible period?), head trauma</td>
<td></td>
</tr>
</tbody>
</table>

### Table 9.3 Effects of hearing loss in children

<table>
<thead>
<tr>
<th>Degree of hearing loss</th>
<th>Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mild: 20–40 dB</td>
<td>Unvoiced consonants and sibilants are not heard clearly</td>
</tr>
<tr>
<td></td>
<td>Articulation problems, delay in language acquisition, dyslalia, possible difficulties in school</td>
</tr>
<tr>
<td>Moderate: 40–60 dB</td>
<td>The majority of speech sounds are not heard</td>
</tr>
<tr>
<td></td>
<td>Abnormal speech development with dysgrammatism, a deficient vocabulary, and poorly intelligible speech</td>
</tr>
<tr>
<td>Severe: 60–90 dB</td>
<td>Absence of spontaneous speech development</td>
</tr>
</tbody>
</table>
9.2 Detection and Investigation of Pediatric Hearing Disorders

Hearing disorders in newborns, infants, and small children are detected with audiometric methods that are adapted to the special difficulties inherent in detecting and quantifying a pediatric hearing problem. Screening tests are best for detecting hearing impairment in newborns. Any suspicion of a pediatric hearing disorder warrants an immediate audiologic work-up that includes objective tests and behavioral audiometric methods that are appropriate for the age of the child.

Screening

Because hearing impairment is not an obvious condition in newborns, it must be detected by screening. If the baby does not pass the screening, a hearing loss should be confirmed and an audiologic evaluation performed during subsequent weeks. Risk screening: This is a selective program that screens only children with features that are known to be associated with an increased risk of hearing loss (see below and 9.1, p.198). Screening is indicated for newborns who:
- Have been in an intensive-care unit for more than 48 hours
- Have a positive family history of hearing impairment; or
- Manifest craniofacial anomalies.

The overall prevalence of a permanent, moderate to severe hearing loss in these three groups is 4%. Consequently, the hearing of all newborns in these groups should be routinely tested (methods: see below). This risk-based screening will detect approximately half of all hearing problems in infants. Universal screening: The universal screening of all newborns can detect up to 80% of all hearing problems and is therefore more effective than risk screening. The best time for screening is 2–3 days after birth (e.g., during the second routine examination), since most babies are easily available for hearing screening at that time, depending on the health-care system. Screening may have to be organized differently in health-care systems in which babies may be available only during the first 24 hours after birth. Additional screening examinations are performed when the child is older—at routine pediatric visits, for example (distraction test, see below), or when the child enters school.

Screening Procedure

Newborn hearing screening is based on objective audiometric test methods that are specially adapted for screening requirements. Useful tests are otoacoustic emissions (OAEs, see pp.188–191) and the auditory brainstem response (early auditory evoked potentials, see pp.186–188). Two-stage screening is performed in most cases, i.e., if a response is not elicited, the test is repeated 1–2 days later or is supplemented by an additional test (e.g., TEOAEs, see p.189, followed by the ABR). Children that do not pass the screening should be referred for a differentiated audiologic evaluation (see below). Approximately one in 10 of these babies has a permanent hearing loss. Screening in older children employs the above methods in addition to behavioral tests such as the distraction test or, in preschool and school-age children, pure-tone audiometry.

Diagnostic Methods in Pediatric Audiology

Objective Methods

The methods of objective audiometry can be used at any age, as long as passive cooperation of the child can be achieved (by sedation if necessary). Hearing loss can be detected, and a site-of-lesion determination can be made. Otoacoustic emissions:
Indication: OAEs provide a rapid method for assessing the function of the cochlear amplifier (see p.162). OAEs cannot be used to determine the degree of hearing loss.
Interpretation: If OAEs are present, it may be assumed that peripheral hearing is satisfactory.

The presence of OAEs does not exclude a hearing disorder. Very rarely, a neural or central hearing disorder may be present.

Auditory evoked potentials:
Indication: Auditory evoked potentials (AEPs) are the most important objective method for the investigation of hearing loss in infants and children. The auditory brainstem response (ABR, see p.187) is most commonly tested. It can be used to determine the hearing threshold, which is done more accurately at frequencies above 1 kHz than at lower frequencies.
Interpretation: If an ABR is not elicited, it should be concluded that severe hearing loss is present.
The latency of the ABR is significantly longer in newborns and infants than in adults. By 2–3 years of age the latency values are approximately equal to those in adults. The maturation process of central auditory structures is reflected in this gradual shortening of ABR latencies.

Imittance measurements: Imittance measurements play a minor role in newborns and infants because it cannot be done accurately using ordinary equipment. It is commonly used in small children, however, for the documentation and follow-up of middle ear effusions with associated conductive hearing loss (see Otitis Media with Effusion, p. 240).

**Behavioral Audiometric Methods**

*Indication:* Behavioral methods are important for testing subjective auditory responses in pediatric audiology. They are also necessary in rehabilitative measures such as hearing-aid fitting.

*Prerequisites:* Behavioral audiometric tests can be performed at virtually any age. The methodology must be age-appropriate, however, and the reliability of the tests is variable.

The examiner must have experience and patience in conducting and interpreting the examination. It is necessary to gain the attention of the child, hold it for as long as possible by adapting the test situation, and correctly recognize and interpret the responses. The examiner must gather as much relevant information as possible within a short time.

**Reflex audiometry:** Nonspecific responses to auditory stimuli can be elicited in normal infants from birth on. Sucking responses, motor responses such as the Moro reflex, and changes in respiratory pattern can be elicited. Generally, these reflexes require loud auditory stimuli of approximately 80 dB.

**Response audiometry:** By about 5 months of age, acoustic stimuli evoke typical response patterns in normal infants that can be used to test hearing. For example, the infant will turn its head toward a sound source that is outside the visual field. This response is initially present in the horizontal plane and later occurs in the vertical plane as well. This response is contingent upon normal maturation of directional hearing.

- **Distraction test:** This test is administered by two testers in a standardized, quiet environment using various standard acoustic stimuli. The setup is shown in Fig. 9.1.
- **Visual reinforcement audiometry (VRA):** This test again utilizes the head-turn response, aided by positive reinforcement, to obtain audiometric threshold measurements. An acoustic stimulus is combined with the activation of a moving toy, such as a dancing bear. After conditioning, the child will turn the head toward the toy when hearing the acoustic stimulus.

*Play audiometry:* This is a variant of pure-tone audiometry (see 8.3, pp. 178–183). By 1–2 years of age, it is possible to incorporate the tasks and responses of pure-tone audiometry into a play setting. For example, the child may stack one building block onto another as soon as a certain acoustic stimulus is withdrawn. The play situation, the tasks, and the presentation of the acoustic stimulus vary with the age of the child and the clinical problem.

*Indication:* These techniques are used for the detection of hearing impairment and for the fitting of hearing aids and cochlear implants.

**Pediatric speech audiometry:** The easiest way to test speech discrimination is by verbally instructing the child to select a certain toy or picture that is on a table along with a number of other toys or pictures. This technique can be used to screen for speech recognition problems by telling the child, at increasing distances, to point to objects displayed in pictures. This kind of test can be administered by 2 years of age.

By about 3 to 4 years of age, the examiner can use audiometric speech tests that have been specially designed for children—e.g., the Pediatric Speech Intelligibility (PSI) test.

By the time the child enters school, basically the same speech audiometric methods can be used as in adults, with minor modifications.
9.3 Treatment of Pediatric Hearing Disorders

The medical treatment of hearing disorders in children is basically the same as in adults. It is merely adapted to the special circumstances in the pediatric age group. Hearing ability is improved by means of hearing aids or cochlear implants whenever possible. To attain satisfactory speech proficiency in small children with permanent hearing loss, however, additional rehabilitative and training measures are necessary in the form of special education services, which should be instituted as early as possible.

**Indications:** There is a general consensus that children with a bilateral, moderate, permanent hearing loss should be treated in order to prevent significant impairment of speech and language development. More controversial are the treatment measures that are appropriate in children with a mild or unilateral hearing loss.

**Basic options:** Conductive hearing loss can often be improved by surgery. For example, malformed elements of the ossicular chain can be reconstructed, or middle ear ventilation can be surgically established in small children with recurrent effusions (see 11.2, p. 234, and 11.3, p. 238). Medical treatment is rarely an option for hearing disorders.

In other cases, every effort should be made to utilize residual hearing as fully as possible while also providing appropriate services to stimulate and support the child’s auditory development. Hearing aids should be prescribed for moderate to severe hearing loss, while cochlear implants are indicated for deafness or profound hearing loss that does not benefit from hearing aids. Other support services are also instituted with the goal of promoting speech and language development. These rehabilitative measures are different from communicative training in adults due to the absence or impairment of speech development in children.

**Auditory Devices**

See also 8.5, Rehabilitation and Hearing Aids, pp. 192–195.

**Hearing Aids**

**Types of aid:** Children are generally fitted with two behind-the-ear hearing aids. This can be done in infants only a few months old. A special case is bilateral aural atresia, in which bone conduction hearing aids are fitted shortly after birth.

The fitting itself is considerably more difficult in children and requires a great deal of experience. Infants and small children are very limited in their ability to cooperate, and these patients lack auditory experience. Children with a hearing aid hear a spectrum of sounds that is unfamiliar to them. With greater degrees of hearing loss, this problem becomes more serious and more time is needed to achieve a proper fitting. The task in hearing-aid fitting is to work with the child to reach both the possibilities and limitations of acoustic amplification.

**Follow-up:** After the hearing aids have been fitted, their function must be regularly tested by parents and educators because initially the child is unable to detect and report malfunctions. But if the hearing aids are of definite benefit, the child will soon ask for them himself and later will make it known if the devices are not functioning properly.

**Other Assistive Devices**

*FM transmitters* are used frequently and successfully in classrooms and often can be coupled to the hearing aid. Placing the microphone close to the teacher allows the teacher’s voice to reach the student with significantly less interference from background noise. Another, rarely used option is the use of vibrators, which can be worn on the wrist and convert the acoustic signal into a synchronous vibratory signal.

**Cochlear Implants**

**Indications and advantages:** Cochlear implants are used in patients who have cochlear damage and an auditory nerve that is responsive to stimulation. They are mainly an option for two categories of hearing-impaired children (see also p. 194):

**Congenital deafness or profound hearing loss:** The earlier cochlear implantation is performed in these children, the better the device can exploit the natural adaptability and receptivity of the central auditory structures. Most children treated early with a cochlear implant learn to understand speech. Many can even understand speech without lip reading (open speech comprehension, see p. 194). It is equally important to train the child’s speaking ability. Cochlear implantation after puberty very rarely provides open speech comprehension or a significant improvement of speech.

**Bilateral acquired deafness** (*e.g.*, after meningitis): Cochlear implantation should be performed as soon as possible so that existing auditory development is not lost and can be exploited with the cochlear im-
plant during rehabilitation. In practice, this means that cochlear implantation should be scheduled as soon as the child is diagnosed with bilateral deafness or profound hearing loss that is not treatable with hearing aids. As a rule, hearing aids should be tried for several months before this determination is made.

**Surgery:** Cochlear implantation surgery is a low-risk procedure even in children. It is very rarely contraindicated for medical reasons.

**Speech processor fitting:** The speech processor is fitted to the child on an individual basis in sessions that are consistent with pedagogic and audiologic principles. Like hearing-aid fitting, it is a process that requires a great deal of patience, empathy, and experience.

**Education Services**

Whenever a child is diagnosed has having a significant, permanent hearing loss, the child should receive special education services from a trained, licensed teacher. At their simplest level, these services will help children learn how to use and care for their auditory devices. Education and counseling should also be provided to parents and other family members on dealing with issues of deafness within the family. For small children, these services are generally provided in the home. Auditory perception is trained and practiced in a play setting, while parents are taught behavioral rules and assistive measures. The goal of early education services is to integrate the hearing-impaired child into public schooling and thus into the society of normal-hearing persons. Specialized kindergartens and schools are available for children who cannot be mainstreamed. Some institutions specialize in a total communication approach that combines oral communication with other modalities such as sign language.

**Rehabilitative Measures for Unilateral Hearing Loss**

Audiologic and education services in children with unilateral hearing loss are generally limited to audiometric surveillance of the healthy side and to counseling. Favorable seating in the classroom is essential. The child should sit as close to the teacher as possible and listed with the good ear turned toward the instructor. It is also important to instruct the teacher in maintaining correct speech habits. The use of an FM system may be necessary in some cases.

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**9.2 Case report: cochlear implantation after meningitis**

A girl was diagnosed with acute meningitis at 19 months of age. *Streptococcus pneumoniae* was detected in the cerebrospinal fluid (CSF). Antibiotic treatment was provided, and the child made a swift recovery.

Up to this point, the child had developed normally. After the acute meningitis, however, the parents noticed a change in behavior. The girl no longer understood speech, and she ceased the “babbling” that had been so common before. ABR testing was performed and indicated a severe, bilateral hearing loss that was worse on the left side than the right.

The child was referred for education services and fitted with hearing aids, but these measures were of little benefit. The child was obviously suffering from her hearing loss, leading at times to behavioral disturbances such as self-injury.

Five months after the meningitis, the child was evaluated for cochlear implantation. By that time no auditory responses could be elicited at all, even when the hearing aids were used. High-resolution computed tomography scans revealed significant intracochlear calcifications on the right side (Fig.). Mild calcifications were also visible on the left side. It was recommended that the child undergo cochlear implantation as soon as possible to forestall further progression of the intracochlear calcification.

The surgery was performed 3 months later, when the girl was 2 years, 1 month of age. Calcifications were found in lumen of the basal cochlea, making it necessary to insert a somewhat shorter-length electrode array.

Subsequent fitting of the speech processor went well, and the girl made rapid progress. Her behavior became more like that before the meningitis, and the parents said that their daughter was “more like her old self” again. The girl asks for her device first thing in the morning and wears it throughout the day.
The External Ear

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10.1 Special Anatomy and Examination of the External Ear

It is necessary to know the special anatomy of the external ear in order to understand the specific manifestations of diseases of the auricle and external auditory canal. Because this anatomy is closely linked to the examination technique, this also deals with the general technique of otoscopy and evaluation of the ear canal. Otoscopic evaluation of the tympanic membrane is covered in Chapter 11 (pp. 228–253).

Anatomy

Auricle

The auricle and the external auditory canal (external acoustic meatus, ear canal) form an anatomical and functional unit. The lateral cartilaginous structures of the external ear are continuous with the medial bony structures. The **medial boundary** of the external ear is the tympanic membrane. The **shape of the auricle** is defined by the elastic cartilaginous plate (Fig. 10.1). The perichondrium of the elastic cartilage forms a unit with the lateral dermis of the auricle; there is no actual subcutaneous tissue in this region. As a result, cutaneous changes and swelling frequently affect the perichondrium and cartilage, which can lead to severe pain, poor absorption, and cartilage destruction with permanent changes in auricular shape.

External Auditory Canal

The auricle and the **cartilaginous portion** of the external auditory canal form a unit both anatomically and in many pathological respects (Fig. 10.2). Diseases of the auricle often spread to the ear canal, and vice versa. The lateral two-thirds of the external auditory canal consists of a fibrocartilaginous framework that is angled downward and forward relative to the bony medial third. This is why, at otoscopy, the mobile cartilaginous part of the ear is pulled upward and backward. The **bony portion** of the ear canal is formed by the tympanic part of the temporal bone. The skin in that region of the ear canal is very thin and directly overlies the peristomeum; this accounts for the temperature and pain sensitivity of the medial canal skin and can influence the pathogenesis and course of ear canal diseases such as necrotizing otitis externa (see p. 221). The bony ear canal grows considerably less than the cartilaginous ear canal. The cartilaginous part is shorter than the bony part in infants, but both parts are of approximately equal length by 5–6 years of age. The ear canal in adults is approximately 2.5 cm long.

Innervation

Most of the external ear receives its sensory supply from branches of the great auricular nerve (from the cervical plexus) and auriculotemporal nerve (from the third division of the trigeminal). Portions of the concha and ear canal are supplied by the auricular branch of the vagus nerve, whose stimulation by otoscopy can induce coughing (vagal irritation). In some
cases the ear canal and concha are also supplied by the 
auricular branches of the facial nerve (somatosensory 
portion).

**Anatomical Relations**

The external ear and particularly the ear canal border 
on the following structures, which can lead to changes in the ear canal:
- Anterior to the cartilaginous and bony ear canal is the *temporomandibular joint*. Trauma to this joint can lead to swelling and bleeding of the ear canal.
- The *parotid gland* borders the cartilaginous ear canal anteriorly and inferiorly. Inflammations and tumors can spread through Santorini’s fissures from the ear canal to the parotid gland or vice versa.
- Posterior to the ear canal is the *mastoid*. The post-
erior bony canal wall forms part of the anterior wall of the mastoid, and the medial part of the canal wall adjoins the tympanic cavity. Inflammation of the mastoid (mastoiditis) can lead to sagging of the posterosuperior canal wall.
- The external ear is bordered superiorly by the tem-
poralis muscle and the squamous part of the tempo-
ral bone. The superomedial portions of the bony ca-
nal wall form the floor of the epitympanum (attic) in the middle ear.

**Examination**

**History and inspection**: The patient should be ques-
tioned specifically about otalgia, aural fullness, or hearing loss (currently or in the past), and particular attention should be given to aural discharge (otor-
rhea). The auricle is a relatively common site of injury due to its exposed location. Generally, these injuries are visible due to the absence of deeper subcutaneous structures. Examination of the ear, then, always begins with a careful inspection of the auricle, its surround-
ings, and the opening of the ear canal.

**Palpation**: Before the ear is examined further and be-
fore otoscopy, the tragus should be palpated. Tragal tenderness signifies an inflammation of the cartilagi-
 nous portion of the ear canal, because displacement of the rigid tragus moves and irritates the perichondri-
um attached to the rest of the meatal cartilage. The same effect can be achieved by pulling gently on the 
auricle.

Otoscopy should be performed with particular care 
when tragal tenderness or otalgia is present.

**Otoscopy**: Examination with a hand-held otoscope or 
aural speculum allows inspection of the external ear 
canal and tympanic membrane. The technique of otos-
copy and the findings in the ear canal are described in 
this chapter and on p. 166. Tympanic membrane find-
ings are described in Chapter 8, pp. 166–191, and 
Chapter 11, pp. 228–233.

When otoscopy is performed with an aural speculum 
shaped so that it normally cannot reach the tympanic membrane, the danger of tympanic membrane injury is very slight. A relatively broad speculum affords good 
exposure of the ear canal and tympanic membrane and provides better illumination. Narrow specula 
may be advanced unnecessarily far into the ear canal.

Touching the medial bony portions of the ear canal 
with the tip of the otoscope is not only painful but 
can easily injure the delicate skin of the canal wall. 

The auricle is gently rotated backward and upward to 
align the cartilaginous ear canal with the bony canal 
for examination (Fig. 10.3). Once the otoscope has 
been introduced under vision and stably aligned with 
the bony canal, the auricle can be released. 

Proceeding laterally to medially, the examiner inspects the course and appearance of the external canal 
as far as the tympanic membrane. Contaminants in 
the form of cerumen, debris, drainage, or foreign bod-
ies are removed by suction with a small instrument or 
by irrigation of the ear canal (see p. 212).

The lateral portions of the ear canal can also be examined with a rhinoscopy-like speculum. Whenever possible, manipulations in the deeper portions of the ear 
canal should be done under stereoscopic vision using 
an otomicroscope.
10.2 Noninflammatory Diseases and Injuries to the External Ear

A very common noninflammatory condition is obstruction of the ear canal by an accumulation of cerumen. Cleansing the ear canal is a common medical service that must be done carefully due to the proximity of middle ear structures and the sensitivity of the ear canal skin. Injuries, foreign bodies, and stenoses are other important medical findings in the external ear.

Deformities and Malformations

Prominent Ears

Prominent or protruding ears are a normal congenital variant that has no functional consequences. The normal angle between the auricle and head is approximately 20–30°. This angle may be increased due to a deep concha or lack of development of the antihelix.

Treatment: Surgical correction consists of reducing the size of the concha and reconstructing the antihelix. In children, the surgery is usually performed at preschool age under general anesthesia. An ear dressing is worn for about 2 weeks after the operation.

Hyperostoses and Exostoses of the Ear Canal

Exostoses of the external ear canal are true osteomas that are most commonly located near the annulus on the superomedial canal wall. They develop from ossification centers and appear otoscopically as pale, rounded bony prominences. Hyperostosis is an appositional growth of the bony ear canal, usually induced by periotic irritation such as frequent contact with cold water (synonym: swimmer’s ear).

Diagnosis: The otoscopic appearance is characteristic (Fig. 10.4). The ear canal is usually narrowed in its bony portion by several smooth, pale prominences. These may form a stenosis that covers the tympanic membrane. The narrowed site is hard and tender when probed with a small, blunt hook.

Complications: A stenosis causing retention of squamous debris and water can lead to recurrent otitis externa. In the absence of inflammation, only high-grade stenoses cause conductive hearing loss.

Treatment: If complications arise, the growths should be surgically removed.

Cerumen and Cerumen Impaction

Physiology: Cerumen (“earwax”) is produced by the cerumen and sebaceous glands in the skin of the ear canal. It forms a protective film in which fatty acids, lysozymes, and the creation of an acid milieu effectively protect the skin of the ear canal. Self-cleansing of the ear canal, with natural removal of accumulated cerumen, is normally accomplished by epithelial migration from the tympanic membrane toward the external meatus.

Pathophysiology: Cerumen impaction may result from a disturbance of the normal self-cleansing mechanism or from excessive cerumen secretion. The cerumen plug consists mainly of secretions from the cerumen glands mixed with sebum, exfoliative debris, and contaminants. Imprudent cleaning of the ear canal (especially with cotton-tipped swabs!) can interfere with the self-cleansing mechanism and displace the cerumen toward the tympanic membrane. Obstruction of the ear canal by cerumen may be caused by the impaction or swelling of a cerumen
10.1 Malformations of the external ear

A continuum exists between a normal variant of auricular shape and an anomaly (see Prominent Ears). The auricle is formed embryologically from six mesenchymal hillocks and migrates in a cranial direction with further development. Low-set ears may therefore signify an anomaly of the external auditory canal or middle ear.

Auricular appendages

These appendages, composed of skin and cartilage, develop from aberrant embryonic cell rests that become trapped in the area of the first branchial cleft. They are usually preauricular and have no functional importance. Rarely, they are associated with other ear malformations or hearing impairment. Treatment consists of excision for appendages that are cosmetically objectionable.

Congenital auricular fistulas and cysts

Aural fistulas, preauricular fistulas and cysts develop from sites of epithelial retention in the area of the first branchial cleft. The fistulous tracts usually terminate blindly. The fistulous openings are most commonly found at preauricular sites on the helical rim. A continuum exists with high cervical fistulas located at an infra- or retroauricular site. Possible complications include infections. In these cases the fistula must be completely excised, based on a knowledge of the possible course of the fistulous tracts and their relations to the facial nerve and external ear canal.

Malformations of the Auricle

Auricular dysplasia is classified into three grades:

Grade I dysplasia (minor anomalies): The structural subunits of the auricle are present but malformed. Prominent ears are an example (see p. 270). Other anomalies are microtia, tubercles (Darwinian tubercles), helical projections (auricular apex), and maccus ear with partial absence of the helix. These anomalies either require no correction or a relatively simple corrective procedure.

Grade II dysplasia (mild microtia): The auricle is small, severely misshapen, and lacks some subunits. An anomaly of the external meatus is often present. Microtia requires meticulous corrective surgery (see 10.2, p. 215).

Grade III dysplasia (microtia and anotia): Normal auricular structures are absent, and the ear canal is almost always atretic (Fig.). Objective audiologic tests should be performed shortly after birth to determine whether acceptable auditory function is present or at least one side. Children with unilateral atresia should receive a bone-conduction hearing aid during the first months of life. Surgical reconstruction of the ear canal and middle ear can be performed by about 10 years of age. Auricular reconstruction is usually done later. Methods of surgical reconstruction are described in 10.2 on p. 215.

Stenosis and atresia of the ear canal

Stenoses and atresias of the external ear canal are frequently but not always associated with auricular anomalies. Unilateral atresia is generally not treated surgically, as this would be unlikely to significantly improve the quality of life.

plug. This often occurs after contact with water. With ageing, drying of the meatal skin and changes in secretions can lead to the formation of a hard cerumen that tends to be retained in the ear, especially with a narrow canal.

Symptoms: Cerumen impaction causes a pressure sensation in the ear with concomitant hearing loss. Some patients complain of vertigo or tinnitus.

Diagnosis: With a cerumen impaction, otoscopy may show obstruction of the ear canal by a yellowish-brown to black material. The consistency of the cerumen is variable. A detailed otologic history should be obtained.

Particular attention is given to tympanic membrane perforations and previous temporal bone fractures or otologic surgery.

Differential diagnosis: An epithelial plug or crust can result from a cholesteatoma in the external ear canal. Occasionally the ear canal is obstructed by a thin skin flap, or cuticle. Tumors, foreign bodies, and crusted blood should also be excluded.

Complications: Otitis externa may develop, but generally complications are very rare.

Treatment: Cerumen and cerumen plugs are removed with a small instrument (hook, curette) or by aural irrigation.

Instrumental cleaning of the ear canal is best done under stereoscopic vision by a specialist using an otomicroscope.
Technique of aural irrigation:

Irrigation is contraindicated in patients with a positive otologic history (see above).

- Hard cerumen can be softened by pretreatment with hydrogen peroxide, a glycerin-containing agent, or other detergents for several days.
- The ear is irrigated with bacteriologically pure water at 37 °C using an ear syringe with a blunt cannula.
- The water jet is directed posterosuperiorly; it is not trained directly on the tympanic membrane (Fig. 10.5).
- Irrigation should be followed by otoscopy and a clinical hearing test (tuning fork test).

Contraindications (referral to a specialist):
- Positive otologic history (see above)
- Single hearing ear affected
- Restless, uncooperative patient
- Foreign body

Prophylaxis: The best preventive measure is to avoid improper cleaning of the ear canal, particularly the regular use of cotton-tipped swabs.

Foreign Bodies in the Ear Canal

Causes: Most cases occur when small children insert small play objects such as beads or Lego pieces into the ear canal. Most foreign bodies in adults consist of noise-reducing ear plugs or objects used for manipulations in the ear canal. Insects may also become trapped in the ear canal.

Diagnosis: The history is usually diagnostic, and generally the foreign body is easily identified at otoscopy. Difficulties can result from secondary injuries, swelling, or inflammation.

Check for signs of associated injury to middle- or inner-ear structures (e.g., tympanic membrane perforation, otitis media, facial nerve lesions, vertigo, nystagmus, or sensorineural hearing loss.

Differential diagnosis: The differential diagnosis includes cerumen impaction, dried blood, tumors of the ear canal, cholesteatoma, and otitis externa (e.g., due to fungal infection).

Complications: A deeply penetrating foreign body can cause middle- and inner-ear damage. The prolonged retention of a foreign body in the ear generally leads to secondary otitis externa, often with a fetid discharge.

Treatment: The foreign body should be carefully removed with a small extraction hook. Care is taken not to push the foreign body deeper into the ear canal or through the tympanic membrane (Fig. 10.6).

Aural irrigation should not be used on foreign bodies in the ear canal.

In children, it is often preferable to extract the foreign body under general anesthesia rather than try hazardous manipulations. Insects can be killed with a 10% lidocaine solution. Very rarely, a surgical incision may be needed to remove foreign bodies lodged tightly in the ear canal.

Injuries and Physical Damage

Auricular Hematoma, Auricular Seroma

Definition: An auricular hematoma or seroma is a collection of blood or serous fluid between the perichondrium and auricular cartilage.

Pathogenesis: Blunt trauma like that commonly occurring in contact and combative sports (wrestling, boxing, water polo) causes the skin and attached perichondrium to separate from the auricular cartilage. If the injury remains closed, a hematoma or seroma may form between these layers. There is very little tendency for the fluid collection to be reabsorbed.

Symptoms: The trauma itself is painful, but typically there is no pain afterward.

Diagnosis: The findings on inspection and palpation are unequivocal (Fig. 10.7). The skin over the lateral auricular cartilage shows swelling and fluctuation.
The examiner should exclude associated injuries to the temporal bone, ear canal, middle ear, and temporomandibular joint and secondary infection of the hematoma.

**Differential diagnosis:** Recurrent polychondritis can give rise to a spontaneous seroma.

**Complications:** A secondary infection, often caused by needle aspiration of the hematoma, can lead to perichondritis. Poor reabsorption of the fluid collection can result in permanent deformity of the cartilaginous framework with the development of an irreversible “cauliflower ear.”

**Treatment:** The hematoma or seroma is surgically evacuated and the perichondrium is reattached to the cartilage. A contoured dressing (e.g., of oil-impregnated cotton) is then applied. Seromas may recur even after surgical treatment.

**Prophylaxis:** Ear protection should be worn for all contact sports.

### Sharp Auricular Injury and Auricular Avulsion

**Definition:** In an open auricular injury, the cartilage is exposed over an area of variable size. An auricular avulsion may be partial (with an intact bridge of skin) or complete (part of the auricle is completely detached).

**Diagnosis:** The injured site is carefully cleaned, and the extent of the injury is determined.

Intact bridges of skin and cartilage should always be left intact.

Associated injuries to the temporal bone, ear canal, middle ear, or temporomandibular joint should be excluded.

**Complications:** A soft-tissue infection and perichondritis may develop secondarily. Crushed or severed parts may succumb to necrosis.

**Treatment:** Primary measures for an auricular avulsion:
- Cover the wound with a sterile dressing.
- Refer the patient to an ear, nose, and throat (ENT) facility right away.
- Send severed ear parts with the patient.
- Cool the part whenever possible, but do not pack it directly in ice (risk of tissue damage!). The ideal solution is to wrap the part in moist gauze, place it in a waterproof plastic bag, and immerse the bag in ice water.

The wound is closed with perichondrial and cutaneous sutures. Exposed cartilage should be covered with
skin, using flap advancement if necessary (see 3.4, p.36). The lateral skin is reapproximated with a contoured ear dressing. With a partial auricular avulsion, the specialist should attempt a reanastomosis if an adequate skin bridge is present, the avulsed part is not too large and is not badly crushed, and the injury is no more than about 6 hours old. As an alternative and for most complete avulsions, the cartilage can be implanted subcutaneously at a retroauricular or cervical site and used 6 months later to perform a secondary reconstruction.

**Prognosis:** The prognosis depends on the extent of the injury. Because the tissue has a good blood supply, parts still attached by a small bridge of skin have a good chance of healing, and even some small detached parts may become revascularized. The prognosis for healing tends to be poor with a complete avulsion, however.

**Burns and Frostbite Injuries**

**Definition:** Thermal injuries to the auricle are graded as follows:

- Damage confined to the skin:
  - Grade I: localized erythema
  - Grade II: blistering of the skin
- Damage involves the entire skin-cartilage unit:
  - Grade III: deep tissue necrosis

**Etiology:** The auricle is a frequent site of frostbite due to its exposed location and inadequate protection from the cold. Burns confined to the auricle are somewhat rare.

**Diagnosis:** A burn is easily diagnosed from the recent history. Frostbite may appear initially as a white skin discoloration that is demarcated from its surroundings. The patient typically does not notice the initial cold injury, which does not become painful until the frostbitten area is rewarmed.

Associated injuries should be excluded, especially with burns. The examiner should check for concomitant involvement of the ear canal and tympanic membrane.

**Differential diagnosis:** Caustic chemical injuries and electrical injuries can produce similar skin changes.

**Complications:** A deep thermal injury may give rise to cartilage necrosis and permanent deformity. Chilblains may develop on the helical rim with ulcerations and itching. Perichondritis can also occur.

**Treatment:** Treatment follows the general principles of surgical wound care for burns and frostbite.

Local treatments (dressings) should not exert pressure on the auricle to avoid further compromise of the auricular blood supply.

**Burns:** Consistent with the general principles of burn care, superficial burns in particular should be cooled immediately and treated with other local anti-inflammatory measures. Surgical debridement may be necessary later for more severe burns.

**Frostbite:** The frostbitten area should be gently warmed (e.g., with a heat lamp), taking care not to burn the tissue. The best strategy for injuries with bulla formation or necrosis is to provide dry treatment and await demarcation of the frostbitten area. Circulatory stimulants such as dextran or pentoxifylline can be used.

**Reconstructive surgery** is deferred until the site has completely healed, which usually takes about 6 months.

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**Fig. 10.8 Partial avulsion of the auricle**

a Auricular injury on the right side with partial avulsion in a 9-year-old girl bitten by a dog.  
b Appearance after surgical treatment and healing.
**Prognosis:** More severe burns or frostbite injuries are often associated with permanent auricular deformity.

**Injuries to the External Auditory Canal**

**Etiology:** Isolated injuries to the external ear canal are caused mainly by foreign bodies or harmful manipulations.

**Diagnosis:** The history usually indicates previous trauma to the ear. The meatal skin is tender, and there is bleeding from the ear canal. Otoscopy may reveal an epithelial injury, bleeding, a hemorrhagic bulla, or crusted blood. Associated injuries to the tympanic membrane, middle ear, temporomandibular joint, and skull base should be excluded.

**Complications:** Ear canal injuries may be complicated by secondary infection. Later, during healing, there may be cyst formation or stenosis of the ear canal due to scarring.

**Treatment:** Detached epithelium should be reapproximated whenever possible. For bleeding, it may be necessary to pack the ear canal with Gelfoam or synthetic sponge.

**Prognosis:** Isolated injuries to the ear canal are usually uncomplicated and show a good healing tendency.

### 10.2 Reconstructive surgery of the auricle

Surgical reconstruction of the auricle may be necessary after trauma, for malformations, or after a tumor resection. The extent and complexity of auricular reconstructive surgery depend on the underlying condition and on what the reconstruction can and must achieve. Total auricular reconstruction is among the most difficult and demanding procedures in plastic and reconstructive surgery.

Auricular reconstructions fall into several categories:

**Shape corrections:** These are relatively simple procedures that are used in the treatment of prominent ears, for example (see p. 210).

**Partial reconstructions:** These may be indicated following a partial resection or for grade II dysplasia. The auricle is reconstructed using available local cartilage whenever possible. The necessary skin coverage is obtained with advancement flaps (see p. 36) or preliminary subcutaneous expansion.

**Total reconstruction:** This is mainly necessary for congenital aplasias and severe anomalies (grade III dysplasia). A total reconstruction is very challenging and complex and is usually staged in several sittings. Missing cartilage is reconstructed with autologous cartilage grafts (costal cartilage) or replaced with synthetic prostheses.

**Epithesis:** An external prosthesis may be considered as an alternative to auricular reconstruction. The epithesis may be attached to implanted titanium screws or an eyeglass frame, or it may be glued directly to the skin.
10.3 Overview: Differential Diagnosis of Inflammatory Changes in the External Ear

Inflammatory changes in the external ear are common and are often treated initially by a primary-care physician. It is important, therefore, to be familiar with the various types of external ear inflammation, their hazards, and their appropriate management. An inflammatory condition of the auricle or external auditory canal is diagnosed clinically; additional tests are rarely needed. Different inflammatory conditions of the external ear are interrelated in their pathogenesis, sometimes making it difficult to differentiate the various forms. Specific diseases are discussed in the next (p. 218). The present unit deals with issues of differential diagnosis, reviewing pathophysiologic relationships and associated problems of differential diagnosis that must be considered in otitis externa.

General

Inflammations of the external ear may manifest acutely with severe pain, subacutely, or may present with chronic complaints such as itching and scaly skin. Acute inflammations are often caused by bacterial infection, while chronic forms more closely resemble eczema.

The term otitis externa usually refers to inflammation of the external auditory canal. This cannot always be clearly differentiated from auricular inflammation, however, since inflammations of the external canal may spread to the auricle, and vice versa. Nevertheless, typical forms of auricular inflammation are distinguishable from inflammatory conditions of the ear canal.

It is not always easy to distinguish among the various inflammatory conditions of the external ear, which are interrelated in their pathogenesis. For example, chronic eczema may give rise to an acute inflammation of the ear canal, or a fungal infection of the ear canal may develop from a subacute bacterial otitis externa. Moreover, otitis media (see 11.3, pp. 238–249) may incite a concomitant inflammation of the external ear.

A profuse, mucopurulent aural discharge often originates in the middle ear and not in the external ear canal.

Pathogenesis

Inflammations of the external ear are often caused by factors that interfere with the normal defenses against infection. The normal cerumen film (acid pH, antibacterial fatty acid content) and the physiologic lateral migration of the epithelium lining the ear canal create an effective barrier to infection. Any of the following factors may disturb the self-cleansing of the ear canal and its protective mechanisms, predisposing to otitis externa:

- Exogenous factors: maceration of the skin by water, the creation of a warm moist chamber, pH changes caused by soaps and shampoos, manipulations with cotton-tipped swabs, insert earphones, or ear plugs.
- Endogenous factors: proneness to eczema, allergies, metabolic disorders such as diabetes mellitus.
- Local changes: exostoses, stenoses, anatomical variants.

It is common for several of these factors to be combined. Treatment and prophylaxis are geared toward preventing or correcting the causal factors.

Differential Diagnosis of Acute Inflammatory Change

Acute inflammations of the auricle or external ear canal are manifested clinically by severe pain. Obstruction of the ear canal by drainage or skin swelling can lead to conductive hearing loss. Generally, there is tenderness on movement of the auricle or tragus. Initial external inspection will show whether the inflammation involves the auricle. If this is the case, a pure perichondritis can be distinguished from a spreading soft-tissue infection (e.g., cellulitis) by its confinement to the cartilage. With an acute inflammation of the ear canal, careful otoscopy can usually differentiate a furuncle of the meatal hairs or glands from acute dermatitis.

If the ear canal is filled with purulent discharge, it can be difficult to distinguish between otitis externa and suppurative otitis media with a perforated tympanic membrane.

A Valsalva maneuver is helpful in diagnosing suppurative otitis media with a perforated tympanic membrane, as it may cause air bubbles to appear in the discharge.
The typical history of *bullous otitis externa* consists of severe otalgia with a bloody discharge. The diagnosis can often be established simply by the otoscopic detection of hemorrhagic bullae on the bony canal wall. *Herpes zoster oticus* manifests with small bullae on the auricle and also in the ear canal.

**Differential Diagnosis of Chronic Inflammatory Change**

The dominant symptom of chronic otitis externa is usually itching, not pain. Inspection will generally reveal redness and scaling or crusting about the meatal orifice.

It is important first to assess the condition of the tympanic membrane and middle ear. A mobile tympanic membrane (Valsalva maneuver, pneumatic otoscopy, see p. 231; tympanography, p. 185) or the absence of conductive hearing loss (positive Rinne test, see p. 168) indicates that the tympanic membrane and middle ear are intact. The skin of the ear canal shows typical changes that may consist of eczema, ulceration, or granulations. Purulent drainage in the ear canal can hamper visual inspection and may have to be carefully removed (e.g., with a fine suction tip). When *chronic otitis media* is present, the chronic inflammatory changes in the external canal are generally caused by drainage from the middle ear. In these cases, too, the drainage in the ear canal often hampers an accurate evaluation and should be carefully removed. Conductive hearing loss is present. The presence of mucosal polyps in the ear canal or of firmly adherent crusts in the superior part of the bony canal often signifies an inflammatory middle ear cholesteatoma (see pp. 243–249).
10.4 Inflammatory Diseases of the External Ear

As described in the preceding, it is common to encounter mixed inflammatory conditions of the external ear. Typical clinical manifestations can be distinguished, however. It is helpful, therefore, to consider inflammations of the auricle and ear canal separately, for at least the onset of the inflammation often presents a typical form. The various forms are described separately in this unit.

Auricle

Eczema and Dermatitis of the Auricle

Definition: an inflammatory condition of the auricle confined to the dermis. The cartilage and perichondrium are not involved.

Pathogenesis: The pathogenesis is the same as in other skin regions and is based on immune/allergic, toxic or physical causes. Eczema and dermatitis of the auricle are frequently caused by:
- Jewelry items (ear rings)
- Soaps and cosmetics (shampoo, hair spray)
- Listening aids (hearing aids, earmolds, insert earphones)
- Thermal injury (sun exposure, radiotherapy, frostbite)

Symptoms: The cardinal symptoms are itching and occasional burning with little pain. The skin is erythematous and may be dry and scaly or moist and weeping, depending on the stage and severity of the eczema. The cartilage and perichondrium are not affected, and the contours of the auricle are unchanged. The changes may also be confined to one region such as the earlobe (jewelry) or retroauricular crease (hearing aid).

Diagnosis: If an allergy is suspected, skin tests should be done to identify the cause.

Differential diagnosis: It is necessary to exclude pyoderma, perichondritis, and cellulitis, which may occur as complications. Differentiation is also required from other dermatoses such as seborrhoeic dermatitis and psoriasis, which frequently occur in the retroauricular crease.

Complications: The cracked skin is susceptible to bacterial complications in the form of pyoderma, perichondritis, or cellulitis.

Treatment: The causes, if known, should be eliminated. Treatment of the skin lesions is based on dermatologic principles. Antibiotics are given only if bacterial superinfection has occurred.

Perichondritis of the Auricle

Definition: Perichondritis is an acute inflammation of the skin and perichondrium that also involves the auricular cartilage. The changes are localized and do not spread beyond the auricular cartilage.

Pathogenesis: Most cases are caused by a bacterial infection stemming from a small injury in the conchal cavity or auricle. Bacterial infection of the lateral portions of the auricle is often associated with perichondritis due to the close attachment of the skin to the perichondrium. The predominant causative organisms are staphylococci and Pseudomonas species. A toxic-allergic or autoimmune etiology is less common.

Symptoms: The clinical picture is characterized by severe pain of rapid onset and a feeling of tension. The auricular contours are effaced, and it is common to find swelling of the concha with marked tenderness. Blisters may develop on the skin. Areas that are devoid of cartilage, like the earlobe, are spared. Regional lymph nodes may be painful and enlarged, and systemic symptoms such as fever may occur.

Diagnosis: The tympanic membrane appears normal at otoscopy, and hearing loss does not occur unless otitis externa is also present. Blood tests generally show leukocytosis and elevated inflammatory parameters—C-reactive protein (CRP), erythrocyte sedimentation rate (ESR), or both.

If raw, weeping skin areas are found, a smear should be taken for bacteriologic examination and sensitivity testing, as a problem organism may be present.

Differential diagnosis: Perichondritis requires differentiation from an inflammation not involving the cartilage (eczema, dermatitis) and from a spreading infection (cellulitis, zoster oticus). Recurrent autoimmune polychondritis (relapsing polychondritis, see 10.3, p. 223) often begins in the auricle. In the absence of other symptoms caused by the destruction of bronchial, nasal or laryngeal cartilage, it is virtually indistinguishable from bacterial perichondritis.
Complications: Unless adequately treated, the inflammation will cause cartilage destruction with permanent auricular deformity.

Treatment: Primary treatment relies on systemic antibiotics that are active against staphylococci. The auricle and ear canal should be carefully and thoroughly cleansed. Antiseptic or antibiotic-containing ointments should be applied locally, and pain is controlled with nonsteroidal anti-inflammatory agents.

Auricular Cellulitis

Definition: Auricular cellulitis is an acute streptococcal infection of the subcutaneous tissue involving the auricle and its surroundings.

Pathogenesis: The streptococci usually gain access to the auricle through small injuries in the concha or external meatus.

Symptoms: The clinical manifestations typically consist of redness, swelling, and warmth of the auricle and its surroundings. The earlobe and adjacent facial skin are also involved. Malaise is present with associated fever and otalgia. The disease progresses swiftly without treatment.

Diagnosis: Inspection and cleansing of the ear canal, with evaluation of the tympanic membrane, are necessary to exclude a middle ear infection.

An attempt should always be made to identify the causative organism in smears and determine its antibiotic sensitivity.

Differential diagnosis: Differentiation is required from eczema, dermatitis, and perichondritis. Dermatitis generally does not produce systemic effects such as fever or an elevated white count. In perichondritis, the surrounding soft tissues and the earlobe usually are not affected. In zoster oticus, there is usually concomitant involvement of the inner ear or facial nerve.

Complications: Rare cases may be complicated by a necrotizing fasciitis (severe, intractable subcutaneous infection, often with the presence of anaerobes). Infections with group A streptococci, known systemic complications such as glomerulonephritis, rheumatic fever, or rheumatic endocarditis may develop (see §3.3 and §4.4, p. 114).

Treatment: The standard treatment is a high-dose regimen of penicillin G (e.g., 4 x 2 mega-IU), preferably by i.v. administration. Other antibiotics with anti-streptococcal activity can also be used. Pain is adequately managed with nonsteroidal anti-inflammatory agents. The auricle and ear canal should be carefully cleansed.

Herpes Zoster Oticus

Pathogenesis: This disease, known also as Ramsay Hunt syndrome, is caused by reactivation of the dormant varicella-zoster virus (VZV) in ganglion cells. Zoster oticus involves cranial nerves VII and/or VIII (and occasionally IX and X).

Symptoms: Patients initially experience ear pain or burning on one side in the absence of physical findings. Typical vesicles erupt shortly thereafter, no more than a few days after symptom onset. This is followed by hearing loss, vestibular complaints (vertigo, dizziness), and often by facial nerve palsy.

Diagnosis: Inspection reveals typical clustered (“herpetiform”) vesicles about the meatus and concha and occasionally on the pinna. There is accompanying lymphadenitis of the high cervical lymph nodes. Clinical examination may demonstrate facial nerve palsy as evidence of seventh and eighth cranial nerve involvement along with sensorineural hearing loss, nystagmus, and unilateral hyporeactibility of the vestibular organ. These signs may occur singly or in combinations.

The diagnosis can usually be made clinically. It can be confirmed by the direct electron microscopic detection of VZV in vesicular aspirate (costly) or later by at least a four-fold titer increase on serologic testing.

Differential diagnosis: In its early stage, herpes zoster oticus is important in the differential diagnosis of sudden hearing loss. The disease requires differentiation from other forms of otitis, especially bullous otitis externa. Other causes of seventh and eighth cranial nerve lesions such as otitis media, mastoiditis, labyrinthitis, cholesteatoma, and tumors of the ear and lateral skull base should be excluded.

Complications: An important local complication is secondary bacterial infection, usually with staphylococci or Pseudomonas species. Zoster meningoencephalitis

![Fig. 10.11 Herpes zoster oticus](image)

Typical bullae with acute inflammation in herpes zoster oticus. The 56-year-old man also displayed clinical signs of vestibular nerve dysfunction and facial nerve palsy.
results from the intracranial spread of the varicella-zoster virus. A severe, refractory post-zoster neuralgia may develop as a late complication, especially in older patients.

**Treatment:** Whenever herpes zoster oticus is suspected clinically, systemic therapy with acyclovir, valaciclovir, or famciclovir should be instituted without delay. The concurrent use of corticosteroids is recommended when facial nerve palsy is present. The lesions are also treated locally with an antiseptic solution.

**Prognosis:** Older patients in particular are apt to have residual morbidity with permanent functional deficits. The prognosis of facial nerve palsy after herpes zoster oticus is poorer than in cases of idiopathic facial paralysis.

**External Auditory Canal**

**Diffuse Otitis externa and Eczema of the Ear Canal**

**Pathogenesis:** An inflammatory condition of the external auditory canal involving the canal skin (eczema, dermatitis due to mechanical injury, toxicity, or allergy) gives rise to an acute bacterial infection of the skin with a mixed flora that includes gram-negative organisms (*Pseudomonas aeruginosa, Proteus mirabilis*) and anaerobes. Primary or secondary fungal infections of the ear canal may also develop (see Otomycosis, p.222). The tympanic membrane is also occasionally involved (myringitis). A warm, moist climate promotes the development of a diffuse otitis externa (swimmer's otitis).

**Symptoms:** The main initial symptom is itching. Pain is present with an acute infection. The patient may notice crusting, and a purulent aural discharge may occur. Obstruction of the ear canal can lead to conductive hearing loss.

**Diagnosis:** With eczema of the ear canal in the absence of acute infection, the canal skin appears dry, cracked, and scaly on otoscopic examination. The skin may be thickened and shows sites of desquamation. The presence of an infection is manifested by diffuse swelling of the canal skin with associated discharge or crusting. A fetid discharge means that anaerobes are involved. Bacteriologic examination is necessary only in cases with persistent or recurrent infection or if the diagnosis is uncertain.

**Differential diagnosis:** Acute otitis media or chronic supplicative otitis media can lead to an accompanying otitis externa and should be excluded by otoscopic examination. A fungal infection (see p.222) or tumor of the ear canal should be considered in persistent cases, and necrotizing otitis externa should be considered if otalgia is also present.

**Complications:** The cracked skin in otitis externa can allow bacterial entry causing perichondritis, cellulitis, or abscess formation. Necrotizing otitis externa may develop in predisposed patients.

**Treatment:** The first and most important step in treatment is meticulous, repeated cleansing and drying of the ear canal followed by the instillation of antiseptic, antibiotic drops that will reduce the swelling.

Steroid- and antibiotic-containing ear drops should be used for no more than two weeks due to the risk of sensitization, antibiotic resistance, and the development of a fungal infection.

Steroid- and antibiotic-containing ear drops are contraindicated in patients with a fungal infection of the ear canal, antibiotic hypersensitivity, or a perforated tympanic membrane. Emphasis is placed on meticulous aural hygiene, which includes protecting the ear from shampoos, soaps, and cotton-tipped swabs.

**Circumscribed Otitis externa**

**Definition:** Known also as a furuncle, this is a circumscribed lesion caused by an acute bacterial infection of the cartilaginous portion of the ear canal.

**Pathogenesis:** Local mechanical trauma and contamination of the ear canal (e.g., from an ear plug, dusty environment, bath water, or attempted self-cleaning of the ear) lead to obstruction of the hair follicles or glandular ducts, followed by a staphylococcal infection of the pilosebaceous units.

**Symptoms:** A furuncle in the ear canal presents as a very painful, tender swelling that can cause mild hearing loss and rarely leads to otorrhea. Patients are generally afebrile.

**Diagnosis:**

- **Inspection and palpation:** tragal tenderness accompanied by a circumscribed, very painful swelling in the cartilaginous portion of the ear canal.
- **Otoscopy:** pronounced swelling of the ear canal with debris in the residual lumen. Frequently the tympanic membrane cannot be seen, but it is normal.
- **Simple hearing test:** The ear canal may be swollen shut, causing some degree of conductive hearing loss.
- **Bacteriologic examination:** The purulent center of the lesion (pus pocket) can be opened with a small, blunt hook to obtain a smear.
Patients with recurrent furuncles should be examined for a predisposing systemic condition such as diabetes mellitus.

**Differential diagnosis:** Various disorders can produce similar clinical manifestations—e.g., foreign bodies in the ear canal, otitis externa accompanying chronic suppurative otitis media, an infected retroauricular atheroma, and tumors of the ear canal.

**Complications:** In rare cases, an abscess in the ear canal with involvement of the surrounding soft tissues may spread to infra-auricular and preauricular sites. Spread to the auricular cartilage may incite perichondritis. Superinfection with *Pseudomonas* can lead to necrotizing otitis externa in predisposed patients (see below). Furuncles may take a particularly severe course in patients with diabetes mellitus.

**Treatment:** The ear canal is *meticulously cleaned* and then treated locally for 1–2 days with 70% alcohol, which is applied hourly to a self-expanding foam or gauze wick inserted into the ear canal (Fig. 10.12). This treatment provides disinfection, reduces local swelling by fluid absorption, and exerts a cooling action that relieves pain. Crusts can be dissolved with antibiotic-containing ointment strips. After the swelling has subsided, antibiotic- and steroid-containing drops can be instilled into the ear. Nonsteroidal anti-inflammatory agents are administered for pain. Abscesses should be incised after they have become clearly demarcated. Systemic antibiotics are used in patients with systemic symptoms and severe local signs of infection.

**Necrotizing Otitis Externa**

**Pathogenesis:** Necrotizing otitis externa, also known as malignant otitis externa, is a dangerous, necrotizing form of otitis externa that occurs almost exclusively in older patients with diabetes mellitus.

Most cases begin with a simple otitis externa that becomes infected with *Pseudomonas aeruginosa*, leading to ulceration and osteitis on the floor of the ear canal. The bone infection may subsequently spread to the middle ear, skull base, retromandibular fossa, and parotid compartment.

**Symptoms:** The initial history is that of an insidious, persistent otitis externa that does not heal. At first there is moderate pain, which may become severe as the condition takes a chronic course. A fetid aural discharge may occur but is not always present.

Medications can be applied to foam wicks in the ear canal, which absorb liquids and exert a gentle pressure.

**a** Above: compressed foam wick before use. **b** Below: foam wick after fluid absorption.

**Diagnosis:**

- **Inspection:** reveals signs of infection in surrounding tissues.
- **Otoscopy:** almost always shows an ulcer on the canal floor with exposed, brownish bone and a fetid discharge.
- **Smear** (with sensitivity testing!): *Pseudomonas aeruginosa*.
- **Radionuclide bone scan, computed tomography:** These studies define the extent of the infection and bone destruction.

Diabetes mellitus is almost always present. Other immune defects should be excluded.

**Differential diagnosis:** Differentiation is mainly required from simple otitis externa and cholesteatoma of the external ear canal. A biopsy should be taken if a tumor is suspected. Complications from chronic otitis media can produce similar findings.

**Complications:** The infection can lead to otitis media, mastoiditis, petrositis, and soft-tissue abscess. Cranial nerve deficits (VII, VIII, IX, X, XI), sepsis, venous sinus thrombosis, or meningitis may occur in the late stage, causing the disease to become life-threatening.

**Treatment:** The ear canal is locally debrided and cleaned at regular intervals. In cases with minimal bone involvement, high doses of an antibiotic effective against *Pseudomonas aeruginosa* can be administered for 6 weeks. Diabetes mellitus should be closely monitored and adequately controlled. If there is poor response to conservative therapy, extensive involvement, or if complications arise, the affected bone should be resected. This surgery may be minor or may consist of petrosectomy, depending on the extent of disease.
Prognosis: The prognosis is guarded and depends on prompt, appropriate treatment. There is only a 50% chance of survival in cases that develop facial nerve palsy or venous sinus thrombosis.

Bullous Otitis Externa

Pathogenesis: Bullous otitis externa (flu-related otitis, hemorrhagic otitis externa) is presumed to have a viral etiology, but the exact causal agent is unknown. The influenza virus has been isolated in sporadic cases. The infection causes toxic capillary damage in the thin epithelial layer of the meatal skin and on the tympanic membrane, leading to the formation of hemorrhagic epithelial bullae.

Symptoms: The disease begins with severe otalgia of sudden onset, often followed by a bloody discharge from the ear canal. Both conductive and sensorineural hearing loss may develop.

Diagnosis: At otoscopy, serous or hemorrhagic bulla formation is observed on the epithelium in the bony portion of the ear canal and on the tympanic membrane. Rupture of the bullae can cause spontaneous bleeding from the ear canal, which contains fresh blood that later dries to form a crust.

Differential diagnosis: Toxic or traumatic injury to the ear canal or middle ear (barotrauma) can produce similar findings. Differentiation is also required from herpes zoster oticus and from tumors of the ear canal.

Complications: Involvement of the middle ear and/or inner ear (labyrinthitis) is relatively common and may be associated with sensorineural hearing loss and vertigo. Rarely, the infection ascends along the statoacoustic nerve and may incite a life-threatening encephalitis in the brainstem.

Treatment: A specific antiviral therapy is not available. Pain is treated with local anesthetic ear drops and nonsteroidal anti-inflammatory drugs. If bacterial involvement of the middle and inner ear is suspected, systemic antibiotics should be administered.

Otomycosis

Pathogenesis: Cerumen often harbors saprophytic fungi that have no specific pathologic significance. Aspergillus, Candida albicans, Mucor, and dermatophytes may, however, aggressively infect the skin of the medial ear canal if the milieu has been altered by the use of steroid- and antibiotic-containing ear drops or other factors. A warm, moist climate is conducive to fungal infections, which are most common during the summer months.

It is rare for a fungal infection of the ear canal to signify a general decline in host resistance.

Symptoms: Otomycosis is manifested less by pain than by severe itching and a feeling of fullness in the affected ear.

Diagnosis: The fungi often appear otoscopically as a white, yellow or black membrane lining the swollen, erythematous skin of the ear canal. The bony portion of the canal is affected almost exclusively. Mycelia can be identified in direct samples. The causative organism is established by microbiologic examination.

Differential diagnosis:

The harmless, superficial fungal colonization of cerumen or drainage should be distinguished from an actual invasive mycosis.

Otomycosis is occasionally difficult to distinguish from other forms of otitis externa, especially diffuse otitis externa. Mixed forms commonly occur. A fungal infection may also develop in the setting of chronic suppurative otitis media.

Course: Otomycosis typically runs a refractory course and has a tendency to recur.

Complications: A fungal infection of the tympanic membrane epithelium can lead to perforation and subsequent otitis media. Spreading fungal infections with necrosis occur only in immunocompromised patients.

Treatment: First it is essential that the ear canal be thoroughly cleaned and dried. Once this has been accomplished, local antifungotics can be administered. It is also necessary in many cases to soften the uppermost epithelial layer with salicylate-containing solutions in order to enhance the antifungal action of specific medications. Systemic antifungal therapy is necessary only in immune-suppressed patients.
10.3 Other inflammatory conditions of the external ear

Recurrent polychondritis
This is a chronic autoimmune disease directed against cartilage tissue. Auricular perichondritis is often seen as the initial manifestation. Involvement of the cartilage in the airways, larynx, and nose occurs in later stages. The disease takes a chronic course that is marked by deformities of the auricle and nose, chronic bronchitis, and occasional dyspnea resulting from cartilage damage in the trachea and larynx.

Synonyms: relapsing polychondritis, systemic chondromalacia, chronic atrophic panchondritis.

Diagnosis: The diagnosis is based on the presence of a systemic inflammatory disease, the detection of antibodies against cartilage tissue, the histologic examination of cartilage tissue, and the course.

Treatment: Treatment consists of oral corticosteroids. Other immunosuppressants such as azathioprine or cyclosporin are also used.

Chronic chondrodermatitis nodularis helicis (Winkler disease)

The diagnosis of chondrodermatitis is based on the presence of a very painful epithelial nodule with an umbilicated center on the free border of the helix or antihelix (Fig. a). This is an inflammatory skin lesion of the cutaneous-perichondrial unit. Its cause is unknown. Older patients are predominantly affected. Treatment consists of complete excision, and subsequent histologic examination confirms the diagnosis. The differential diagnosis includes tumors and gouty tophi. Recurrence is not uncommon even after complete excision.

Gouty tophi

Gouty tophi may form near the joints and on the auricular cartilage. They appear as small, pale, freely movable subcutaneous nodules on the helical rim. Generally, there is no need for local treatment.

Lymphadenosis cutis benigna (Bäverstedt disease)

This lesion appears as a firm, reddish nodule on the earlobe (Fig. b). The lymphadenosis represents a cutaneous manifestation of infection with *Borrelia burgdorferi*, and patients often give a prior history of tick bite. The treatment is the same as for a *Borrelia* infection.

Granulating otitis externa

Circumscribed or diffuse granulations may form on the skin of the bony ear canal and on the tympanic membrane, occurring spontaneously or as a sequel to otitis externa or ear surgery. Often the precipitating cause cannot be determined, and the inflammation usually takes a refractory course. Treatment consists of removing the granulations followed by the topical application of antibiotics and corticosteroids. Necrotizing otitis externa and a tumor should be excluded.

Specific otitis externa

Syphilis (mainly stage II), *Mycobacterium tuberculosis*, and atypical mycobacteria are very rare causes of otitis externa.
10.5 Tumors of the External Ear

Tumors of the auricle are relatively common, easily recognized and treatable, whereas tumors of the ear canal are rare and are often misinterpreted. It is important to make an early diagnosis and to differentiate benign tumors from premalignant and malignant lesions of the external ear.

Tumors of the Auricle

The auricle is heavily exposed to weathering effects due to its location. As a result, the auricle is a common site of occurrence for epithelial skin tumors that are caused by, or related to, actinic exposure. Men over 60 years of age are predominantly affected. Table 10.1 reviews the principal tumor entities, which together account for more than 90% of auricular tumors. Fibromas, histiocytomas, chondromas, ceruminomas (arising from the cerumen glands), cylindromas (Spiegler tumor, carcinoma of the eccrine sweat glands; not to be confused with adenoid cystic carcinoma), hemangiomas, and lymphangiomas are examples of rare nonepithelial tumors that may affect the external ear. It is also important to distinguish true neoplasms from nevoid lesions, cysts, inflammatory changes such as chondrodermatitis and gouty tophi (see 10.3, p. 223), and deformities (see 10.2, pp. 210–215).

The diagnosis is usually made histologically following excision of the tumor. This surgery may require reconstruction of the auricle, depending on the location and extent of the lesion. The specific reconstructive measures (see 10.2, p. 215) depend on the overall situation, including the need for postoperative radiotherapy. A suspected malignant tumor should be excised and its margins assessed by frozen tissue histology. Extensive tumors may necessitate a complete auricular resection.

The differential diagnosis includes skin lesions such as cysts, keloids, otophymas, and nevoid lesions. Nevus mainly require differentiation from melanoma. The prophylaxis of auricular tumors consists of adequate protection from sun exposure.

Tumors of the Ear Canal

Isolated tumors of the external auditory canal are rare. Involvement of the ear canal most commonly occurs in association with an auricular tumor. This concomitant involvement of the ear canal by an auricular malignancy generally alters the approach to treatment, as a simple excision is no longer possible in most cases. Unlike tumors of the auricle, isolated tumors of the ear canal are frequently misinterpreted and are treated for some time as otitis externa. The most common malignant tumor of the ear canal is carcinoma of the canal skin. Less common are adenoid cystic tumors, adenocarcinomas, and basal cell carcinomas. These tumors usually present as a painful, ulcerated, nonhealing lesion in the skin of the ear canal. Many cases manifest bleeding and secondary infection with chronic, purulent otitis.

For this reason, every nonhealing, ulcerated or granulating lesion of the ear canal should be biopsied under the operating microscope.

The differential diagnosis should include chronic otitis externa, especially necrotizing otitis externa, chronic otitis media with mucosal polyps, a middle ear tumor, and a penetrating parotid tumor. Further investigation relies on computed tomography, which can define the extent of tumor infiltration in the bone, parotid compartment, and middle ear.

Treatment is generally surgical, with or without postoperative irradiation. The prognosis depends largely on the extent of disease but tends to be unfavorable compared with auricular tumors.

Fig. 10.13 Tumors of the auricle
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11.1 Anatomy, Physiology and Examination of the Middle Ear

The middle ear, with its complicated anatomy, occupies a central position in the temporal bone. Sound transmission by the ossicles and the ventilation of the temporal air cells via the eustachian tube are complex mechanisms, and disturbances of these mechanisms account for much of the pathology of the middle ear. This unit explores the anatomy and function of the middle ear in greater detail as a follow-up to 7.1 (pp. 155–156). Understanding these principles forms the basis for examination of the middle ear.

Special Anatomy and Physiology of the Middle Ear

As noted in 7.1 (p. 155), the middle ear consists of three parts: the tympanic cavity bounded laterally by the tympanic membrane, the system of temporal bone air cells, and the eustachian tube. When the middle ear transmits sound waves (alternating variations in air pressure), it performs impedance matching by equalizing the differences in impedance (acoustic resistance) between the air and perilymph. The eustachian tube equalizes static differences in air pressure between the middle ear and external auditory canal.

Tympnic Membrane

*Function:* The tympanic membrane (eardrum) has two functions: it gathers sound like the membrane of a microphone (see 7.1, p. 155), and it provides sonic shielding of the round window membrane. Sound waves that directly impinge on the round window can counteract the perilymphatic fluid displacement induced by the stapes, reducing the sensitivity of the cochlea.

*Anatomy:* The normal anatomy of the tympanic membrane is shown in Fig. 11.1. It consists of two portions called the pars tensa and pars flaccida. The much larger *pars tensa* is a funnel-shaped area stretched between the malleus handle and the bony ear canal. It is composed of three layers (Fig. 11.2):

- The outer layer of the pars tensa, called the *cutaneous layer*, consists of smooth, stratified squamous epithelium that normally reflects light.
- The inner layer bordering the tympanic cavity, called the *mucosal layer*, consists of a single layer of squamous epithelium.
- Between the outer and inner layers is the *lamina propria*. It consists of two layers of connective-tissue fibers: an outer layer of radially directed fibers (*radiate layer*) and an inner layer of circular fibers (*circular layer*). These fibers blend with the fibrocartilaginous ring at the circumference of the tympanic membrane. This ring anchors the tympanic membrane in the tympanic sulcus of the bony ear canal.

The *pars flaccida* (synonym: Shrapnell membrane) is located superior to the malleolar folds. It cannot always be clearly identified at otoscopy and may blend with the superior canal wall. Microscopically, the pars flaccida lacks the reinforcing fibrous layers that are present in the pars tensa. As a result, this portion of the tympanic membrane often retracts first in response to negative pressure in the middle ear, creating an epithelial pocket.
**Tympanic Cavity**

The aerated tympanic cavity (synonyms: middle ear space, middle ear cleft) allows for unrestricted mobility of the tympanic membrane, which is necessary for sound transmission. Most of the air enters the tympanic cavity through the eustachian tube, but some gases diffuse directly into the middle ear through blood vessels in the mucosa.

**Levels of the tympanic cavity:** The tympanic cavity (see Fig. 7.1, p.154, and Fig. 11.3) is divided into three levels relative to the plane of the tympanic membrane:

- The portion of the tympanic cavity at the level of the tympanic membrane is called the *mesotympanum*. It contains the round window, the oval window with the stapes, and the promontory (bony prominence overlying the basal turn of the cochlea).

- Above the plane of the tympanic membrane is the *epitympanum* (synonyms: attic, epitympanic recess). The tympanic part of the facial nerve defines the boundary between the epi- and mesotympanum on the medial wall of the middle ear. The epitympanum contains the principal mass of the auditory ossicles with their associated ligaments and several mucosal folds. The epitympanum is small and contains little air, and inflammations may become encapsulated in that area. The epitympanum communicates with the mastoid antrum via the aditus ad antrum and also with the air cells of the mastoid process. The antrum contains the bony prominence of the lateral semicircular canal, which is often the first part of the labyrinth to be attacked by an osteoclastic disease process in the middle ear. A thin bony layer, the tegmen tympani, forms the roof of the epitympanum and separates it from the middle cranial fossa.

- Below the level of the tympanic membrane is the *hypotympanum* (synonym: hypotympanic recess). It borders on the bulb of the jugular vein and contains cells (tympanic cells) that communicate with the mastoid air cells.

**Auditory ossicles:** The ossicles in the middle ear have several distinctive features. They are the smallest bones in the human body and are freely suspended, being nourished entirely through their periosteal attachments. They are attached by thin tendons to the smallest muscles in the body, the intra-aural muscles (tensor tympani and stapedius).

The handle (manubrium) of the *malleus* is attached along its length to the tympanic membrane. The tip of the handle forms a central spoon-shaped depression, the umbo, which is an important landmark for evaluating the tympanic membrane.

The next bone in the ossicular chain is the *incus*, which articulates with the malleus. The head of the malleus and the body of the incus are located in the epitympanum and comprise most of the mass of the auditory ossicles.

The long process of the incus articulates with the *stapes*. The footplate of the stapes is attached to the
11.1 Function of the articular connections and mass of the auditory ossicles

The articular connections and mass of the auditory ossicles are not necessary for sound transmission. In birds, for example, sound is transmitted from the tympanic membrane to the inner ear by one thin ossicle, the columella. The extra mass of the ossicles probably serves more to adjust the resonance of the middle ear system as a means of reducing sound transmission by bone conduction. Chewing, for example, produces loud noises that can mask sounds from external sources. The intra-aural muscles may also help reduce this “internal noise.” The joints in the ossicular chain are less for improving sound transmission than for adjusting the position of the ossicles in response to atmospheric pressure changes (see also 7.1, p. 155). The intra-aural muscles may assist in this function by keeping the ossicular joints mobile. By promoting “articular hygiene,” the muscles help to maintain the important pressure-equalizing function of the joints. Sound-induced movements of the ossicles are too small in themselves to keep the joints functioning.

rim of the oval window by the elastic annular ligament, creating a movable interface between the stapes footplate and the perilymphatic space. 

**Intra-aural muscles:** The stapedius muscle inserts onto the head of the stapes and occupies a bony canal parallel to the mastoid part of the facial nerve, which also innervates the muscle. The tensor tympani muscle lies parallel to the eustachian tube and is innervated by the trigeminal nerve. It inserts onto the head of the malleus. Reflex contractions of the intra-aural muscles (stapedial reflex, see 8.4, Objective Hearing Tests, pp.184–191) may serve to moderate the internal noise level produced by mastication and speech, and it may also be a mechanism to protect the inner ear from high, sustained external noise levels.

**Anatomical relations of the tympanic cavity:** The cavity of the middle ear is surrounded by functionally and clinically important structures:

- The lateral wall is formed by the tympanic membrane and the bony ear canal.
- The medial wall borders the cochlea.
- The inferior wall borders the bulb of the jugular vein.
- The superior wall borders the dura of the middle cranial fossa.
- The anterior wall borders the internal carotid artery.
- The posterior wall borders the mastoid part of the facial nerve.

The tympanic cavity is traversed by the chorda tympani, which arises from the facial nerve and contains gustatory fibers for the anterior two-thirds of the tongue (see 14.1, Clinically Relevant Anatomy and Function of the Facial Nerve, p.290).

**Blood supply and innervation:** The tympanic cavity receives its blood supply from various branches of the external carotid artery (middle meningeal artery, ascending pharyngeal artery, maxillary artery, and stylo-mastoid artery).

It derives most of its sensory innervation from the tympanic nerve, which arises from the glossopharyngeal nerve (see Fig. 16.3, p.315). This connection accounts for the “referred otalgia” that can occur in association with pharyngeal processes. The tympanic nerve also has connections with parasympathetic portions of the glossopharyngeal nerve, sympathetic fibers of the internal carotid plexus, and with the trigeminal nerve.

**Air Cells of the Temporal Bone**

The mucosa-lined air cells of the temporal bone all communicate with the tympanic cavity and, like the middle ear itself, are aerated via the eustachian tube. The pneumatization of the mastoid and other parts of the temporal bone develops gradually during childhood, similar to the development of the paranasal sinuses (see also pp. 4–6). The mastoid antrum and several adjacent air cells are generally present in infants. The degree of pneumatization that subsequently develops is variable and depends partly on the ventilation of the tympanic cavity, the function of the eustachian tube, and the inflammatory history of the middle ear. As a rule, few pneumatized cells develop in children who have a history of chronic otitis media. On the other hand, “normal” pneumatization of the temporal bone can have a broad range of appearances, analogous to the paranasal sinuses. With extensive pneumatization, the air cells may extend into the zygomatic arch, the temporal squama, and the petrous apex.

The function of the temporal bone air cells in humans is unknown. Perhaps a greater air volume is better for equalizing pressure differences and thus can help to protect the middle ear. The air volume behind the tympanic membrane is unimportant for sound transmission in humans (unlike small rodents, for example, in which the air space of the tympanic cavity contributes greatly to the physical properties of the sound-conducting apparatus in the middle ear).

**Eustachian Tube**

The eustachian tube connects the tympanic cavity with the nasopharynx, where the inlet of the tube forms a funnel-shaped orifice behind the choana (see Fig. 1.5, p. 4).

**Functions of the eustachian tube:**

- Ventilates the tympanic cavity and air cells
- Equalizes pressure differences between the tympanic cavity and the atmosphere
The eustachian tube consists of a bony part (lateral third) and a cartilaginous part (medial two-thirds). At their junction is the narrowest part of the tube, the isthmus.

- Drains the middle ear spaces
- Creates a barrier to ascending infection

The eustachian tube runs more horizontally in infants and small children than in adults. It is considerably shorter and broader and consists of softer cartilage. Presumably this compromises the overall function of the eustachian tube, accounting for the higher incidence of otitis media in children. By about 7-10 years of age, the eustachian tube closely approximates the adult tube in its anatomy and function.

**Anatomy (Fig. 11.4):** The lateral third of the eustachian tube framework consists of a bony canal that conveys the tube in addition to the tensor tympani muscle. The medial framework is a patent cartilaginous tube that is suspected from the skull base. The narrowest portion of the tube, called the isthmus, lies at the junction of the bony and cartilaginous parts. Inflammatory stenosis may develop at that location. The medial part of the tube is surrounded by fatty tissue, glands, veins, and muscles. It is actively opened by the tensor veli palatini muscle (see p. 72). Eustachian tube function in general relies on a complex balance between opening forces such as muscle tone, middle ear pressure, and cartilage resilience and closing forces generated by tissue pressure, mucosal surface tension, and a negative pressure in the middle ear.

**Examination**

**History**

There are no complaints that are specific for middle ear disease. Otalgia, aural discharge (otorhea), a feeling of pressure, or hearing loss can occur with diseases of the external ear or inner ear. But a history of chronic inflammatory ear diseases usually signifies otitis media. This type of inflammation can lead to scarring and decreased ventilation of the middle ear.

Before any otologic manipulations are performed, the patient should be asked specifically about tympanic membrane perforation, previous trauma to the ear, or any surgery of the middle ear. Most patients will be able to provide this information.

**Otoscopy**

Otoscopy with evaluation of the tympanic membrane is the mainstay for the investigation of middle ear diseases.

The examination begins with inspection of the external ear and its surroundings, particularly the mastoid. The technique of otoscopy and normal tympanic membrane findings are covered in 8.1 (pp. 166–167) and 10.1 (pp. 208–209). For the otologist, evaluation of the tympanic membrane with a stereoscopic microscope is essential for the diagnosis of middle ear diseases.

**Interpretation:** Normally, otoscopy will reveal nothing more of the middle ear than the lateral aspect of the tympanic membrane. A normal tympanic membrane has differentiated features, is mobile, and reflects light (see p. 167). The mobility of the tympanic membrane provides clues to the condition and ventilation of the tympanic cavity.

**Function tests:** Tympanic membrane mobility can be tested actively or passively during otoscopy.

- **Passive mobility test:** The external ear canal is sealed with the speculum, and a positive pneumatic pressure is produced in the ear canal, causing movement of the tympanic membrane (Siple pneumatic otoscopy).
- **Active mobility tests:** In active tests of tympanic membrane mobility, air is forced up the eustachian tube into the middle ear, inducing an outward movement of the tympanic membrane. Usually this movement is most clearly appreciated in the posterosuperior quadrant (see Fig. 11.1, p. 228). Since this also tests eustachian tube patency, the active mobility tests are particularly important for examination of the middle ear.

- In the Valsalva maneuver, the patient is told to swallow, then pinch the nostrils and “bear down” to produce a positive pressure in the pharynx.
Fig. 11.5  Politzer maneuver

Air is forced into the nasopharynx by occluding one nostril and squeezing a Politzer bag. Meanwhile the patient is told to say “cuckoo” or “Coca Cola” (to close the soft palate). The positive pressure in the nasopharynx travels through the eustachian tube into the middle ear, and the patient hears a snapping sound.

- In the Politzer maneuver, air is forced into one side of the nose by squeezing an air bag while the other nostril is occluded and the soft palate is closed off (Fig. 11.5).
- In the Toynbee maneuver, a negative pressure is created in the pharynx by having the patient swallow with the nostrils pinched shut. This induces an inward movement of the tympanic membrane.

None of these maneuvers should be used in patients with an acute inflammation of the middle ear or nasopharynx.

**Interpretation:** A negative Valsalva maneuver may be due to:
- Improper technique
- A nonpatent eustachian tube
- A thickened, scarred tympanic membrane
- A perforated tympanic membrane

When the tympanic membrane is perforated, there may be an audible rush of air through the opening. With drainage in the ear canal with a perforated tympanic membrane, the Valsalva maneuver may produce visible air bubbles—a strong suggestive sign of otitis media.

**Hearing Tests**

The typical clinical presentation of middle ear disease includes **conductive hearing loss**, in which the Weber test is lateralized to the affected ear and the Rinne test is negative (see 8.1, p. 167).

**Pure-tone audiometry** shows a difference between bone conduction and air conduction thresholds (air-bone gap, see 8.3, p. 178).

**Tympanometry**, which can be performed only with an intact tympanic membrane, may show a flattening or shifting of the peak into the negative pressure region (type B and C tympanograms, see 8.4, p. 185). If the tympanogram is normal, the function of the ossicles can be assessed by testing the **stapedial reflex** (see p. 185).

Middle ear disease generally leads to an absence of **otocoustic emissions** (OAEs, see pp. 189–191), which cannot be transmitted laterally across the diseased middle ear.

The following findings, then, are indicative of **normal middle ear function**:
- Normal otoscopic appearance of the tympanic membrane
- Positive Rinne test
- Normal tympanogram (type A) and a positive stapedial reflex
- Detectable OAEs (assuming normal cochlear function)

**Imaging Studies**

Since visual inspection of the middle ear is generally limited to the tympanic membrane, imaging studies are an important adjunct to function tests. Radiographic studies are the most rewarding due to the preponderance of bony structures.

**Standard projections** of the temporal bone will invariably superimpose numerous structures. This problem can be reduced by obtaining **special views** of the temporal bone in which fewer structures are superimposed. The most important of these views are:
- The Schütter view: projection along the ear canal to demonstrate the mastoid air cells
- The Stenvers view: projection angled 45° forward to demonstrate the petrous ridge and petrous apex

X-ray films are routinely obtained on both sides for purposes of comparison.

Even with special views, a number of structures are still superimposed and some essential fine details in the middle ear may be obscured. Thus, conventional radiographs are of limited value due to their relatively poor sensitivity and specificity and are unnecessary when computed tomography is performed.
Computed tomography: The most important modality for temporal bone imaging is **high-resolution thin-slice computed tomography (CT)**, which has superseded conventional radiographs owing to its ability to define specific bony structures for specific investigations. The study consists of axial and/or coronal scans, usually obtained with a slice thickness of 0.5–1.5 mm. Bone windows are used to define the fine structural details and boundaries of osseous structures. Contrast administration is rarely necessary in examinations of the middle ear.

Other imaging modalities: **Magnetic resonance imaging (MRI)** is used less frequently for middle ear imaging because it is inferior to CT for defining bony structures. It can be helpful, however, for selected indications such as tumors. This also applies to **angiography**, which is used for investigating tumors, suspected vascular lesions, pulsatile tinnitus, etc.

**Diagnostic Tympanotomy**

Surgical exploration of the middle ear (tympanotomy) is also available as a diagnostic option. The tympanic cavity is opened for inspection through the ear canal under an operating microscope by incising the canal skin in front of the tympanic membrane and reflecting the skin and membrane as a flap. The bony canal wall may also have to be taken down, depending on the pathology. In most cases, diagnostic tympanotomy is combined with the surgical correction of any abnormalities that were noted in preoperative studies.
11.2 Pathophysiology and Otoscopic Features of Otitis Media

This explores the relationships between the development of chronic inflammatory middle ear diseases and the resulting chronic structural changes ("scars") that are accessible to otoscopic examination. The central elements in the pathophysiology of otitis media are impaired ventilation and mucosal inflammation in the middle ear. They lead to typical changes in the tympanic membrane, various clinical forms of chronic otitis media, and functional sequelae. It is important to distinguish between the inflammatory disease process itself and its sequelae, which often persist after the disease has resolved. The disease processes and associated clinical manifestations of otitis media are described in the next (pp. 238–249).

Pathophysiology of Otitis Media

The most common pathologic process in the middle ear is inflammation, termed otitis media. Regardless of its clinical presentation, otitis media is the product of fundamental pathophysiologic processes that can produce a number of typical otoscopic changes (see pp. 235–237).

General

The pathophysiology of most chronic middle ear diseases is based largely on two functional disturbances: impaired middle ear ventilation and inflammation. These mechanisms are closely interrelated and often cannot be separated from each other in any given case. A chronic impairment of middle ear ventilation leads to inflammation of the mucosa, which in turn compromises eustachian tube function and middle ear ventilation (Fig. 11.7). This vicious cycle is a pivotal element in most inflammatory conditions of the middle ear.

Causes of Impaired Ventilation

Impaired ventilation of the middle ear is almost always caused by eustachian tube dysfunction. Decreased tubal patency is more common in this regard than excessive patency. Various factors can compromise the ventilating function of the eustachian tube:

- Stenosis of the tube lumen due to inflammatory mucosal swelling (e.g., caused by an upper respiratory viral infection). The air in the tympanic cavity is absorbed, and a negative pressure develops that further compromises eustachian tube function.
- A negative pressure can also develop in a healthy middle ear due to a rapid rise of ambient air pressure, as during an aircraft landing (see 11.4, Barotrauma, p. 251). The mucosa of the eustachian tube collapses, and the negative pressure itself incites mucosal swelling.
- Extrinsic obstruction of the tube, as by a tumor.
- Deficient active opening of the tube by the tensor veli palatini muscle. The special anatomy of the eustachian tube in small children (see 11.1, p. 231) hampers the function of the tensor veli palatini. Malformations of the jaw and palate can further compromise or even disable the tube-opening muscles, resulting in chronic inflammation of the middle ear.
- A congenital or acquired bony stenosis or stricture due to scarring.

Excessive patency of the eustachian tube may also cause a negative pressure to develop in certain compartments of the middle ear. The negative pressure during inspiration is transmitted through the patent tube into the tympanic cavity. This negative-pressure effect is magnified when air is forcibly inhaled through the nose (sniffing), which may be done habitually. As mucosal folds are raised in the middle ear, a chronic negative pressure may develop, especially in the epitympanum, leading to retraction of the tympanic membrane and chronic inflammation.

Infection and Inflammation

Infections and noninfectious inflammations play another key role in the pathogenesis of the otitis media. **Adenoiditis:** Initial contact with microorganisms and environmental agents in infants and small children can induce intensive immunologic-inflammatory changes affecting the tissues in Waldeyer’s ring. The adenoids (pharyngeal tonsils) are particularly important in the development of otitis media. The size of the adenoids is of relatively minor importance because the adenoids rarely if ever cause direct mechanical obstruction of the eustachian tube. The problem, rather, is chronic adenoiditis, which creates a reservoir for pathogenic micro-organisms and induces adenoid hyperplasia. Adenoiditis is promoted in turn by obstructed nasal breathing and by rhinitis or rhinosinusitis.

**Infections of the middle ear mucosa:** Viral and bacterial infections of the upper respiratory tract, which are common in children, can also directly affect the middle ear mucosa. They have a tendency to ascend
The central pathophysiologic process is eustachian tube dysfunction interacting with a chronic inflammation of the middle ear mucosa (shown in green). Various causal factors (red) can initiate this cycle, which in turn can have various sequelae (blue).

Through the eustachian tube into the middle ear (tubogenic infection). With a perforated tympanic membrane, gram-negative bacteria in particular can also enter the middle ear from the external canal, inciting an acute otitis media or perpetuating a chronic inflammation.

**Noninfectious inflammations:** Allergic or toxic inflammations of the upper respiratory tract contribute to adenoiditis and nasal airway obstruction or (less commonly) may spread to involve the middle ear mucosa. The reflux of gastric juice may also contribute to the inflammation. This mechanism may be particularly important in infants and small children who have a short eustachian tube that offers little protection.

**Otoscopic Features of Otitis Media**

The pathophysiologic processes described above are difficult to detect directly by clinical means. They do produce typical effects, however (see Fig. 11.7), that can be detected by otoscopic inspection. We are dealing with a descriptive classification in which specific pathologic significance cannot be ascribed to the observed changes.

First, it is helpful to distinguish changes in the external ear canal from changes in the middle ear (see 10.1, p. 208). Both can produce similar tympanic membrane changes, and the cause is not always easy to recognize. The history and other findings generally provide an adequate basis for differentiation, however.

Otoscopic changes in the middle ear are classified into **acute inflammatory**, **chronic inflammatory** and **cicatricial** changes. The presence of one type of change does not exclude any of the others. For example, a cicatricial change (scar) may also be acutely or chronically inflamed.
Discharge perforation with tympanic membrane

Umbo

Bubbles

Fig. Partial middle ear effusion

Serous effusion behind an uninfamed tympanic membrane, which is slightly retracted. Air bubbles are clearly visible.

Acute Inflammatory Changes

An acute inflammation of the middle ear generally incites acute inflammatory changes in the tympanic membrane. The eardrum is thickened, erythematous, and less transparent than normal (Fig. 11.8).

But the transparency of the tympanic membrane shows individual variations even when the membrane is not inflamed, and so this is not a reliable sign. The swelling and thickening of the tympanic membrane give it an irregular surface that is no longer smooth and reflective, but opaque. The typical light reflex is either absent or fragmented. When fluid collects in the middle ear, the tympanic membrane may bulge outward. This bulging is often first evident in the pars flaccida. As the inflammation progresses and the fluid pressure rises, the tympanic membrane may become perforated. Generally, the perforation begins in a small area of the eardrum, allowing the fluid to drain into the ear canal. If an open connection exists between the acutely inflamed middle ear mucosa and the discharge in front of the tympanic membrane, pulsations may be transmitted from the inflamed mucosa. Pulsatile discharge in the ear canal is indicative of acute otitis media, therefore.

Middle Ear Effusion

A middle ear effusion is a fluid collection in the middle ear without a perforated tympanic membrane and with no signs of acute inflammation (Fig. 11.9).

Partial middle ear effusion: Air bubbles or an air-fluid level behind the tympanic membrane signify the entry of air, confirming at least some degree of ventilation through the eustachian tube. Generally, then, a partial effusion in the middle ear signifies a subacute, serous inflammation with a good prognosis. This type of effusion is typically found after viral infections of the upper respiratory tract.

Complete middle ear effusions devoid of air can be detected indirectly by noting an immobile tympanic membrane at otoscopy or by recording a flat tympanogram (see Fig. 8.19b, p.185). Most of these effusions are chronic and become increasingly mucoid as the mucosa forms increased numbers of goblet cells. The color of the effusion changes from amber to grayish, dark, or blue.

The color of the effusion, however, is not a reliable criterion for assessing the duration or composition of a middle ear effusion.

Chronic Tympanic Membrane Perforation

Findings: A tympanic membrane perforation without acute inflammatory changes is usually easy to detect at otoscopy (Fig. 11.4, p.231). Basically the tympanic membrane has a strong capacity for self-healing, and even large perforations can heal spontaneously. But various cicatricial and atrophic changes in the middle ear can result in a nonhealing perforation, which is considered a hallmark of chronic otitis media. The tympanic cavity is visible through the perforation. The remaining tympanic membrane and the tympanic cavity can have various otoscopic appearances:

- Cicatricial, noninflammatory change
- Chronic inflammatory changes with thickened, hyperplastic mucosa and a mucous discharge
- Acute inflammatory change with marked erythema and a purulent discharge (Fig. 11.14, p.231)

Sequela of tympanic membrane perforation: A chronic tympanic membrane perforation can contribute to the resolution of chronic otitis media, but it can also pose a hazard. The following adverse sequelae may occur:

- Recurrent infections of the middle ear may develop via the external canal, with associated recurrent otitis. This may also adversely affect the inner ear (serous labyrinthitis, see 12.2, p.263).
- The degree of conductive hearing loss depends on the size and location of the perforation.
- A cholesteatoma may form due to the ingrowth of squamous epithelium into the middle ear. Chole-
steatomas usually develop from atrophic epithelial pockets in the tympanic membrane or from a marginal perforation. They rarely develop with a dry, central perforation that does not reach the fibrocartilaginous ring.

A chronic tympanic membrane perforation can also have desirable effects:
- A chronic perforation can contribute to the healing of mucosal inflammation, since it provides a route for middle ear ventilation.
- The perforation also provides a route for fluid drainage from the middle ear, resulting in relief of pain.

The adverse sequelae predominate in most cases, however, and surgical closure of the perforation (especially with intact eustachian tube function) is indicated (myringoplasty or tympanoplasty, see 11.2, p. 245–246).

**Atrophic Scars of the Tympanic Membrane**

A perforated tympanic membrane may undergo incomplete healing and scarring that cover over the perforation. This is particularly likely to occur after a chronic inflammation of the tympanic membrane. The lamina propria can no longer reconstitute its normal fibrous structure, and consequently this layer is thin or absent; the tympanic membrane loses its normal strength and tension. Various conditions may ensue, depending on the size of the atrophic scar and the ventilation of the middle ear, and these conditions have varying otoscopic appearances:

**Circumscribed atrophic scar** (Fig. 11.10): Only part of the tympanic membrane is atrophic, usually an area located in the posterosuperior quadrant. With normal middle ear ventilation and eustachian tube function, the scar lies in the anatomical plane of the tympanic membrane. The atrophic area is abnormally mobile in response to pneumatic otoscopy or a Valsalva maneuver.

**Retraction pocket**: A circumscribed atrophic scar in the tympanic membrane is retracted inward by a negative pressure in the tympanic cavity. A sustained negative pressure causes the pocket to expand in the middle ear. At some sites in the pocket the epithelium can no longer migrate laterally into the ear canal. The trapped, stagnant squamous debris may then become infected, resulting in the formation of a cholesteatoma (see pp. 243–244).

**Middle ear atelectasis**: The entire tympanic membrane is very thin and markedly retracted. The tympanic cavity is no longer aerated, and the fine squamous epithelium of the tympanic membrane lines the mesotympanum. Occasionally it is difficult to distinguish a complete tympanic membrane perforation from middle ear atelectasis by otoscopic examination.

**Sclerosis and Fibrosis**

Besides atrophic scars, fibrous and sclerotic scars can also form in the middle ear. The following types are distinguished according to the nature and location of the scars.

**Sclerotic tympanic membrane scar**: The fibrous lamina propria of the tympanic membrane shows typical calcific deposits that are clearly recognized as white areas (Fig. 11.11). With a normal position of the tympanic membrane and normal middle ear ventilation, sclerotic scars have no adverse sequelae and do not cause hearing impairment.

**Tymanosclerosis**: The sclerosis predominantly affects the tympanic cavity in addition to the tympanic membrane. Calcium deposits form in the middle ear mucosa and can cause ossicular fixation with associated impairment of sound conduction. Tymanosclerosis has a strong tendency to recur after surgical treatment, which often provides only a temporary improvement in hearing.

**Fibrosis**: This refers to extensive scarring of the tympanic membrane or, more commonly, the tympanic cavity in the absence of calcifications. The sequelae and otoscopic findings are similar to those in tymanosclerosis.
11.3 Otitis Media

The most common diseases of the middle ear are inflammations. Infections play a major role in these diseases. As explained in the previous section, otitis media and impaired middle ear ventilation are closely interrelated. This focuses on the various inflammatory conditions that can affect the middle ear and their immediate sequelae in the form of inflammatory complications. Chronic structural sequelae (“scars”) were described in the preceding.

Classification and Terminology

The classification of otitis media (middle ear inflammation) is not standardized, and the terminology is diverse and not always logical. We showed in the previous section that the various forms of otitis media are on a continuum. For example, acute otitis media is generally followed by several weeks of middle ear effusion like that occurring in serous or otitis media with effusion. But unlike acute otitis media, otitis media with effusion does not necessarily present with initial pain, and it may progress to acute otitis media when an acute infection supervenes. Nevertheless, various inflammatory conditions of the middle ear can still be distinguished from a clinical and learning standpoint, and these entities will be described separately below. The terminology in this unit is based on an international consensus of proposed and accepted terms.

Myringitis

As the boundary between the external and middle ear, the tympanic membrane rarely develops an isolated inflammation. An inflammation of the tympanic membrane, called myringitis, usually develops in association with otitis externa or otitis media (see below). The principal diseases that can lead to a more or less isolated form of myringitis, such as bullous otitis, were covered in Chapter 10 (see 10.4, pp. 218–223).

Acute Otitis Media

Epidemiology: Acute otitis media is a common disease in infants and small children but can occur at any age. More than 50% of infants experience one or more episodes of acute otitis media during their first year of life. This increases to approximately 80% by 3 years of age.

Pathogenesis: The disease is generally caused by an infection that ascends to the middle ear through the eustachian tube. Bacteria can be isolated from the middle ear in approximately two-thirds of cases. The main causative organisms are Streptococcus pneumoniae, Haemophilus influenzae, and Branhamella catarrhalis.

One-third of cases are probably caused by respiratory viruses. Adenoids are a frequent nidus for middle ear infections in children, even if they are unenlarged.

Factors that increase or reduce risk: Besides craniofacial anomalies, several risk factors for acute otitis media have been identified in children. A previous episode of acute otitis media or the presence of chronic serous otitis media will increase the risk for a recurrence of acute otitis media. Parental smoking is a proven risk factor. Leaving infants and small children at day-care centers is also known to increase the risk and may expose children to a particularly harmful microbiological flora.

On the other hand, extending the period of breast-feeding has been shown to lower the childhood risk of acute otitis media.

Symptoms: Acute otitis media is often preceded by a viral infection of the upper respiratory tract. The initial symptom is a severe earache, which in babies may be manifested by rubbing the affected ear or by nonspecific disease symptoms. Fever is usually present during the first 24 hours. In infants, nonspecific symptoms such as irritability, vomiting, or diarrhea may be the dominant features. Perforation of the tympanic membrane is manifested by aural discharge and by an improvement or resolution of otalgia.

Diagnosis: On physical examination, the mastoid shows no swelling but may be moderately tender to pressure. Otoscopy reveals an opaque, thickened, erythematous, and sometimes bulging tympanic membrane (see Fig. 11.12, p. 239, and Fig. 11.8, p. 236). The tympanic membrane is immobile by pneumatic otoscopy; a Valsalva maneuver should not be performed. Otoscopy may be difficult in children due to cerumen in the ear canal and fussy behavior. The typical features of conductive hearing loss are also present. Generally, a bacteriologic examination is not performed when the tympanic membrane is intact, but it should always be done in patients with a spontaneous perforation or paracentesis. Paracentesis for a bacteriologic examination is indicated in immunocompro-
Recurrent Acute Otitis Media

Definition: The occurrence of five or more acute middle ear inflammations in 1 year, or three inflammations in 6 months, is classified as recurrent otitis media. The middle ear heals between episodes, and there is no effusion present in the tympanic cavity.

Epidemiology: Recurrent acute otitis media is a disease of infants and small children. It occurs in approximately 10% of children who have had a previous bout of acute otitis media.

Etiopathogenesis: The pathogenesis of the individual episodes is basically the same as in acute otitis media (see 112, pp. 234–237), but with a greater prevalence of risk factors.

Symptoms, diagnosis, and course: The individual episodes run the same course as a single bout of acute otitis media. An allergy should be excluded if the history is suspicious for an allergic reaction. The course and complications of the episodes are basically the same as described for acute otitis media.

Differential diagnosis: It is difficult to distinguish recurrent acute otitis media from repeated acute infections in a setting of chronic secretory otitis media. This differentiation can be made only by detecting a normally aerated tympanic cavity after each acute episode. The exclusion of acquired or congenital cholesteatoma is particularly important in these cases, but is often difficult.

Chronic otitis media with a perforated tympanic membrane and recurrent flare-ups can also have a similar clinical presentation.

Treatment: The episodes are treated the same as an isolated bout of acute otitis media. Additionally, there are several actions that can be taken to prevent new recurrences:

- Prophylactic antibiotics are effective but controversial due to the development of resistance.
- Vaccinations against pneumococci can help to prevent new episodes.

Otitis media is caused by different Haemophilus strains than meningitis and therefore is not prevented by Haemophilus vaccinations.

- An adenotomy can decrease the bacterial burden in the nasopharynx and improve eustachian tube function. The placement of a ventilation tube in the tympanic cavity can improve middle ear ventilation.

mised patients, treatment failures, and if complications arise.

Differential diagnosis: Acute otitis media must be distinguished from other forms of otitis. It should be differentiated from otitis externa and from an acute exacerbation of chronic otitis media, particularly in a draining ear with a perforated tympanic membrane.

Course: Spontaneous perforation of the tympanic membrane may occur. After the acute phase of the inflammation has subsided, a residual inflammatory effusion will persist in the tympanic cavity for several weeks, with associated conductive hearing loss.

Complications: Complications are rare. The most common is acute mastoiditis, but any of the otopenic complications described below may occur (p. 247).

Treatment: Nonsteroidal anti-inflammatory analgesics or acetaminophen are given for pain relief. Decongestant nose drops or irrigations may be necessary for relieving nasal airway obstruction.

Unobstructed nasal breathing improves the drainage function of the eustachian tube.

As a rule, antibiotic therapy is started right away though spontaneous improvement may occur in 1–2 days without antibiotics. The antibiotic should be continued for 7–10 days and should be active against the main pathogens listed above. If there is no response or if signs and symptoms worsen within 48 hours, a different antibiotic should be tried. If improvement is unsatisfactory, paracentesis should be performed to obtain a fluid sample for bacteriologic examination.

Prophylaxis: Risk factors should be avoided whenever possible, and the breastfeeding of infants should be prolonged.
Otitis Media with Effusion

Synonyms: secretory otitis media, serous otitis media, mucoid otitis media, glue ear

**Definition:** Otitis media with effusion (OME) refers to an inflammatory effusion behind an intact tympanic membrane that is not associated with acute otologic symptoms or systemic signs. The process may be classified as acute (effusion lasting up to 3 weeks), subacute (up to 3 months), or chronic (more than 3 months).

**In Children**

**Epidemiology:** OME is the most common ear disease in preschool-age children and one of the most common diseases overall. Approximately 3–4% of children have the chronic form. Generally, both ears are affected.

**Etiopathogenesis:** see 11.2, Pathogenesis, pp. 234–237.

**Symptoms:** Symptoms of acute inflammation are usually absent in children. The major symptom is **hearing loss**, but the children themselves rarely report this. Speech and language developmental delay and perceptual impairment may occur in bilateral cases.

**Diagnosis:** The diagnosis is made otoscopically. The tympanic membrane often appears opaque, thickened, and occasionally retracted. Its color may be pale, reddish, yellowish, or bluish, depending on the effusion. Pneumatic otoscopy showed decreased or absent mobility of the tympanic membrane. The tympanogram is a graphic record of tympanic membrane mobility. It may show a flat curve (type B) or occasionally a negative-pressure peak (type C) in mild and acute cases (see 8.4, p. 185). A tympanogram should be recorded to provide a baseline for follow-up.

**Differential diagnosis:** Other diagnoses are rare in children, but a cholesteatoma behind an intact tympanic membrane should be considered if the case has an unusual appearance or runs an atypical course.

**Complications:** The most frequent complication is acute otitis media, from which other otogenic complications may arise.

**Treatment:** The **acute or subacute form** is treated conservatively in an effort to improve nasal breathing and eustachian tube function. This may include the short-term use of decongestant nose drops, moisturizing and hygienic measures, or occasional topical steroids. The value of antibiotic therapy is controversial. Middle ear ventilation can also be improved by having the patient inflate balloons with the nose. Risk factors for otitis media should be curtailed or eliminated as much as possible.

The **chronic form** of OME should be treated surgically if significant hearing loss is present. Paracentesis (incision of the tympanic membrane) provides access for aspirating the effusion, which will immediately restore normal hearing. Ordinarily the incision will close spontaneously in 1–2 weeks, allowing a new fluid collection to form. This can be prevented by inserting a ventilation tube or myringotomy tube (Fig. 11.13): this creates a “chronic” perforation and provides ventilation of the tympanic cavity through the external ear canal. A myringotomy tube will not impair hearing but is associated with a risk of middle ear infection from the ear canal.

Generally, an adenotomy is performed at the time of paracentesis. Various strategies can be applied using various combinations of these measures.

**Prognosis:** **Unilateral** OME has a good prognosis and should resolve within 3 months. Since speech hearing is not impaired, a conservative approach can often be taken in these cases. The detection of air in the middle ear (bubbles, air-fluid level) is also considered a good prognostic sign, as it confirms partial function of the eustachian tube (see also p. 236).

A greatly thickened tympanic membrane and a symptom duration that exceeds three months indicate a protracted course for most cases. A rapid cure is unlikely, and generally these cases should be managed surgically.

**In Adults**

**Etiopathogenesis** (see also 11.2, pp. 234–237): OME is basically the same disease in adults as in children. In adults, however, it is necessary to consider certain causes of eustachian tube dysfunction that are not a factor in children, such as sleep apnea syndrome and tumors of the nasopharynx. Conversely, adenoid hyperplasia and chronic adenoiditis have little if any importance in adults.

**Symptoms:** Patients complain of a clogged or pressure sensation in the affected ear. Pain is rarely present. Some patients complain of a popping or sloshing sound. Unlike small children, adults tend to find the hearing loss and pressure sensation very troublesome.

**Diagnosis:** A **history** of a cold often precedes the complaints. OME is diagnosed **otoscopically** (see also 11.2, pp. 234–237). This examination reveals an opaque tympanic membrane with very poor mobility. A Val-salva maneuver either does not force air into the tympanic cavity or does so only with difficulty. The effu-
sion in the tympanic cavity is easily recognized when air is present (bubbles, level) but is often difficult to appreciate with a complete middle ear effusion. **Hearing tests** show the presence of conductive hearing loss. The Weber test is lateralized to the affected ear in unilateral cases, and the Rinne test is negative. Tympanography yields a flat curve. Pure-tone audiometry indicates an air-bone gap, and otoacoustic emissions are absent.

If a middle ear effusion persists for more than 3 weeks, an endoscopic examination of the nose and nasopharynx should be performed to exclude a tumor.

**Differential diagnosis:** Bullous otitis externa should be considered as a special cause of middle ear effusion. The differential diagnosis also includes other causes of conductive hearing loss with an intact tympanic membrane, such as otosclerosis and ossicular chain disruption. Effusions can also result from barotrauma, a cerebrospinal fluid (CSF) leak from the lateral skull base, or tumors of the temporal bone. A perilymphatic fistula generally leads to inner ear symptoms. The patent eustachian tube syndrome can produce similar complaints, but conductive hearing loss is absent and the tympanic membrane is mobile.

**Complications:** A bacterial, generally ascending infection can develop leading to acute otitis media and its complications. A serous labyrinthitis may develop via the round or oval window.

**Treatment:** Acute and subacute OME are treated conservatively as in children. Emphasis is placed on relieving nasal airway obstruction and treating infections of the nose and paranasal sinuses. The patient should be instructed to perform regular Valsalva maneuvers. If the disease persists for more than 3 months and the tympanic membrane becomes markedly thickened, there is little chance that the inflammatory effusion will resolve any time soon. Surgical treatment is recommended for these cases, and a myringotomy tube should be inserted if conductive hearing loss is present.

**Prognosis:** The prognosis is good in general and in cases with a nonspecific cause. Unilateral middle ear effusions with air bubbles or an air–fluid level almost always have a good prognosis, and the otitis media should resolve within 3 months.

### Chronic Suppurative Otitis Media

**Definition and general information:** A tympanic membrane perforation will usually heal spontaneously in a few weeks. A nonhealing perforation is almost certainly the result of chronic inflammation. Chronic otitis media, then, should be diagnosed in the presence of a **chronic tympanic membrane perforation**, even if there are no active signs of mucosal inflammation. If a specific infection or cholesterol plaque can be excluded, the disease should be classified as **chronic suppurative otitis media**. This process may be **dry**, and thus without active inflammatory signs such as pain, discharge, and swelling of the mucosa. If discharge is present, it is classified as a **wet** or **draining form**. Generally, this means that bacteria have infected the middle ear through the nonintact tympanic membrane. This type of infection may occur **acutely** and resolve quickly, or it may become **chronic** (chronic mucosal suppuration). Thus, the clinical manifestations and otoscopic appearance of chronic suppurative otitis media tend to change over the course of the disease.

**Etiopathogenesis** (see also 11.2, pp. 234–237): The pathogenesis is usually multifactorial. The following factors play a role:

- Chronic inflammation secondary to eustachian tube dysfunction
- Genetic and constitutional factors that affect the healing capacity and resistance of the mucosa
• Special anatomic characteristics of the middle ear spaces such as pneumatization and relative sizes
• The nature, pathogenicity, virulence, and resistance of the infecting organisms

**Symptoms:** Chronic suppurative otitis media presents initially with chronic otorrhea—generally a mucopurulent discharge—through the nonintact tympanic membrane. After the infection clears, the patient has few or no symptoms other than a variable degree of hearing loss. The recurrence of infection may cause pain, but this is not always present. Aural discharge reappears and may be creamy or mucopurulent in the presence of an acute infection. Chronic drainage may consist of odorless, stringy mucus or it may have a fetid smell due to chronic infection with *Pseudomonas* or anaerobes.

**Diagnosis:** The diagnosis is made from the **history** and **otoscopic findings** (Fig. 11.14). Examination reveals a central perforation in the tympanic membrane that does not involve the fibrocartilaginous ring. Often this can be appreciated only in a dry ear. The extent of the perforation can be highly variable. The tympanic membrane and middle ear may show additional features of chronic inflammation such as calcifications, atrophic areas, retractions, or ossicular destruction.

Eustachian tube patency should always be tested and documented (Valsalva maneuver).

In a **draining ear,** the external ear canal contains secretions and may also be inflamed and swollen. Occasionally the perforation is difficult to see due to drainage or general inflammatory changes involving the ear canal and middle ear. A Valsalva maneuver may cause air bubbles to appear in the secretions. A **smear** should be taken for bacteriologic examination.

**Conductive hearing loss** is more pronounced in the draining ear.

**Differential diagnosis:** In cases with an acute infection and pronounced inflammatory changes, chronic suppurative otitis media often cannot be positively distinguished from a **cholesteatoma.** Even imaging studies such as CT are not helpful in this regard. Imaging studies are not indicated in the acute stage and are necessary only for preoperative planning or if complications arise. An expert can differentiate the conditions by otoscopic examination following treatment and resolution of the acute stage. The differential diagnosis also includes otitis externa, specific infections (e.g., with mycobacteria), inflammatory causes such as Wegener granulomatosis, and tumors of the middle ear.

**Course:** A tympanic membrane perforation may be dry for years and cause few if any complaints. Other cases may present with recurrent or persistent otorrhea. This depends in large part on the patient’s diligence in protecting the ear and practicing aural hygiene.

**Complications:** The occurrence of infectious complications such as mastoiditis or abscess formation is rare and atypical in chronic suppurative otitis media. In chronic cases, conductive hearing loss is generally accompanied by the development of cochlear hearing loss, probably the result of a toxic serous labyrinthitis.

**Treatment:** Acute suppurative episodes occasionally require treatment with systemic antibiotics, but this is not consistently necessary. The selection of a specific agent should be directed by antibiotic sensitivity testing.

**Otherwise,** the treatment of otitis media with drainage through the perforated tympanic membrane consists of local measures:

• Repeated, meticulous cleansing and drying of the ear is essential.
• Ear drops that contain ototoxic substances (aminoglycoside antibiotics), if used at all, should be used only to treat acute swelling and only for a brief period (no more than 3 days).
• It is important to provide adequate ear protection while bathing or showering, for example, to keep soap and water out of the ear. This can be done by inserting petrolatum cotton wads or commercially available ear plugs. Otherwise the ear canal should remain clear and should not be packed with cotton.

Adequate ear protection is also important in dry ears to prevent reinfection.

When the ear has been dry for approximately three months, surgical closure of the tympanic membrane can be performed (tymanoplasty, see 11.2, pp. 245–246). Chronic, intractable suppuration requires ablative surgery of the middle ear consisting of a mastoidectomy or a modified radical operation.
Chronic Otitis Media with Cholesteatoma

Definition of Cholesteatoma

**General definition:** A cholesteatoma has two characteristic features:
- Keratinizing squamous epithelium is found in bony spaces at an abnormal location.
- Bone is destroyed through an inflammatory osteoclastic process.
Thus, chronic otitis media with cholesteatoma can be defined as an osteoclastic inflammation of the mucosal spaces in the middle ear. Often there is a coexisting infection, usually with gram-negative and anaerobic bacteria leading to a fetid aural discharge. Infection is not always present, however.

**Types of cholesteatoma:** Cholesteatoma may be congenital or acquired. The **congenital** or **true cholesteatoma** is very rare and is usually found behind an intact tympanic membrane. It can occur anywhere in the temporal bone and will not be discussed here further. **Acquired cholesteatoma** arises in connection with inflammations and ventilation problems of the middle ear. Two forms are distinguished:
- Primary acquired cholesteatoma, known more accurately as **pars flaccida cholesteatoma** (Fig. 11.15), develops from a squamous epithelial pocket in the pars flaccida and initially expands in the epitympanum, where it tends to destroy the lateral attic wall. An unfortunate synonym for this type of cholesteatoma is a “true” cholesteatoma. Other synonyms are epitympanic cholesteatoma, attic cholesteatoma, and attic retraction cholesteatoma.
- Secondary acquired cholesteatoma, known more accurately as **pars tensa cholesteatoma**, originates either from a perforation of the pars tensa with destruction of the fibrocartilaginous ring (marginal perforation) or from a retraction pocket in the pars tensa, usually located in the posterosuperior quadrant (Fig. 11.16). This cholesteatoma initially develops in the mesotympanum but often expands from there into the epitympanum.

Clinical Aspects of Cholesteatoma

**Epidemiology:** A cholesteatoma can occur in any age group but is rare in small children.

**Pathogenesis:** The primary cause of acquired cholesteatoma is probably an impairment of middle ear ventilation (see also p. 234). Eustachian tube dysfunction causes a negative pressure to develop in the middle ear, which may be continuous (due to decreased tubal patency) or transient and recurrent (due to inadequate tubal closure and a negative pressure in the nasopharynx, as in snifﬁng). A retraction pocket forms in the tympanic membrane. The pocket is lined by squamous epithelium that tends to migrate on the tympanic membrane and in the external canal. On entering the middle ear, it causes inflammation and bone resorption. Secondary infection of the squamous debris can further intensify these effects.

**Symptoms:** Acquired cholesteatoma usually presents as chronic otitis media (see p. 240). Less commonly it presents with complications such as labyrinthitis, facial nerve palsy, or intracranial infection.

A **dry, uninfected cholesteatoma** does not cause otalgia or otorrhea. It is manifested clinically by functional deficits. A common initial symptom is aural pressure signifying impaired middle ear ventilation. This is followed by hearing loss and later by potentially severe symptoms that require immediate treatment. These alarming symptoms include facial nerve palsy and signs of vestibular dysfunction such as rotary vertigo and dysequilibrium.

A more common condition, however, is an **infected cholesteatoma with discharge**. It presents as chronic otitis media with otorrhea, generally fetid, and with hearing loss. Pain may supervene with an acute exacerbation, but this is somewhat rare. Again, functional deficits such as facial nerve palsy or vestibular dysfunction may eventually occur. Other possible complications are abscess formation and meningitis.

**Diagnostic work-up:**

**Establishing the diagnosis:** The diagnosis is made by otoscopy (Figs. 11.15–11.17), which typically shows white epithelial debris in a retraction pocket in the attic or in the posterosuperior quadrant of the tympanic membrane. Occasionally, bone erosion is also noted in the posterosuperior canal wall close to the tympanic membrane. With a **dry cholesteatoma**, brownish-black crusts are usually found on the superior canal wall. When **acute inflammatory changes** are present, it may be difficult or impossible to make an accurate interpretation of the otoscopic findings. These cases should first be treated using the same regimen as for chronic otitis media so that an accurate diagnosis can be made (see p. 240).

**Screening for complications:** Hearing tests will generally indicate a conductive hearing loss. The presence of sensorineural hearing loss may mean that a complication has already developed. **Facial nerve function** should be tested whenever a cholesteatoma is suspected (see 14.1, pp. 290–293).

A **labyrinthine fistula** can be confirmed by eliciting the “fistula sign”: a Politzer bag is used to generate a negative and positive pressure in the ear canal. Meanwhile, the eyes are observed with Frenzel lenses. If a
labyrinthine fistula is present, the pressures will evoke rotary vertigo and nystagmus. Imaging studies: Imaging studies do not advance the diagnosis of cholesteatoma. Thin CT slices of the temporal bone can, however, define the extent of bone destruction, detect intracranial complications, and offer presumptive evidence of a labyrinthine fistula. Preoperative CT scans also provide information on the degree of pneumatization of the temporal bone. If complications are suspected, CT scans with contrast-agent administration should be obtained.

Differential diagnosis: Not infrequently, a dry cholesteatoma is mistaken for ordinary cerumen. A cholesteatoma with discharge mainly requires differentiation from chronic suppurative otitis media.

Chronic postinflammatory changes in the tympanic membrane (e.g., retraction pockets) are on a continuum with an actual, acquired cholesteatoma.

Course and complications: Without treatment, it is likely that the bone destruction will progress and complications will arise such as labyrinthine fistula, facial nerve palsy, or intracranial processes.

Treatment: Surgical treatment is necessary due to the bone destruction caused by cholesteatoma. The main goal of the surgery is to eradicate the destructive inflammatory process in the mastoid and tympanic cavity (“radical operation,” 11.2, p.245). A second-line goal is to improve hearing. This can be accomplished with a tympanoplasty, which may be performed concurrently with the ablative surgery or deferred until a later time. Acute inflammatory changes are treated with the same local measures recommended for chronic suppurative

A cholesteatoma develops from a retraction pocket in the pars tensa (usually in the posterosuperior quadrant). Above: otoscopic appearance of the tympanic membrane. Expansion of the retraction pocket in the epitympanum is shown diagrammatically (not visible otoscopically). Below: coronal section through the external ear canal and epitympanum at the level of the retraction pocket and cholesteatoma. Yellow: facial nerve.
11.2 Surgical treatment of chronic otitis media: curative and reconstructive surgery of the middle ear

Ablative middle-ear surgery
In cases of active inflammation like that associated with cholesteatoma, the middle ear must be surgically cleared of disease prior to reconstruction. This involves opening the pneumatized portions of the temporal bone in varying degrees and removing the inflammatory changes.

Mastoidectomy
Principle (Fig. a): The cells of the mastoid and antrum are drilled out through a retroauricular incision. A broad communication is established between the epitympanic space of the tympanic cavity and the mastoid.
Indication: chiefly mastoiditis.

Radical mastoidectomy
Principle (Fig. b): The mesotympanum is opened in addition to the mastoid cells. In the closed or canal wall up technique, the posterior bony canal wall is left intact; in the open or canal wall down technique, it is removed. This operation creates a broad connection between the mastoid, tympanic cavity, and ear canal lumen, called a radical cavity. Today the radical mastoidectomy is almost always combined with a tympanoplasty and the creation of a new tympanic cavity.
Indication: Definitive treatment of cholesteatoma generally requires a radical operation.

Subtotal petrosectomy
Principle (Fig. c): A radical cavity is created, and additional portions of the temporal bone are removed. The surgical cavity then is obliterated with fatty tissue from the abdominal wall. The eustachian tube is closed, and the external ear canal is permanently closed with sutures.
Indication: This operation is rarely performed for inflammatory changes and is used mainly for tumors and post-traumatic conditions (see 15.2 and 15.3, pp. 302-309). It completely disrupts sound conduction across the middle ear.

Reconstructive operations
The permanent changes caused by otitis media can be treated by microsurgical reconstruction. The goal is to seal the tympanic cavity from the external ear canal and restore a largely physiologic mechanism of sound transmission. This type of operation is generally called a tympanoplasty. It usually consists of reconstructing the tympanic membrane (myringoplasty) and the ossicular chain (ossiculoplasty), but occasionally only one of these operations is necessary.
The curative surgery may be combined with a tympanoplasty in the same sitting, or the reconstruction may be staged, depending on the situation. The nature and extent of the reconstruction are tailored to the individual case.

Repairing a tympanic membrane defect: myringoplasty
Principle (Fig. d): The tympanic membrane is reconstructed with a free graft that is usually harvested from the auricle or its surrounding area in the same operation. The most commonly used graft materials are temporalis fascia, perichondrium from the auricle (tragus or concha), or thin cartilage slices. The tissue may adhere to the recipient site by surface tension, or it may be attached with fibrin glue.

Indication: Myringoplasty is necessary in most reconstructive procedures for otitis media and often must be combined with reconstruction of the ossicular chain.

Result: Because myringoplasty is performed with an avascular graft, postoperative healing depends on various local factors. There is an approximately 80% success rate of graft healing with permanent closure of the tympanic membrane, depending on the initial situation.
11.2 Surgical treatment of chronic otitis media: curative and reconstructive surgery of the middle ear (continuation)

Reconstruction of the ossicular chain: ossiculoplasty

Principle: The purpose of ossicular chain reconstruction is to restore sound-pressure transformation from the tympanic membrane to the oval window. The nature and outcome of the reconstruction depend partly on the extent of the ossicular damage. The condition of the stapes is particularly important in this regard. Three main situations are encountered:

- The stapes is intact. The reconstruction establishes a connection between the tympanic membrane (or malleus handle) and the head of the stapes. A synthetic prosthesis that creates this connection is called a partial ossicular replacement prosthesis (PORP) (Fig. e).

- The stapes crura are absent, but a mobile footplate is present. The reconstruction establishes a connection between the tympanic membrane and the footplate. The prosthesis in this case is called a total ossicular replacement prosthesis (TORP) (Fig. f). Both of these reconstructions correspond to a Wullstein type III tympanoplasty.

- The footplate is absent or fixed. A new window is constructed (Fig. g; see also Fig. 11.20, p. 252, and type V below).

Materials: The ossicular chain can be reconstructed with exogenous material, allograft material, or synthetic implants made from various materials such as metal (gold, titanium), plastic, or ceramic. Remnants of the true ossicles are used whenever possible.

Result: As a general rule, the function of the ossiculoplasty (sound transmission) is best accomplished with an intact stapes. Normal values are never achieved when a new window is constructed. Postoperative scarring and further inflammation contribute to a wide range of hearing results.

Wullstein classification of tympanoplasties

The microsurgical technique that is now used in many areas was first developed in 1950 for reconstructive surgery of the middle ear. Two important pioneers were the German otologists Zöllner and Wullstein. In 1952, Wullstein introduced a scheme for classifying the five basic types of tympanoplasty, which is still in limited use today. It is based on the reconstruction of various functional components, i.e.:

- The tympanic membrane (type I = myringoplasty, Fig. d)

- The lever mechanism of the ossicular chain (type II, Fig. h)

- Sound transmission without a lever mechanism (type III = ossicular chain reconstruction, Figs. e, f)

- Sonic shielding of the round window membrane (type IV = creating a small tympanic cavity without restoring sound transmission to the oval window, Fig. i)

- Constructing a new inner ear window (type V, Fig. j)

Since today the lever mechanism of the ossicles is considered to have little importance in sound transmission and operations such as fenestration of the semicircular canal are no longer performed, the classification of tympanoplasties into five “classic” types is often imprecise and confusing. Only types I and III have practical clinical importance.
otitis media. Patients with generalized symptoms should receive systemic antibiotics.

**Otogenic Complications of Otitis**

Otogenic complications may originate from the external ear or middle ear. If a complication arises from the ear canal or auricle, there will almost always be a prior or concomitant infection of the middle ear spaces.

Because otogenic complications are both rare and hazardous, it is better to consider these complications too early than too late (i.e., maintain a high “index of suspicion”).

Otogenic complications are otologic emergencies that should be investigated and treated by a specialist without delay. The patient should be evaluated as soon as possible by audiometry and CT; frequently these studies will demonstrate a need for immediate surgery.

Besides acute complications, otitis can also lead to chronic sequelae that are more difficult to detect. The most important is sensorineural hearing loss due to an accompanying serous labyrinthitis.

This section reviews the typical acute local complications of otitis media. They are most common in association with cholesteatoma but may also occur in the setting of an acute or chronic otitis media.

Local acute complications are an important warning sign indicating the need for immediate (usually operative) treatment.

The earlier surgery is performed, the better the chance of curing the complication or at least preventing further functional deterioration. In rare cases, a chronically infected otitis media may also cause acute systemic complications such as endocarditis (in patients with valve defects), sepsis, or a severe local infection in immunocompromised patients.

**Mastoiditis**

**Definition:** Mastoiditis is an inflammation of the air cells in the mastoid process. Involvement of the mucous membranes of these air cells consistently occurs in acute otitis media and is referred to as associated mastoiditis.

*Mastoiditis in the strict sense* is present when the inflammatory process is focused on the mucous membranes and bony structures of the mastoid. When the temporal bone is well pneumatized, the inflammation may also involve the cells of the petrous bone (petrositis) or zygomatic arch (zygomaticitis).

**Etiopathogenesis:** Mastoiditis usually originates from an infection of the middle ear. It is the most frequent complication of otitis media, but its overall incidence is low. Important pathogenic factors are the degree of mastoid pneumatization, the virulence of the infecting organism, host immune status, and the treatment that has been provided for otitis media. Inadequate antibiotic treatment can predispose to mastoiditis.

Besides infection and abscess formation, an infrequent cause of mastoiditis is an inflammatory destructive process like that occurring in Wegener disease.

**Symptoms:** Patients present clinically with fever and local pain. In infants, mastoiditis or antritis may manifest with malaise, abdominal pain, and anorexia (occult mastoiditis or antritis).

**Diagnosis:** The classic clinical triad consists of:

- A prominent auricle with retroauricular swelling
- Tenderness over the mastoid
- Otorrhea

This classic presentation is somewhat unusual, however. Mastoiditis should be suspected in cases where acute otitis media fails to improve or worsens over a 2–3-week period.

**Otoscopy** reveals the features of acute or subacute otitis media with or without tympanic membrane perforation. The posterior wall of the external auditory canal may be erythematous and swollen (“sagging of the posterior canal wall”). The diagnosis is best established by CT, which can detect other complications as well. Besides clouding of the mastoid air cells and middle ear spaces, scans demonstrate erosion of the mastoid bone structure.

The inflammatory parameters whole blood cell count (WBC), C-reactive protein (CRP), and erythrocyte sedimentation rate (ESR) are markedly elevated.

**Differential diagnosis:** Otitis externa with abscess formation behind the ear can mimic mastoiditis (pseudomastoiditis). Inflamed retroauricular lymph nodes can
also produce tenderness and swelling over the mastoid like that seen with mastoiditis. **Tumors** of the temporal bone such as eosinophilic granuloma, sarcoma, metastases (breast carcinoma, bronchial carcinoma, renal tumors), and lymphomas can mimic the features of mastoiditis.

**Complications:** Other otogenic complications (see below) may arise from mastoiditis, and so the risk of additional complications is increased. The potential complications of mastoiditis are reviewed in Fig. 11.18.

**Treatment:** Treatment generally consists of mastoidectomy (see 11.2, pp. 245–246), which is always combined with culture-directed intravenous antibiotics. Paracentesis and the placement of a myringotomy tube are frequently necessary to decompress the middle ear. Antibiotics without surgery are sufficient only in an early stage of mastoiditis. These cases require intravenous antibiotics and inpatient observation.

### Intracranial Complications

**Meningitis:**

*Etiology:* Otogenic meningitis can result from a clinically overt otitis media, especially with cholesteatoma. It can also arise from an occult process involving the lateral skull base, which may be difficult to detect or undetectable by otoscopy.

With meningitis arising from an unknown focus, especially when the infecting organism is *Streptococcus pneumoniae*, the middle ear and lateral skull base should be considered as potential sites of origin and should be investigated by CT.

*Routes of spread:* Meningitis can result from the spread of a middle ear infection through preformed channels (blood vessels, diploic veins), through the labyrinth (tympanogenic labyrinthitis), through bone gaps caused by laterobasal fractures, or by contiguous spread from infected osteitis or cholesteatoma.

*Symptoms:* Severe headache, fever, clouding of consciousness, and nuchal stiffness become evident within a matter of hours.

The immediate *diagnostic work-up* includes CT scanning of the temporal bone with contrast agent administration and lumbar puncture.

*Treatment* with antibiotics and, if indicated, corticosteroids should be instituted without delay. With processes involving the middle ear or lateral skull base, surgery should be performed after the patient’s general condition has been stabilized. The surgery is both diagnostic to establish the cause of the infection and therapeutic to eradicate the infectious focus and eliminate its route of spread.

*Course:* Otogenic meningitis can lead to inner ear disorders or even bilateral deafness.

### Intracranial Abscesses

**Intracranial abscesses:** These lesions form via the same pathways as meningitis (see above). Otogenic abscesses are generally located in the temporal region or posterior cranial fossa (cerebellar abscess).

The following types are distinguished:

- An *epidural abscess* (between the temporal bone and dura) does not communicate with the subarachnoid space, and so it often produces few symptoms. The dominant signs are those of the underlying otitis media. Otosurgical eradication is often sufficient.
- A *subdural abscess* is located in the subarachnoid space and therefore communicates with the CSF. This lesion is on a continuum with subdural *empyema* and meningitis.

- An *intracranial abscess* leads to systemic signs such as fever, headache, nausea, and vomiting, which may be an expression of raised intracranial pressure. Focal neurologic signs depend on the location of the abscess. A temporal abscess leads to speech disturbances, epileptic phenomena, and cranial nerve palsies (I–VIII). The effects of a cerebellar abscess may include increased intracranial pressure, cranial nerve deficits, balance disturbances and ataxia.

Subdural (subarachnoid) and intracranial brain abscesses should be managed by a combined otosurgical and neurosurgical approach. Effective treatment gen-
Cranial Nerve Deficits

Facial nerve: Peripheral facial nerve palsy (see 14.2, p.294) is the most frequent cranial nerve complication of otitis media. The deficit results from direct inflammatory involvement of the peripheral nerve. The tympanic segment is most commonly affected.

- Even “ordinary” otitis media in children can lead to facial nerve palsy. *Treatment* with antibiotics, combined if necessary with corticosteroids, may be sufficient. With concomitant mastoiditis, it is usually necessary to proceed with mastoidectomy and drainage of the middle ear spaces.

- In adults, cholesteatoma is the most common cause of otogenic facial nerve palsy. Surgery should be performed as soon as possible in these cases following an appropriate audiologic and imaging work-up.

Petrous apex syndrome: The occurrence of trigeminal symptoms and abducens nerve palsy accompanied by otopathologic symptoms is known as petrous apex syndrome or Gradenigo syndrome. Lesions of the petrous apex, usually inflammatory but occasionally neoplastic, cause irritation of the trigeminal nerve root with trigeminal neuralgia or hypesthesia of the trigeminal nerve. They can also irritate the abducens nerve, causing diplopia. Associated deficits involving cranial nerves VII and VIII are often present.

Other cranial nerve deficits: Almost all the cranial nerves may be affected individually or in various combinations by an otogenic intracranial abscess. Deficits most commonly involve the abducens nerve, trochlear nerve, and trigeminal nerve.

 inflammatory sinus thrombosis and otogenic sepsis: Mastoiditis can lead to inflammatory thrombosis of the sigmoid sinus or jugular bulb. With an infected thrombus, dissemination and extension of the thrombus can give rise to otogenic sepsis. The thrombus may also cause intracranial venous outflow obstruction with increased intracranial pressure (especially if there is unilateral hypoplasia of the sigmoid sinus).

Septic phenomena combined with otitis should always raise suspicion of sinus thrombophlebitis.

The *diagnosis* is established by CT or MRI. Treatment generally consists of mastoidectomy with surgical removal of the thrombus.

Labyrinthitis

Tympanic labyrinthitis is described more fully in 12.2, pp. 262-263.

The occurrence of vestibular symptoms in otitis media (vertigo, dysequilibrium, nystagmus) is a warning sign that points to labyrinthitis. An immediate otologic work-up should be performed, and appropriate treatment should be provided.

erally requires the surgical eradication of middle ear disease and intracranial evacuation of the abscess.

Complications: All abscesses can lead to meningitis. This is particularly common with subdural abscesses.
11.4 Injuries and Noninflammatory Diseases of the Middle Ear

This deals with noninflammatory diseases that are confined to the middle ear. They may consist of injuries, ventilation problems, or bone diseases. These conditions may also compromise the function of the inner ear and cause changes involving the entire lateral skull base. There is some overlap with the material presented in Chapter 15 (pp. 298–301), which deals with less localized diseases of the lateral skull base.

Injuries

Injuries to the Tympanic Membrane and Middle Ear

Etiology: Injuries to the tympanic membrane through the external auditory canal may be direct penetrating injuries from a pointed object, or they may be caused indirectly by pressure changes. Indirect injuries are more common. Typical cases involve slaps to the ear or sports-related injuries as in diving. Direct injuries are typically caused by sharp branches, self-cleaning manipulations (cotton-tipped swabs), or welder’s slag burns, in which molten droplets may penetrate the ear canal and drum and enter the middle ear. More violent trauma may affect both the middle ear and inner ear. This is more common with direct injuries.

Pathogenesis: Rupture of the tympanic membrane due to a rise or fall in air pressure depends basically on the rapidity of the pressure change. Atrophic scars in the tympanic membrane are sites of predilection and can perforate more easily than normal tissue in response to relatively small pressure changes. Besides the tympanic membrane, injuries to the middle ear usually affect the ossicles in the form of fractures or dislocations. Injuries that involve the stapes often lead to involvement of the inner ear. The facial nerve is also vulnerable to direct trauma.

Symptoms: A traumatic perforation of the tympanic membrane causes pain of brief duration. The pain may be followed by a clogged sensation and slight bleeding from the ear canal. The patient may feel ear escaping in response to a Valsalva maneuver or nose blowing.

The presence of vestibular symptoms (vertigo, dizziness, nausea) or facial nerve palsy signifies trauma that is not confined to the tympanic membrane.

Diagnosis: A tympanic membrane perforation is detectable at otoscopy, appearing as a slitlike or triangular opening with jagged margins and usually located in an inferior quadrant of the membrane (Fig. 11.19). A hearing test should always be performed. With an injury confined to the tympanic membrane, hearing tests show only mild conductive hearing loss (Weber test lateralized to the affected ear, Rinne test weakly negative or even positive in some cases). Injuries that involve the ossicles, more pronounced conductive hearing loss is found.

Sensorineural hearing loss, nystagmus, or facial nerve palsy are signs of severe middle ear trauma with inner ear involvement. Immediate surgical exploration of the middle ear is advised.

Inner ear involvement may also occur in the form of acute noise trauma (see 12.2, pp. 260–265).

Treatment: The tympanic membrane has a strong capacity for self-healing, and most perforations heal by themselves. They can be covered with various materials such as plastic film or cigarette paper, and any curled edges of the perforation should be straightened. The ear should be protected from water, shampoo, and soap. Isolated tympanic membrane perforations generally heal without sequelae, but conductive hearing loss often persists when middle ear trauma has occurred.

Fig. Traumatic rupture of the tympanic membrane

Fresh tympanic membrane rupture in the right ear. Otoscopy reveals a triangular perforation with hemorrhagic margins.
These cases require later surgical exploration of the middle ear with restoration of sound transmission.

**Barotrauma of the Middle Ear**

**Definition:** A rapid change in air pressure can have acute traumatic effects on the tympanic membrane and middle ear due to a negative pressure in the tympanic cavity (p. 234). The Teed scale classifies barotrauma into five grades of severity based on otoscopic findings:

I  Erythema of the pars flaccida
II  Erythema of the entire tympanic membrane
III  Hematoma of the tympanic membrane
IV  Hematom tympanum
V  Rupture of the tympanic membrane

**Etiopathogenesis:** Barotrauma is caused mainly by compression events associated with airplane landings or diving. When the ambient pressure rises and there is inadequate pressure equalization through the eustachian tube, a negative pressure will develop in the tympanic cavity relative to the environment. This can lead to swelling and bleeding of the middle ear mucosa. These changes, combined with a collapse of the eustachian tube walls, create a further obstacle to tubal opening. With a further rise of ambient pressure, the negative pressure in the middle ear also increases. The tympanic membrane is retracted inward, and the round window bulges into the middle ear. This can result in a perforation of the tympanic membrane or round window membrane.

Inadequate primary opening of the eustachian tube may be caused by any kind of tubal obstruction, but simple rhinitis is the most common precipitating cause.

**Symptoms:** The initial symptom is severe ear pain that is not relieved by a Valsalva or similar maneuver. Conductive hearing loss usually goes unnoticed because of the pain. Vertigo and nystagmus may occur in some cases.

**Diagnosis:** The diagnosis is based on the typical history and otoscopic findings. Otoscopy may reveal tympanic membrane changes, a middle ear effusion, or a rupture of the tympanic membrane, depending on the grade of the injury.

**Differential diagnosis:** The differential diagnosis should include a perilymphatic fistula without actual barotrauma. These cases will generally require surgical exploration of the tympanic cavity. A temporal bone fracture, acute otitis media, chronic secretory otitis media, and bullous otitis externa can have similar otoscopic features.

**Complications:** A tympanic membrane perforation is most hazardous during diving. Cold water can enter the middle ear, causing caloric irritation of the vestibular organ with associated vertigo, loss of orientation, and possible vomiting.

Bacterial infection of the middle ear can also occur through a perforated tympanic membrane or by the tubogenic route based on mucosal swelling and discharge. Involvement of the inner ear via the oval or round window and the formation of a perilymphatic fistula can lead to permanent cochleovestibular dysfunction.

**Treatment:** Paracentesis immediately relieves the negative pressure in the middle ear. In milder cases, an effort is made to reopen the eustachian tube by reducing swelling with medications such as nonsteroidal anti-inflammatory agents or steroids and by optimizing nasal breathing. Nonsteroidal anti-inflammatory drugs (NSAIDs) are given to control pain.

When there is clinical suspicion of an inner ear disorder due to rupture of the round window membrane, the negative pressure should be relieved (paracentesis) and the patient immobilized (bed rest). Surgical exploration of the tympanic cavity may also be required.

**Prophylaxis:** Patients with rhinitis, sinusitis, or otitis media should be prohibited from diving. Predisposed individuals making an airline flight should maintain good nasal breathing by the use of decongestants and moisturizing saline sprays. Patients should also try to equalize pressures frequently and avoid the development of a strong negative pressure in the middle ear.

**Bone Diseases**

**Otosclerosis**

**Definition:** Otosclerosis is a disease of the bony otic capsule in which structural changes in the bone often cause stapes fixation resulting in conductive hearing loss.

**Epidemiology:** The disease occurs predominantly between 20 and 50 years of age. It is about twice as common in women as in men and approximately 10 times more common in whites than in blacks and Asians. Histologically, it is relatively common to find the structural changes described below (in up to 10% of autopsies) in the absence of clinical symptoms.

**Pathogenesis:** The cause of the structural bone changes is unknown. A genetic disposition is presumed to exist based on a positive family history in approximately half of patients. Hormonal changes in women (pregnancy) have also been implicated. Local infection with
the measles virus is assumed to be a precipitating factor. The fact that the abnormality of bone metabolism is limited to the otic capsule is probably related to the special embryonic developmental history of that structure.

The otic capsule affected by otosclerosis initially undergoes localized resorption with a spongiotic structural change. Later the bone at the affected sites becomes sclerotic. A site of predilection is the anterior portion of the oval window niche, where involvement of the footplate and anterior crus of the stapes can cause stapes fixation.

**Symptoms:** Patients notice a slowly progressive hearing loss in one or both ears. With bilateral involvement, one ear is usually affected more than the other. Tinnitus may be present, but vestibular dysfunction due to otosclerosis is rare. The disease may cause cochlear function impairment, however (cochlear form).

**Diagnosis:** Typical signs of conductive hearing loss are accompanied by normal otoscopic findings. The Weber test is lateralized to the affected ear, the Rinne test is negative, and pure-tone audiometry shows that the air conduction threshold is considerably higher than the bone conduction threshold. The bone conduction threshold itself may also be increased in the pure-tone audiogram, especially in the range of 2 kHz (*Carhart notch*).

High-resolution CT scans can occasionally define the otosclerotic foci in the otic capsule, demonstrating them not as sclerotic foci but as circumscribed sites of decalcification. CT is not generally needed to confirm otosclerosis, however, except in the rare cochlear form.

**Differential diagnosis:** The differential diagnosis of conductive hearing loss with an intact tympanic membrane includes the following:
- Middle ear anomalies.
- Ossicular chain disruption due to:
  - Inflammatory changes such as aseptic necrosis of the long process of the incus or tympanosclerosis; the tympanic membrane in these cases usually shows residual signs of chronic inflammation.
  - Traumatic dislocation of the ossicular chain, usually evident from the history; again, typical otoscopic findings are noted.
- Generalized disorders of bone metabolism (see below).

**Complications:** When otosclerotic foci occur at other sites in the otic capsule (“capsular sclerosis”), they can produce a variable degree of cochlear hearing loss ranging to deafness. These additional foci may be caused by toxic products from the bone changes that gain access to the inner ear spaces. Even in cases with stapes fixation, cochlear hearing loss may eventually develop.

**Treatment:** There is no known treatment for the structural bone changes. With cochlear involvement and in other special situations, sodium fluoride may halt the progression of inner ear changes when taken orally over a period of several months.

The rehabilitation of conductive hearing loss may rely on surgical treatment or a hearing aid. Surgical treatment consists of replacing the fixed stapes with a prosthesis that can transmit acoustic vibrations to the inner ear. This requires establishing a connection with the inner ear by creating an opening in the footplate (stapedotomy, Fig. 11.20). The inevitable opening of the perilymphatic space carries a small degree of surgical risk. It can result in peri- or postoperative cochlear hearing loss and vestibular symptoms. While these problems can never be completely excluded, they are very rare when the stapedotomy is performed by an experienced surgeon. By and large, stapes replacement surgery is very successful and is beneficial for the patient.

**Other Bone Diseases**

Systemic bone diseases such as osteogenesis imperfecta, Paget disease, and acromegaly can also lead to stapes fixation. Cochlear hearing loss is also common in patients with osteogenesis imperfecta.
Besides fixation of the stapes footplate, the head of the malleus may be fixed to the bony walls of the epitympanic recess (malleus head fixation). This may occur idiomatically, postoperatively, in the setting of tympanosclerosis, or in patients with anomalies. Treatment consists of surgically reestablishing a vibratory connection for sound transmission (see above and 11.2, pp. 245–246).

Other Diseases of the Middle Ear

Patulous Eustachian Tube

Etiopathogenesis: Normally, the eustachian tube is opened only by the action of the tensor veli palatini muscle. When the amount of fat and connective tissue surrounding the eustachian tube is diminished, a permanently open connection may be established between the nasopharynx and the tympanic cavity. The cause of this may be weight reduction (decreased fat), a reduced venous pressure, or an anatomical change in the cartilage, muscles, or skull base. This type of change may occur in the postpartum period, in association with cachexia or anorexia, after radiotherapy or in acromegaly, to cite a few examples.

Symptoms: The history is characteristic. Women are predominantly affected and complain of aural fullness, vague hearing problems without objective hearing loss, and occasional roaring tinnitus. When questioned, many patients describe autophony, or hearing one's own voice “inside the head” with an echo-like distortion. Lying down or assuming a head-down position improves or relieves the symptoms by raising the venous pressure. Physical or mental effort tends to exacerbate the symptoms. A history of weight reduction is occasionally elicited.

Diagnosis: Otoscopy and hearing tests are normal. Otoscopy or immittance measurements may demonstrate movements of the tympanic membrane synchronous with respirations. Endoscopic examination of the nasopharynx and eustachian tube orifice is necessary to exclude pathology (see 2.1, p. 17).

Treatment: Underlying causes should be treated whenever possible. The condition is harmless in most cases, however, and it is often sufficient to reassure the patient by explaining the cause. Regular, adequate fluid intake may be beneficial.

If the symptoms are persistent and very distressful for the patient, the submucous injection of collagen at the tubal orifice in the nasopharynx may be considered as an option. This carries a subsequent risk of decreased middle ear ventilation, however, predisposing to otitis media with effusion.

Tumors

Tumors of the middle ear are rare. The most common is paraganglioma (synonym: glomus tumor), which presents either with typical otoscopic findings or with functional deficits of the basal cranial nerves (see 15.3, p. 306). Other tumors such as eosinophilic granulomas, carcinomas, carcinoids, plasmacytomas, giant cell tumors, sarcomas, lymphomas, and metastases are usually manifested by nonspecific inflammatory symptoms.

Many of these tumors are misdiagnosed and treated as otitis media for a prolonged period of time.

Because the treatment of tumors often affects the entire lateral skull base, they are discussed more fully in 15.3 (pp. 306–309).
Inner Ear and Retrocochlear Disorders

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12.1 General Clinical Aspects of Cochlear and Retrocochlear Hearing Loss

The function of the cochlea as the basis for hearing is described in 7.2 (pp. 161–163). This unit deals with inner ear disorders that lead exclusively or predominantly to hearing impairment in the form of sensorineural loss. The disorders that lead exclusively or predominantly to vestibular symptoms such as vertigo and dysequilibrium are covered in 13.4 (pp. 282–287). This describes the general clinical features of cochlear dysfunction and explores the distinction between cochlear and neural (retrocochlear) hearing loss.

Introduction

The cause of sensorineural hearing loss most commonly resides in the cochlea. This condition is described as cochlear or sensory hearing loss. A similar type of impairment may be caused by an auditory nerve lesion (neural or retrocochlear hearing loss) or by a lesion of the central auditory pathway (central hearing loss).

It is often difficult clinically to distinguish between a cochlear and retrocochlear cause of hearing loss.

Pathogenic influences may act both on the cochlear sensory structures and on neural structures. Moreover, the causes of sensorineural hearing loss are frequently unknown. Hence the term sensorineural hearing loss, which includes disturbances of neural function at cochlear sites such as the synapses or ganglion cells. Cochlear disturbances may also be associated with disturbances of the vestibular system in patients with labyrinthine damage or systemic disease.

Symptoms of an Inner Ear Disorder

The clinical hallmarks of an inner ear disorder are hearing impairment, tinnitus, and vestibular symptoms in the form of dysequilibrium and vertigo (see 13.2 and 13.3, pp. 275–281). These symptoms may be isolated or combined.

Hearing Impairment

Hearing impairment with a cochlear cause usually involves a diminished hearing ability, or hearing loss (synonym: hypacusis), which is generally accompanied by a distortion of hearing (synonym: dysacusis). The latter type of hearing impairment is perceived as imprecise comprehension and distorted sounds. Hearing impairment due to an inner ear disorder can vary in its duration of onset and natural history:

- An abrupt, unilateral hearing impairment is also known as sudden hearing loss. It is described as symptomatic if the cause can be ascertained. Often, however, the cause of sudden hearing loss is indeterminate and the condition is described as idiopathic sudden hearing loss (12.3, pp. 267–269).
- More commonly the hearing impairment takes a slowly progressive course, which is typical of hereditary causes, for example, and also of presbyscusis (hearing loss with ageing).
- Stable hearing disorders may occur after the cause (e.g., drugs or noise) has been eliminated.

Tinnitus

Definition and causes: Tinnitus is an auditory sensation that occurs in the absence of an external acoustic or electrical stimulus and has no subjective information content. Tinnitus is a general, nonspecific symptom of an auditory system abnormality (“noise in the system”) whose cause can range from an obstruction of the external auditory canal to a brain tumor. Most cases, however, are caused by a disturbance of peripheral sensorineural structures (Table 12.1).

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<th>Table 12.1 Tinnitus: classification and causes</th>
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<tr>
<td>Classification</td>
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<td>“Subjective” tinnitus</td>
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<td>Conductive tinnitus</td>
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<td>Sensorineural tinnitus</td>
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<td>Central tinnitus</td>
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<td>“Objective” tinnitus</td>
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<td>Vascular tinnitus</td>
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<td>Myogenic tinnitus</td>
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Traditionally, a distinction is drawn between “objective” and “subjective” tinnitus. This distinction is flawed, however, because tinnitus is a symptom similar to pain or pruritus, it is always subjective. As a result, tinnitus cannot be measured objectively (see also 8.3, p. 183). Occasionally, however, a patient may hear vascular or muscular sounds originating in the body, and these sounds can be amplified to objectively audible levels with a stethoscope or microphone probe. This perception is then referred to as “objective” tinnitus.

Tinnitus that adversely affects the quality of life has a prevalence of approximately 0.5% in the general population. Tinnitus is associated with sleep disturbance in approximately 4% of these persons, and the symptom of tinnitus can become a disease in these patients. The disease is compensated if it does not compromise the activities of daily living. It is decompensated if it causes significant disability, suffering, or suicidal ideation.

Tinnitus is classified by its duration as acute (up to 3 months), subacute (4–12 months), or chronic (longer than 1 year).

The treatment of tinnitus is challenging. Intravenous lidocaine is of temporary benefit in approximately 70% of cases, underscoring the frequent neural component of the condition.

Acute tinnitus can be treated with circulatory stimulants or oxygen therapy, although it has not been proven that these measures are effective. Aside from anecdotal reports, the efficacy of medical treatments in general has not been positively established.

Addictive treatment agents should be avoided, therefore.

Chronic tinnitus with decompensation is best managed with a combination of behavior modification and psychotherapeutic measures. The hyperacusis that often coexists with tinnitus can be treated acoustically with noise generators that are worn like hearing aids (noisers, maskers). The goal of this “tinnitus retraining therapy” is to integrate the tinnitus into the acoustic perception of the patient in a way that is no longer disturbing or distressful.

Diagnostic Evaluation

History

The subjective symptoms of cochlear or auditory-nerve damage described above are nonspecific and do not permit an etiologic diagnosis. Often only a detailed history can provide evidence of etiologic factors. The following points in particular may have etiologic significance:

• Former and current noise exposure during work or leisure activities, also previous acute noise trauma

• Cranial trauma, which may be associated with acute noise trauma or with a direct contusion of the cochlea

• A prior history of chronic otitis media, which may be an indirect cause of cochlear damage

• Family history of hearing disorders

• Current or previous use of ototoxic medications or exposure to toxic substances

The actual handicap imposed by a hearing disorder can be appreciated only from the patient’s history, as it depends only partly on the degree of hearing impairment that can be measured by audiometry. Occupational and personal demands on hearing, the personality itself, and social behavior contribute just as much to the actual hearing handicap.

It is particularly important to recognize a tendency toward social withdrawal based on the hearing disorder.

Because sensorineural hearing disorders are made noticeable by difficulties in speech recognition, especially in a noisy environment, people with hearing loss tend to avoid acoustically challenging but socially important situations such as group discussions and large gatherings in order to avoid misunderstandings, ridicule, and embarrassment. This can lead to social isolation and, when combined with comprehension difficulties, to mistrust.

Clinical Examination

Inspection and otoscopy will show no abnormalities in patients with an isolated lesion of the cochlea or auditory nerve. With a moderate, symmetrical disturbance, suprathreshold tuning fork tests will also be normal. With a unilateral or asymmetrical disturbance, the Weber test is laterализed to the better-hearing ear and the Rinne test is normal.

The whispered speech test usually shows significant impairment in the ability to repeat whispered numbers—especially those with sibilants (e.g., “66”) due to the high frequency spectrum of these numbers. Numbers spoken at a normal conversational level have a considerably lower frequency spectrum and are easier to understand.

Audiometry

The following findings are noted in the typical audiometric profile of a sensorineural hearing disorder (Fig. 12.1):

• The hearing threshold in the pure-tone audiogram is increased for air and bone conduction. This generally occurs first and most conspicuously in the high-frequency range (“high-tone loss”).
- Sound conduction is not impaired—i.e., the air and bone conduction thresholds are equal in the pure-tone audiogram, and immittance measurements are normal.
- Otoacoustic emissions are absent because the function of the cochlear amplifier is impaired.
- Speech recognition is abnormal, particularly in background noise, and there is loss of speech discrimination. Speech is heard, but it is misunderstood or not understood. In monosyllabic word tests (see 8.3, p. 180), all of the words are not understood even at highly amplified levels, and higher levels may even be associated with poorer comprehension.

All of these findings cannot reliably distinguish between a cochlear and retrocochlear hearing disorder. Even the suprathreshold audiometric tests that were once widely used are not effective for this purpose (see 8.3, p. 178). A reliable differentiation can be made audiometrically by eliciting the auditory brainstem response (ABR, brainstem auditory evoked potentials; see 8.4, p. 184). This test is of limited value for greater degrees of hearing loss, however, and imaging procedures (MRI) should be used.

**Imaging Studies**

High-resolution thin-slice computed tomography (CT) scans of the temporal bone can detect changes in the bony labyrinth like those associated with malformations, trauma, or bone diseases such as osteogenesis imperfecta (Fig. 12.2), Paget disease, or advanced otosclerosis.

MRI is a particularly good modality for examining the auditory nerve. It is the method of choice for diagnosing a retrocochlear lesion in patients with severe hearing loss.

MRI can only detect structural changes in the auditory nerve or brainstem, however, and so a normal MRI study does not exclude neural dysfunction.
CT of the inner ear in osteogenesis imperfecta

Thin axial CT slice of the petrous bone demonstrates the rarefied bone structure around the cochlea and vestibular labyrinth. The patient is deaf in both ears (compare with normal cochlea in Fig. 7.4, p. 157).

Retrocochlear Disorders

Etiology: The main known causes of retrocochlear hearing impairment are tumors of the internal auditory canal and cerebellopontine angle, compression of the auditory nerve by vascular loops, changes in the auditory nerve or its entry zone due to multiple sclerosis, and other inflammatory processes. Cerebellopontine angle tumors are discussed in 15.3 (see pp. 306–309).

Diagnosis: In many cases, a sensory cochlear disorder cannot be distinguished from a retrocochlear neural disturbance of inner ear function on the basis of clinical findings. Moreover, combined lesions are not uncommon. Generally speaking, however, cochlear disturbances are predominant. It is difficult to detect a bilateral retrocochlear component of sensorineural hearing loss, but a unilateral retrocochlear disorder is of particular clinical importance.

In patients with unilateral sensorineural hearing loss, a retrocochlear disorder should be excluded by ABR testing (Fig. 12.3, pp. 187–191) or MRI. The subjective complaints associated with a retrocochlear disorder are basically the same as those of a cochlear disorder (hearing impairment, tinnitus, and possible vestibular symptoms). The patient may exhibit particularly poor speech recognition, although this is a nonspecific sign and may be subtle when the symptoms are unilateral.

Audiometry exhibits the typical features of sensorineural hearing loss. A speech hearing loss that is quite pronounced in relation to the pure-tone audiogram may direct attention to a retrocochlear disorder. Because lesions of the auditory nerve cause a change in nerve conduction velocity, a disturbance of auditory temporal resolution is often noted as an early sign of a retrocochlear disorder. Pure-tone audiometry is performed using tones of longer duration, which are still clearly audible even if temporal resolution is impaired. Temporal resolution is essential for speech recognition, however, which involves the perception of very rapid spectral changes.

Since the conduction velocity of the auditory nerve is abnormal, the measurement of ABR latencies is the most sensitive and useful audiometric test for detecting a retrocochlear disorder (see pp. 187–191). Most of the lesions mentioned above can be detected by MRI, and therefore this study should be included in the work-up of retrocochlear impairment.

Auditory evoked brainstem potentials in a patient with a left-sided retrocochlear hearing disorder. The right side shows normal ABR potentials with a normal latency. On the left side, only potential I is clearly defined. The potentials were evoked with a brief click stimulus approximately 90 dB above the hearing threshold.

[Diagram of auditory brainstem response in retrocochlear disorders]
12.2 Cochlear Hearing Loss with a Known Cause

The cause of an inner ear disorder often remains unknown. Some etiologic factors such as inheritance, noise exposure, inflammations, and toxic substances are well known, however. Even so, it can be difficult clinically to relate specific causal factors to sensorineural hearing loss.

**Hereditary Sensorineural Hearing Loss**

**Classification:** Hereditary inner ear disorders are clinically and genetically diverse. A basic distinction is drawn between syndromic and nonsyndromic hereditary inner ear disorders. **Syndromic hereditary hearing loss** is usually present at birth and is described in 9.1 (p. 198).

**Hereditary, nonsyndromic inner ear disorders** are classified into two forms based on the time of onset of the hearing loss:

- A congenital form: see Causes and Effects of Pediatric Hearing Disorders, pp. 198–201.
- A later-onset form: the inner ear disorder is manifested after birth (occasionally not until adulthood).

**Incidence:** Hereditary inner ear disorders are a frequent cause of cochlear hearing loss. It is estimated that approximately one-third of all cases of sensorineural hearing loss have a genetic cause or a contributing genetic cause. The later-onset forms are significantly more common than the congenital forms.

**Inheritance:** Hereditary inner ear disorders may have a dominant, recessive, sex-linked, or mitochondrial mode of inheritance. Whereas sensorineural hearing loss that is already present at birth usually has an **autosomal-recessive** inheritance, the later-onset forms tend to be **autosomal-dominant.** **X-linked** hearing disorders predominantly affect males, often begin at puberty, and may present as a mixed sensorineural and conductive hearing loss. Hearing disorders that are transmitted by **mitochondrial** inheritance are progressive and are often associated with muscular diseases or other symptoms of mitochondrial pathology.

**Symptoms:** Nonsyndromic, hereditary hearing disorders are reasonably symmetrical—i.e., the hearing loss affects both ears to an approximately equal degree and over the same range of frequencies. Hereditary hearing loss of later onset may progress intermittently with recurrent attacks of sudden hearing loss (especially in children), or there may be a gradual progression of hearing loss over time (most common in adolescents and adults).

This deals with forms of cochlear hearing loss that occur predominantly in adults and have a known etiology. The forms and effects of congenital and early childhood cochlear hearing loss were reviewed in 9.1 on pp. 198–201.

**Diagnosis:** A positive family history can be elicited in cases with a dominant mode of inheritance. Audiometry exhibits the typical features of cochlear hearing loss (see p. 257). The pure-tone audiogram is nonspecific. A saucer-shaped curve with greater hearing loss at the middle frequencies does suggest hereditary hearing loss, but the audiogram may also show other configurations such as a low-tone or high-tone loss. There have been increasing reports of gene loci at which changes can lead to hearing impairment. Some genetic mutations, as in the connexin gene, can be detected with laboratory tests.

**Treatment:** Treatment is limited to the prevention of further acquired damage and to rehabilitation with hearing aids and other assistive devices. Genetic counseling may be advised.

**Noise-Induced Hearing Loss**

**Pathogenesis and classification:** Excessive noise exposure can cause direct mechanical trauma to the delicate structures of the cochlea. When the exposure is prolonged, metabolic injury is added to the traumatic changes. The nature and extent of the damage depend on the acoustic energy of the injurious noise and the duration of exposure. The higher the energy (i.e., the level or loudness level) and the steeper the initial up-slope of the energy, the greater the likelihood that mechanical trauma will occur. Several types of noise-induced hearing loss are distinguished on the basis of exposure time. They are not separate and distinct entities but are on a continuum:

**Acute acoustic trauma:** The ear is exposed to a sudden, intense sound event of short duration. The loudness level exceeds 140 dB SPL, and the duration of the pressure rise is very short—less than 1.5 ms. The most frequent noise source is a gunshot, but acute acoustic trauma may also be caused by airbags, firecrackers, or aerial fireworks.

**Blast injury:** In this type of injury the ear and body are subjected to the pressure wave from an explosive blast. Again, the loudness level exceeds 140 dB SPL, but the duration of the pressure rise is longer than with gunshot trauma (> 2 ms) and the frequency spec-
Fig. Safe weekly exposure times to noise

Relationship between noise exposure measured in dB(A) and the exposure time that can cause measurable hearing loss in less than 5% of normal-hearing subjects. Because the sound level is measured logarithmically, an increase of 3 dB corresponds to a 50% reduction of exposure time.

- **Acute noise-induced hearing loss**: The ear is exposed to high levels of continuous or intermittent (pulsed or impact) noise lasting from a period of seconds (e.g., jet engine) to hours (e.g., rock concert). The higher the sound pressure level, the shorter the exposure time needed to produce injury.
- Acute noise-induced hearing loss is often reversible or partially reversible. Little attention may be given to the symptoms of slight deafness and tinnitus. The most frequent sources are loud power tools, musical performances, or engine noise.
- **Chronic noise-induced hearing loss**: This is an irreversible cochlear hearing loss whose severity depends on the noise level, exposure time, and individual factors. Exposure to levels below 85 dB(A) for 8 hours per workday is considered safe (Fig. 12.4).
- While potentially harmful noise exposure once occurred mainly in the workplace, noise-induced hearing loss is now a greater hazard in young people exposed to music and other noises that are associated with leisure activities (socioacusis).

- **Symptoms**: The typical symptoms of noise-induced inner ear damage are a muffled sensation and tinnitus. These symptoms occur immediately in acute events and develop more gradually in response to less intense noise exposure. The symptoms improve after the noise is withdrawn and may disappear completely after less intense exposure.
- **Irreversible, chronic noise-induced hearing loss** presents with the typical features of sensorineural hearing loss—increase, loss of speech discrimination, especially in background noise, and tinnitus. The tinnitus is constant, and the annoyance factor is variable and does not correlate with the degree of hearing loss.

- **Diagnosis**: The audiometric hallmark of noise-induced hearing loss is a drop-off in the hearing threshold between 3 and 6 kHz (Fig. 12.5). Initially, a notch appears in this frequency range, indicating that the hearing threshold is better at 8 kHz than at 4 kHz. In cases of **acute noise exposure**, the threshold curve declines over a relatively broad frequency range immediately after the exposure, but the effect is still most pronounced in the 3–6-kHz range. This disturbance, called a temporary threshold shift (TTS), generally resolves within a few hours.
- With **sustained noise exposure**, hearing undergoes a permanent threshold shift (PTS) without recovery. Over time the patient develops an irreversible, progressive high-tone hearing loss that may also affect the middle frequencies around 1–2 kHz.
- The pitch of the tinnitus is often perceived at the frequencies of the declining pure-tone threshold curve.

- **Complications**: The tinnitus may become decompensated, resulting in significant psychological distress.

- **Treatment**: There is no known effective treatment for noise-induced hearing loss. It is important to avoid further damage to the inner ear from noise or other influences. Acute exposure can be treated with measures to improve metabolic conditions, the microcirculation, and the oxygen supply to the inner ear. Corticosteroids are also recommended.

- **Prophylaxis**: The lack of an effective treatment for noise-induced hearing loss underscores the importance of prevention. The most effective **primary prophylaxis** is the reduction of noise emissions, which is an engineering and legislative concern. Hearing protection should be con-
Persistently worn in situations where there is exposure or potential exposure to injurious noise. The physician should support prophylaxis by recommending the use of hearing protection and the reasonable control of noise levels in the home (e.g., music).

For secondary prophylaxis, people with occupational noise exposure should be regularly screened so that any noise-induced hearing loss can be detected at an early stage and appropriate measures can be taken such as hearing protection or a change of workplace.

**Traumatic Injury to the Inner Ear**

**Synonyms:** labyrinthine concussion and contusion, labyrinthine trauma

**Etiology and pathogenesis:** The inner ear is vulnerable to direct or indirect trauma, with most injuries occurring in association with acoustic trauma. The damage to the inner ear may be functional (labyrinthine concussion) or structural (labyrinthine contusion):

- Temporal bone fracture (see 15.2, pp. 302–305): the labyrinth may lie within the fracture zone.
- An impact to the skull that does not directly injure the temporal bone can still cause strong accelerating and decelerating forces to act on the labyrinth, causing bleeding or other structural and functional changes.
- Barotrauma (perilymphatic fistula, see 13.4, p. 286): the inner ear is damaged by pressure changes acting through the round or oval window.

**Symptoms:** Initially there are nonspecific vertiginous complaints and hearing impairment that are often obscured by the associated symptoms of craniocerebral trauma or other injuries. Later testing often indicates only a hearing disorder, which is often more pronounced in one ear than the other. Tinnitus may be present.

**Diagnosis:** The diagnosis is based on the history and associated clinical and audiometric findings. Generally, a shallow notch appears in the 3–4-kHz range as a manifestation of additional acute acoustic trauma. Vestibular findings (see 13.2, pp.275–279, and 13.3, p.280) can also help to advance the diagnosis.

Structural injuries to the labyrinth cannot be detected directly; they can only be presumed on the basis of functional disturbances.

**Differential diagnosis:** In patients with concomitant vertiginous symptoms, it is difficult to distinguish traumatic lesions of the inner ear from central vestibular and auditory lesions like those associated with a brain contusion.

**Treatment:** No specific treatment is available. The ear must be protected from additional harmful influences such as noise and ototoxic antibiotics.

**Labyrinthitis**

**Pathogenesis and classification:** Labyrinthitis can result from an infection or other inflammatory process affecting either the labyrinth itself or its surroundings. There are three routes by which an infection or inflammation can spread to the labyrinth:

- **Tymanogenic labyrinthitis:** An infection or inflammation of the middle ear may be transmitted through the round or oval window:
  - Acute toxic form (serous labyrinthitis): The labyrinth itself is not infected; presumably it becomes inflamed by substances that are released in the middle ear (e.g., inflammatory mediators or bacterial toxins). This pathogenic mechanism is probably also involved in viral tymanogenic infections (e.g., in bullous otitis externa).
  - Acute suppurative form: A bacterial infection of the middle ear spreads to involve the labyrinth.

  This form is rare but very dangerous and often results in permanent loss of hearing and vestibular function. It can lead to meningitis.

  - Chronic labyrinthitis: This may eventually be manifested as inner ear damage. Chronic otitis media is among the possible causes.
Meningeal labyrinthitis: The labyrinth may be infected bilaterally (often with *Streptococcus pneumoniae*) from the intracranial space, possibly through a patent cochlear aqueduct. In these cases the labyrinthitis develops as an accompanying feature of meningitis, predominantly affecting infants and small children but also occurring in adults. The process may lead to complete deafness and calcification of the labyrinth.

Hematogenous labyrinthitis: Viruses and bacteria can infect the labyrinth by the hematogenous route, resulting in hearing loss and dysequilibrium. Typical causative organisms are mumps and measles viruses, human immunodeficiency virus (HIV), cytomegalovirus (perinatally), and spirochetes (syphilis, borreliosis). Other hematogenous bacterial infections of the labyrinth are rare.

**Symptoms and diagnosis**: Labyrinthitis is manifested clinically by cochlear hearing loss, tinnitus, and vestibular symptoms (vertigo, dysequilibrium, nystagmus).

The appearance of vestibular symptoms in a patient with otitis media is a warning sign of labyrinthitis and necessitates an immediate otologic work-up and appropriate treatment.

Audiometry shows signs of sensorineural hearing loss. With tympanogenic labyrinthitis, high-resolution CT scans of the temporal bone are generally necessary to exclude a labyrinthine fistula. Cerebrospinal fluid (CSF) sampling should be performed if there is the least suspicion of meningitis. Serologic screening for syphilis, borreliosis, and labyrinthogenic viruses should be performed in all patients with suspected labyrinthitis due to an unknown cause.

**Treatment**: Tympanogenic labyrinthitis in acute otitis media requires careful decompression of the middle ear with a myringotomy tube, either alone or combined with a mastoidectomy (see 11.2, p. 245). A labyrinthine fistula also warrants immediate surgical treatment (see p. 286).

A bacterial infection is treated with high doses of antibiotics, usually administered intravenously. An antibiotic should be used that will enter the subarachnoid space. Corticosteroids are probably beneficial in cases with a viral or toxic etiology.

**Prognosis**: Some recovery of inner ear function is possible, but most patients are left with permanent residual damage. Recovery from severe functional deficits is rare.

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**Ototoxicity**

**Etiology and pathogenesis**: Toxic damage to the inner ear may affect both cochlear and vestibular functions, or each of the functions may be affected more or less in isolation. The toxic effect may be reversible or irreversible.

The causative agents of ototoxic effects may be *endogenous* (see Metabolic Disorders, p. 265) or *exogenous*. The principal exogenous toxins are listed in Table 12.2. A great many drugs have been named in isolated reports as producing ototoxic effects.

Ototoxic effects are generally symmetrical.

An exception to symmetrical ototoxicity is the local toxic effect produced by substances in the tympanic cavity. Examples are the local application of ototoxic medications like those contained in ear drops (exogenous toxin) and effects from the local endogenous toxins that form in chronic otitis media.

**Symptoms and diagnosis**: The *cochlear disorder* is frequently accompanied by tinnitus, which may be the initial presenting symptom. Generally, the degree of hearing loss is similar on both sides. Audiometry displays the typical features of cochlear hearing loss (see 12.1, p. 257). Since *vestibular input is symmetrical*, there is no labyrinthine imbalance and consequently no nystagmus. The typical vestibular symptoms consist of oscillosia and dysequilibrium (see 13.2, p.277).

**Differential diagnosis**: Neurotoxicity requires differentiation from labyrinthine toxicity. Neurotoxic damage to cranial nerve VIII produces symptoms similar to those of a labyrinthine disorder. Many ototoxic substances are also neurotoxic, often making it difficult to distinguish specific inner ear damage from diffuse neurologic injury. For example, the central neurotoxic effect of alcohol is usually predominant, but alcohol also has a specific effect on the labyrinth.

**Ototoxic Drugs**

Aminoglycosides damage the cochlear hair cells. Streptomycin and gentamicin are more likely to affect the vestibular system, whereas amikacin tends to affect the *cochlea*.

The critical factor in terms of ototoxicity is the concentration of the agent in the perilymph and endolymph, which correlates only indirectly with the serum concentration.

A number of other factors, some genetic and metabolic, appear to influence the ototoxicity of aminoglycosides. Impaired renal function is of particular significance.
Some ear drops contain aminoglycosides as antibiotics, and consequently these products should not be used in patients with a perforated tympanic membrane. Applied locally in the middle ear, they can exert a toxic effect on the inner ear by way of the round or oval window. This effect is applied therapeutically in the treatment of Ménière’s disease.

Patients often notice a high-pitched tinnitus as the initial subjective symptom. Tests reveal the typical signs of cochlear hearing loss, which affects the higher frequencies first, followed by the lower frequencies.

**Cisplatin**: The ototoxic effect of the cytostatic drug cisplatin is directed mainly against the hair cells. Cisplatin also has neurotoxic properties. The hearing loss associated with cisplatin use is a typical *cochlear high-tone loss*, which may be reversible to some degree. Deafness is rare. Particular attention should be given to renal function impairment.

Other platinum-based cytostatic drugs such as carboplatin appear to be less ototoxic, although hearing loss may still occur. **Loop diuretics** such as furosemide have an ototoxic effect on the stria vascularis at high doses and cause a decrease in the endocochlear potential. This leads to a *hearing loss at all frequencies*, which is generally reversible.

**Quinine**: The use of quinine has long been linked to the development of *tinnitus, hearing loss, and dysequilibrium*. The ototoxic effect, consisting of a hearing loss at *all frequencies*, is generally reversible. Other antimarial drugs such as quinidine, chloroquine, and mefloquine can produce similar adverse effects, but less frequently than quinine.

**Salicylates**: Salicylic acid derivatives produce an oblique, dose-dependent ototoxic effect that is always reversible. The results are a roaring tinnitus and a mild to moderate impairment of *cochlear function at all frequencies*.

**Prophylaxis**: If the use of potentially irreversible ototoxic medications is planned, the patient’s cochlear function should first be tested if at all possible. Useful tests are pure-tone audiometry, high-tone audiometry, and otoacoustic emissions.

Inner ears that have preexisting damage are generally more susceptible to ototoxic effects. Very strict selection criteria should be applied for using ototoxic medications in these patients.

During treatment with ototoxic medications, special attention should be given to dosage, renal function, and adequate hydration. Regular measurements of serum levels are helpful. For secondary prophylaxis (= early detection of disease), inner ear function should be tested by high-tone audiometry, otoacoustic emissions, or pure-tone audiometry. At the latest, these tests should be conducted when the patient first complains of tinnitus, vestibular problems, or subjective hearing difficulties.
Other Acquired Inner Ear Disorders

Immunologic Causes of Inner Ear Disease

Recognized autoimmune syndromes can lead to inner ear disease. These disorders present clinically with bilateral, often asymmetrical cochlear hearing loss that may have a different time of onset in each ear and generally takes a rapidly progressive, occasionally fluctuating course. There may be accompanying symptoms of vestibular dysfunction. 

Cogan syndrome is invariably associated with a bilateral, progressive loss of cochleovestibular function along with interstitial keratitis. The disease is rare, occurring predominantly in young women in their teens or twenties. Therapy consists of immediate treatment with high-dose corticosteroids, which may be supported with immunosuppressants such as cyclophosphamide or cyclosporin. 

Wegener’s granulomatosis and other forms of vasculitis can also lead to immune-mediated inner ear disorders. Recurrent polychondritis can affect inner ear function in addition to the auricle. 

Primary autoimmune diseases arising in the inner ear itself are presumed to occur, but at present they cannot be confirmed clinically or by specific laboratory tests. A trial of high-dose corticosteroids is recommended in clinically suspected cases.

Decreased Blood Flow

An acute blood flow deficit in the vertebrobasilar territory (labyrinthine infarction) can lead to acute, severe, usually unilateral disturbances of inner ear function. A selective decrease of blood flow may occur in the cochlear branch or labyrinthine artery. These cases do not display associated brainstem symptoms. The degree to which a chronic reduction of blood flow like that associated with abnormal blood pressure regulation, generalized atherosclerosis, diabetes mellitus, or nicotine abuse can lead to functional impairment of the inner ear is uncertain and is difficult to establish as causative in any given case. Epidemiologically, statistical correlations have been found between cochlear disorders and the factors cited above.

Metabolic Disorders

Like chronic circulatory insufficiency, metabolic changes are known to be associated with hearing loss and other labyrinthine pathology. The link to hyperlipidemia is very well established. For this reason, the correction of such disorders is definitely advised in patients who present with unexplained chronic hearing loss. It is also believed that uremia can be a direct cause of labyrinthine damage.

Cochlear Otosclerosis

Otosclerosis can lead to progressive cochlear hearing loss when it involves the labyrinthine capsule outside the oval window niche (see 11.4, pp. 251–253). The damage is most likely caused by toxic metabolic products from the diseased bone.
12.3 Cochlear Hearing Loss with an Unknown Cause

An effort should always be made to establish the cause of sensorineural hearing loss. The known causes were discussed in the preceding. In the majority of cases, however, even a thorough diagnostic work-up will be unable to establish a definitive cause for sensorineural hearing loss. The hearing loss in these cases is often assigned to a typical clinical presentation that serves as a “diagnosis,” and these presentations are the subject of the present. It should be kept in mind that these “diagnoses” can have various unknown causes and are generally made by a process of exclusion. Presbycusis, sudden sensorineural hearing loss, and chronic progressive idiopathic hearing loss are examples of diagnoses with an unknown cause.

Presbyacusis

**Definition:** Presbycusis is an age-related, apparently idiopathic and symmetrical sensorineural hearing loss that affects persons over 50 years of age. High frequencies are affected more than low frequencies, and speech recognition is impaired more than pure-tone hearing, especially in a noisy environment.

**Occurrence and incidence:** Initial signs of presbycusis may appear as early as the fourth decade. Approximately one-third of persons over 65 years of age (50% of men, 25% of women) show significant hearing loss averaging 35 dB or more by pure-tone audiometry.

**Etiology and pathogenesis:** Presbycusis is a collective term that encompasses various disorders of the auditory system (peripheral and central) and various etiologic influences (multifactorial etiology). Because the hearing loss is nonspecific, a purely age-related, endogenous hearing loss is virtually indistinguishable from hearing loss that has contributory exogenous causes:

- **Ageing processes** at the cellular level (hair cells and neurons) and organ level (basilar membrane, organ of Corti, stria vascularis) can have various manifestations. Postulated changes include autointoxication by metabolic products, a loss of elasticity due to connective-tissue changes, and an increase in mass due to nondegradable deposits in the cells and organs.

- **An endogenous genetic predisposition** is probably also a factor in many patients. Presbycusis in these cases is on a continuum with hereditary sensorineural hearing loss of late onset.

- **A cumulative exposure to exogenous factors that can damage hearing** also has causal significance with ageing. These factors include noise, middle ear diseases, toxic substances, as well as tobacco use (socioacusis).

One feature of presbycusis is a disturbance of the sensory elements in the cochlea. This is unlike age-related visual deterioration, or presbyopia, which mainly involves a disturbance of the transmission system (decreasing elasticity of the lens). A more valid ophthalmologic counterpart to presbycusis is age-related macular degeneration.

**Symptoms:** Patients initially complain less of hearing loss than of problems with recognizing speech in background noise. Tinnitus may also be present.

**Diagnosis:** Otoscopy and impedance audiometry yield normal findings. Pure-tone audiometry demonstrates a symmetrical sensorineural hearing loss that exceeds the normal degree of age-related hearing loss, which is defined in the pure-tone audiogram by international standards (ISO 1999, Fig. 12.7). Pure-tone audiometry mainly indicates a high-tone loss. The speech audiogram often shows greater impairment of speech recognition than would be expected based on the pure-tone audiogram.

**Differential diagnosis:** The differential diagnosis includes virtually all known factors that lead to gradual, bilateral hearing loss (see 12.2, pp. 260–265). The most important of these are hereditary sensorineural hearing loss, chronic noise-induced hearing loss, and hearing loss with an ototoxic or metabolic cause.

**Course:** Generally, the hearing loss is progressive, but its degree and time course are highly variable and cannot be predicted.

**Treatment:** No specific medical or surgical treatment is available. Bilateral hearing-aid fitting and other rehabilitative measures to improve communication may be necessary, depending on the auditory handicap (see 8.5, pp. 192–195).
Sudden Sensorineural Hearing Loss

**Definition:** Sudden sensorineural hearing loss refers to an immediate, unilateral hearing loss of sensorineural origin that has no apparent external cause. There are a number of known causes of sudden hearing loss. If a cause is identified and a diagnosis can be made, the condition is described as **symptomatic sudden hearing loss.** Often the cause remains unknown, however, and a pseudodiagnosis of **idiopathic sudden hearing loss** is made.

This section deals mainly with idiopathic sudden hearing loss, since the management of symptomatic hearing loss is dependent on the cause.

**Etiology and pathogenesis:** Idiopathic sudden hearing loss is believed to have a viral, vascular, or autoimmune cause. A disturbance in the homeostasis of the inner ear is probably present, and psychosomatic factors may occasionally play a role.

**Symptoms:** The symptoms appear within seconds to hours. The hearing loss is variable in its affected frequencies and degree, ranging from a mild loss to sud-
den deafness. Tinnitus is often present as an associated feature; vestibular symptoms are less common.

**Diagnosis:** The clinical diagnosis of “sudden sensorineural hearing loss” is made in patients who present with sudden, unilateral hearing loss of sensorineural origin. The history rules out an external causal event. Otoscopy is normal, and the Weber test is lateralized to the healthy ear. **Examination by a specialist** should not be delayed and should include inspection of the external ear canal and tympanic membrane with an otomicroscope, tests of vestibular functions, and pure-tone audiometry. The work-up may also include otoacoustic emissions and impedance audiometry. All of these tests are intended to exclude known causes of a symptomatic sudden hearing loss. This process can be aided by additional tests for arterial hypertension, hyperlipidemia, borreliosis, syphilis, toxoplasmosis, and neurotropic viruses such as mumps, herpes zoster, and HIV. A neurologic or immunologic work-up should be completed in patients who also have neurologic or inflammatory symptoms. ABR testing or MRI should be scheduled at a later date to exclude a retrocochlear lesion, especially a cerebellopontine angle tumor. These tests subject the inner ear to an acoustic stress, and therefore they should be done following a recovery period of at least one week.

**Differential diagnosis:** Because idiopathic sudden hearing loss is a diagnosis of exclusion, the differential diagnosis basically includes all causes of symptomatic sudden hearing loss, which are listed in Table 12.3. It is particularly important to detect a perilymphatic fistula after microtrauma, infections such as borreliosis and herpes zoster, and vestibular schwannoma (see 15.3, p. 307), as these conditions require specific therapies.

The possibility of a psychogenic hearing disorder should always be considered as well.

**Treatment:** Causal treatment for **symptomatic sudden hearing loss** should be initiated as soon as possible. This particularly applies to cases caused by labyrinthitis (see 12.2, p. 262), herpes zoster oticus, or perilymphatic fistula (see p. 286). **Idiopathic sudden hearing loss** is treated with measures aimed at improving the microcirculation and oxygenation of the inner ear. Steroids are also prescribed. No therapy has proven to be effective, however.

**Prognosis:** More than 50% of patients with idiopathic sudden hearing loss show a spontaneous improvement or resolution of complaints, regardless of treatment. This usually occurs during the first few weeks.
Chronic Progressive, Idiopathic Sensorineural Hearing Loss

Synonym: idiopathic bilateral sensorineural hearing loss

Definition: Chronic progressive idiopathic hearing loss is a bilateral sensorineural hearing loss that usually begins between 30 and 50 years of age.

The disease is defined as having its onset before age 50 to distinguish it from presbycusis.

Etiology: The cause of the hearing loss is unknown—i.e., there is no clinical evidence of toxic, metabolic, genetic, infectious, or other intrinsic or extrinsic causes. Nevertheless, causes such as a recessive genetic defect cannot be definitely excluded.

Symptoms: Both sides are affected, but the course is variable. The disease may begin with sudden hearing loss or may progress gradually. The progression of the hearing loss is variable but often culminates in severe bilateral hearing loss or deafness over a period of years or decades. Hearing impairment is the dominant finding and is frequently accompanied by tinnitus. Vestibular symptoms are generally absent, however.

Diagnosis: The audiometric detection of typical cochlear hearing loss, together with the progression of symptoms and the exclusion of other etiologic factors, confirm the diagnosis.

Differential diagnosis: Chronic progressive idiopathic hearing loss is a diagnosis of exclusion, and therefore the differential diagnosis includes all known causes of chronic sensorineural hearing loss—metabolic, genetic, autoimmune, infectious, physical, and toxic. A retrocochlear, psychogenic, or central hearing loss should also be excluded.

Treatment and prognosis: Specific treatment cannot be offered in cases with an unknown cause. Rehabilitative measures in patients with an auditory handicap should be instituted early, given the progressive nature of the disease. Since bilateral disease may culminate in severe hearing loss or even complete deafness, these patients should be promptly referred for comprehensive rehabilitation that includes training in lip reading. A cochlear implant is often beneficial in cases in which hearing aids no longer give acceptable improvement.
Vestibular Disorders

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13.1 Clinically Relevant Anatomy and Function of the Vestibular System

The clinical anatomy of the petrous bone and inner ear–labyrinthine complex was covered in 7.1 (pp. 154–157). This unit provides additional specific and clinically important details on the anatomy and physiology of the vestibular system.

Vestibular End Organ

The vestibular labyrinth is part of the inner ear (see also p. 156). It consists of the semicircular canals, which are sensitive to angular acceleration, and the otolithic apparatus, which is sensitive to linear acceleration (Fig. 13.1).

Semicircular Canals

The three semicircular canals occupy three spatial planes that lie at right angles to one other (Fig. 13.1). The posterior semicircular canal is directed along the axis of the petrous bone (approximately 45° to the sagittal and coronal plane; Fig. 13.1 a) and is roughly vertical. The lateral (horizontal) semicircular canal is tilted approximately 30° upward from the horizontal plane at its anterior end when the head is in a normal upright position. When caloric testing is done in the supine position, the head must be elevated by about 30° so that the lateral canal lies vertically.

A rotational stimulus is amplified in the vestibular nuclei by a push–pull mechanism resulting from the mirror-image arrangement of the right and left semicircular canal systems. The inhibition of neural discharges in the semicircular canals on one side (“push”) is accompanied by an increased discharge rate on the opposite side (“pull”).

Each semicircular canal has a dilation at its utricular end called the ampulla (Fig. 13.2). The ampulla contains the sensory cell system of the associated semicircular canal, consisting of the crista and cupula. Due to the inertial lag of the endolymph, angular acceleration of the head causes a deflection of the cupula that displaces the sensory cilia within it. The cupular motion is a bowing rather than a swinging-door type of movement, and it is this deflection that stimulates the vestibular sensory cells.

Otolithic Apparatus

The vestibular apparatus contains two additional sensory cell regions called the static maculae (Fig. 13.3). The hair-cell cilia in these regions are embedded in a gelatinous material called the otolithic membrane. This membrane is studded with ototholiths (synonyms: otoconia, statoliths), which are calcium carbonate
crystals ranging from 0.1 to 30 μm in size. With the head in an upright position, the macula of the utricle (see Fig. 7.3, p.156) is roughly horizontal while the macula of the saccule is approximately perpendicular to it (see Fig. 13.1b).

The otolithic organs are sensitive to linear accelerations that are not perpendicular to the macula. These forces cause the relatively inert otolithic membrane to shift in relation to the layer of sensory cells. Because the gravitational force produces a static linear acceleration and the two maculae are at approximately right angles to each other, at least one macula is stimulated at all times. In this way, the otolithic apparatus senses the position of the head in space. If normal gravitation is absent (as during space flight), this function is lost and severe vestibular disturbances may arise. Translations and other dynamic linear accelerations are registered equally on both sides, resulting in a summation effect at the central level as opposed to the push–pull effect occurring in the semicircular canals.

**Central Vestibular Structures**

The vestibular nerve is the part of the eighth cranial nerve that connects the sensory cells of the end organ to the vestibular nuclei in the brainstem. Its ganglion cells are located in the distal third of the internal auditory canal (vestibular ganglia, synonym: Scarpa ganglia). The vestibular nuclei process and integrate both vestibular and nonvestibular information, thereby influencing the function of the oculomotor and spinal motor systems. The left and right nuclei are interconnected by functionally important commissures that play a key role in the maintenance of equilibrium. The central vestibular motor system is shown schematically in Fig. 13.4. The principal nonvestibular afferent projections of the vestibular nuclei are as follows:

- **Visual afferents**: Information on the movement of visual images across the retina (“retinal slip”) is conveyed directly to the vestibular nuclei. This ex-
Vestibular Functions

Normal vestibular function cannot be perceived or described in isolation. This is because the vestibular organ is at least tonically active at all times and is constantly interacting with other sensory systems, especially the visual and proprioceptive systems. The major functions of the vestibular system include the following:

- Fixation of the visual horizon for spatial orientation during rapid head movements. This function relies on interaction with the visual system.
- Maintenance of posture and equilibrium. This function relies mainly on interaction with the proprioceptive and motor systems.

Vestibulo-Ocular Reflex

Some of the most important efferents of the vestibular nuclei are projected to the motor nuclei of the eye muscles. Reflex corrections in the movement and position of the eyes by the vestibular organ help to maintain spatial visual orientation through a compensatory adjustment of the visual horizon, allowing a fixed object to stay within the visual field. With the rapid, small-amplitude head movements that occur during many activities, corrections by the visual system alone would be too slow for stable fixation. To make these corrections more rapidly, the vestibular signals are relayed to the eye muscles by direct efferent projections from the vestibular nuclei to the oculomotor nuclei. The vestibulo-ocular reflex (Fig. 13.5) is a reflex that involves three neurons and has no direct feedback. The vestibular nuclei can receive feedback from retinal receptors for fine control, but this feedback is absent in darkness or when the eyes are closed. In this case the vestibulo-ocular reflex is an “open-loop reflex” and the corrective response is generally smaller than the actual movement. The stimuli for the vestibulo-ocular reflex come mainly from the semicircular canals. The otolithic organs, however, appear to have a role that has not been fully appreciated in the past. The reflex is also influenced by polysynaptic connections between the two vestibular nuclei (e.g., convergent neurons) and by connections with the cerebellum.

Vestibulospinal Reflexes (VSRs)

The vestibular nuclei can affect spinal motor activity (head position, postural stability, upright gait) directly via the lateral and medial vestibulospinal tracts and indirectly via the reticulospinal tract. The vestibulocervical reflex (VCR, called also the vestibulocollicular reflex) is particularly important for stabilizing the head position and thus for maintaining visual orientation. Spinal motor function is controlled by proprioceptive, visual, and vestibular reflexes. These three sensory systems are mutually complementary and are subject to mutual controls. An upright gait is generally possible when two of the systems are functioning. If two of the systems are damaged, upright posture and movement are impaired. In the clinical testing of spinal motor function, the vestibular influences cannot be tested in isolation. Other sensory systems must always be tested as well.
13.2 Examination of the Vestibular System

Vestibular function tests are used in the investigation of vertigo and dysequilibrium. A detailed history is of key importance and is supplemented by simple clinical tests. Based on this information, the physician can then select further instrumented diagnostic procedures. The history and differential diagnosis in patients with vertigo are covered in the next (pp. 280-281).

History

A detailed history will often allow for a preliminary diagnostic classification of dizziness or dysequilibrium. The following information should be elicited:

**Nature:** systematic complaints such as vertigo or loss of balance with directional components (see p. 281 for details)?

**Onset:** acute or gradual onset?

**Duration:** spells lasting for seconds, minutes or hours, or more prolonged occurrences lasting for days or more?

**Course:** diminishing, constant, or increasing?

**Provocative factors:** are complaints brought on by certain positions or movements (position-dependent or posture-dependent)?

**Accompanying symptoms:**

- **Otolologic** complaints such as hearing loss, tinnitus, or otorrhea?
- **Neurologic** complaints such as headache, loss of consciousness, or dysarthria?
- **Visual** complaints such as visual impairment or diplopia?
- **Autonomic** complaints such as nausea or vomiting?

The history should also cover the following:

- Use of drugs and medications
- Trauma to the skull or cervical spine
- Diseases of the cardiovascular system
- Neurologic diseases

Clinical Examination

The clinical examination focuses on the patient's ear, nose and throat status, with particular emphasis on otologic findings and cranial nerve functions. The rest of the examination consists of:

- Testing spinal motor function and coordination
- Oculomotor function testing
- Nystagmus testing

Spinal Motor Function and Coordination

The evaluation of spinal motor function should test the spontaneous coordination of the body as a whole and of the upper extremities. It should precede provocative vestibular testing, therefore. The **standard vestibular test battery** should include the following tests:

**Romberg test:** The patient stands upright with the eyes closed and the feet parallel and close together (to reduce the stance area). A subject with a normal vestibular system can stand in this position for 30 seconds without significant body sway.

**Unterberger stepping test:** With eyes closed, the patient takes 50 steps in place, bringing the thighs up to a horizontal position with each step. A tonus discrepancy due to a vestibular lesion will cause the patient to rotate toward the side of the affected labyrinth (up to 45° of rotation is normal).

**Finger-to-nose test:** With eyes closed and the arm extended forward, the patient slowly brings the forefinger to the tip of the nose. Ataxia, intention tremor, and action myoclonus indicate a cerebellar lesion.

If these results are equivocal, **additional diagnostic tests** can be performed that basically test the same global reactions and their coordination:

**Walking straight with eyes closed:** The patient walks forward for 4 meters along a straight line with the eyes closed, endeavoring to place one foot directly in front of the other. A normal subject will not stray from the line. A patient with a vestibular disorder will deviate constantly toward the affected side.

**Walking forward and backward:** The patient alternately takes two or three steps forward and backward. As in the Unterberger test, a patient with a vestibular disorder will rotate toward the side of the lesion, in this case tracing out a star-shaped path.

**Arm tonus and drift test:** The patient sits with the eyes closed and both arms extended forward and supinated. Sagging of one arm indicates a cerebellar lesion (arm tonus reaction), while a parallel drift of both arms toward one side indicates a vestibular lesion (drift response). The tonus discrepancy can be accentuated by having the patient move the extended arms up and down between the knees and the horizontal (modified Barany pointing test).
13.1 Nystagmography and related tests

**Definition:** Nystagmography is a procedure for recording eye movements and evaluating spontaneous or induced nystagmus in darkness (without visual fixation) and during rotation. It is used to document and calculate quantitative parameters. **Technique:** Electronystagmography (ENG) utilizes the dipole properties of the eye to record ocular movements with electrodes affixed to the skin (Fig.). Another nystagmographic technique is **video-oculography,** in which the eyes are observed with an infrared camera and pupillary movements are automatically tracked and recorded.

**Interpretation:** The recorded nystagmus can be automatically analyzed with a computer, the slow phase velocity being the most important parameter for further quantitative study. The standard ENG test battery includes the following tests:
- Recording spontaneous nystagmus: Spontaneous eye movements are recorded with the patient looking straight ahead and with gaze directed 30° to the left and to the right, with visual fixation and in darkness (see p. 277).
- Tracking test for the analysis of slow eye movements (see below)
  - Optokinetic testing (see p. 279)
  - Rotational testing (see p. 279)
  - Caloric stimulation (see p. 279)
Other, specialized tests are also available for further investigation of the vestibular system:
- Tests of the otolithic organs (e.g., determining the subjective vertical or eccentric rotation)
- Posturography: objective recording and analysis of spinal motor function and coordination on stationary or moving (dynamic) platforms
- Saccadic analysis: induction and oculographic analysis of saccades (fast eye movements)
- Neck rotation test: tests for nystagmus induced by rotating the body while the head remains stationary
- Vestibular evoked myogenic potentials (VEMP): stimulation of the sacculus by acoustic clicks and registration of evoked muscle potentials of the sternocleidomastoideus

**Diadochokinesis:** Both hands are rapidly pronated and supinated to test cerebellar function and fine motor skills.

These tests always evoke a cumulative response that is only partly influenced by the vestibular system (see p. 273). The results should be interpreted accordingly, taking into account the overall situation.

As a general rule of thumb, a vestibular disorder leads to directional instability, usually toward the side of the affected vestibular organ, while other neurologic disorders lead to nondirectional instability or ataxia.

**Eye Movements**

Abnormalities of eye movements, especially oculomotor palsy, should be distinguished from nystagmus (see below). Simple clinical tests are helpful in making this differentiation:

**Ocular motility:** The patient tracks the tester’s forefinger or some other target moving in an H-shaped pattern (Fig. 13.6). Observing the eye movements provides a cumulative test for all the eye muscles. Normal motility is reflected in a complete, bilateral freedom of eye movements, which are coordinated (equal on both sides).

**Smooth pursuit** (Fig. 13.7): A target is slowly moved back and forth at a comfortable fixation distance from the patient’s eyes. The tester should not move the target more than 30° to either side. A normal response consists of a smooth, coordinated tracking movement.
Oscillopsias can be detected with the Halmagyi test. The patient and examiner sit opposite each other, about 1 meter apart. The examiner moves the patient’s head rapidly back and forth over about a 30° arc while the patient keeps the gaze fixed on the examiner’s nose. If the vestibular control of eye movements is weak, definite saccade-like corrective movements will be noted due to difficulties in maintaining fixation.

**Nystagmus: Definition, Classification, and Testing**

**Definitions:** Nystagmus in the broad sense refers to any rhythmic eye movement, which may be spontaneous, provoked, or induced. The eye movement may be physiologic or may be symptomatic of an abnormal condition.

Nystagmus in the strict sense (Fig. 13.9) refers to a conjugated, coordinated eye movement about a certain axis, which can be subdivided into rhythmically alternating slow and fast phases. The slow phase is the tonic eye movement induced physiologically or pathologically by a vestibular stimulus. The opposing fast phase is a saccade-like refixation movement induced by the central oculomotor system. Visual perception is suppressed during the refixation movement, producing the constant rotatory sensation that is characteristic of nystagmus.

The direction of the nystagmus is defined by the easily observed fast component. Nystagmus that is present at rest within a 30° visual field is called spontaneous nystagmus (see “Types of spontaneous nystagmus,” below). It has pathologic significance.

Provoked nystagmus is a pathologic nystagmus that can be produced by certain maneuvers. It generally has a vestibular cause and may be transient (lasting less than 60 seconds) or persistent (lasting more than 60 seconds) in response to a provocative maneuver. This differs from induced nystagmus, which is a physiologic phenomenon (e.g., “train nystagmus”)

**Diagnosis:** Frenzel glasses are an extremely helpful aid in the diagnosis of nystagmus. The patient’s eyes are illuminated, and glasses with a power of 20 diopters eliminate visual fixation. Generally, a minimum of three beats are needed to detect nystagmus.

**Spontaneous Nystagmus**

**Screening for spontaneous nystagmus:** First the patient fixes on a well-defined target about 1 meter away that is presented straight ahead and also at a 30° angle to the left and right. The patient may additionally be tested at 15° upward and downward gaze. The eyes are observed while the targets are presented.
Next, Frenzel glasses are put on to suppress visual fixation, and the same gaze directions are tested as before. Any regular, rhythmical eye movement is classified as spontaneous nystagmus, but not the nondoncational, irregular, jerky movements (“square waves”) that often occur when fixation is suppressed.

**Types of spontaneous nystagmus**: Three main types can be distinguished:

- **Vestibular spontaneous nystagmus**: Any directional nystagmus that always beats toward the same side and can be abolished or markedly suppressed by visual fixation is classified as vestibular spontaneous nystagmus. It is caused by a peripheral or central vestibular lesion and generally beats toward the healthy side. Horizontal nystagmus can be graded according to its spontaneous presence (Alexander grading scheme):
  - **First-degree nystagmus** is present only during gaze in the direction of the nystagmus.
  - **Second-degree nystagmus** is also present on central gaze.
  - **Third-degree nystagmus** is present in all gaze positions.

- **Gaze-evoked nystagmus**: The fast phase of the nystagmus is always in the direction of gaze—i.e., to the right during gaze toward the right side, and to the left during gaze toward the left side. This condition is a central gaze disturbance that causes a slow deviation of gaze toward the center of the visual field. The gaze intention produces the recentering movement, and so the nystagmus generally becomes more intense as the deviation increases. Gaze-evoked nystagmus can also result from toxicity such as barbiturate, phenytoin or alcohol poisoning.

Differentiation is required from the physiologic phenomenon of end-point nystagmus, which occurs through muscular action when the gaze deviates by approximately 40°.

**Fixation nystagmus** (pendular nystagmus, congenital nystagmus) is defined as nystagmus that is most pronounced in the central position and during fixation. Usually it displays a high frequency with no clearly distinguishable fast and slow components (hence the name “pendular nystagmus”) and is present during binocular fixation. A rare variant occurs only during monocular fixation and beats toward the side of the fixation (latent fixation nystagmus). Fixation nystagmus is often based on a congenital abnormality of the oculomotor centers and causes no subjective complaints. Acquired lesions are a less frequent cause. Fixation nystagmus may also occur in association with visual disorders (amblyopic nystagmus).

**Provocation of Nystagmus**

As before, the nystagmus is observed through Frenzel glasses and can be positively identified after a minimum of three beats. **Common provocative maneuvers are listed below:**

- **Head shaking**: Gentle, passive, horizontal shaking of the patient’s head may “unleash” a spontaneous nystagmus.
- **Positional testing (static)**: This involves the slow-motion assumption of various body positions (supine, lateral decubitus, head hanging). The vestibular apparatus and especially the otolithic organs are exposed to various gravitational stimuli in the different positions.
- **Hallpike–Dix maneuver**: The patient is moved swiftly from an upright sitting position to a head-hang-
If the tympanic membrane is perforated, warm or cold air can be used for caloric stimulation. A difference of 7° between the irrigating water and body temperature will induce endolymphatic movement through conduction. This causes a unilateral stimulation of the vestibular labyrinth and induces a typical vestibular nystagmus, which generally lasts approximately 2–3 minutes.

Warm-water irrigation induces nystagmus toward the same side: hot = homolateral or warm = same. Cold irrigation induces nystagmus toward the opposite side: cold = contralateral or cold = opposite.

In a normal test, both labyrinths are stimulated equally. If the test is performed using Frenzel glasses, the examiner waits 30 seconds after the irrigation and then counts the number of beats for an additional 30 seconds.

• **Rotational stimulation:** The rotational stimulus represents the physiologic stimulus for the vestibulo-ocular reflex (see p. 274). Rotation about an axis passing through the head stimulates one or more semicircular canals on each side, depending on the head position. The left and right sides are stimulated in an opposing fashion: Rotating the head in one direction (e.g., to the right) induces a nystagmus toward the same side (to the right; rotatory nystagmus). When the rotation ceases, the nystagmus reverses to the opposite side (in this example, to the left; postrotatory nystagmus).

The test is conducted in darkness using nystagmography (13.1, p. 276). Rotational testing is particularly useful for quantitative analysis of the vestibulo-ocular reflex and is therefore used in follow-ups.

In contrast to this vestibular nystagmus, rotation of the visual field while the head remains stationary induces an **optokinetic nystagmus.** The patient observes a striped pattern that preferably fills the entire visual field and is alternately rotated to the left and right, in each case inducing a nystagmus in the opposite direction.

### Induction of Nystagmus

**Vestibular nystagmus** can be induced in two ways.

• **Caloric stimulation:** The patient lies with the head elevated 30° (so that the lateral semicircular canal is vertical, see p. 272), and the external auditory canals are irrigated with warm (44°C) or cold water (30°) for 30 seconds.

Make sure that the tympanic membrane is intact before performing this test.
13.3 Diagnosis and Differential Diagnosis of Vertigo and Dysequilibrium

Vertigo and balance disorders (dysequilibrium) are common ambiguous symptoms that can be clarified by taking a systematic history and applying targeted diagnostic procedures. The present explores options for this type of approach. Dysequilibrium generally takes precedence over vertiginous complaints. Vertigo may well be present in the absence of dysequilibrium, or a balance disorder may occur without vertigo, as in patients with predominantly proprioceptive or motor disturbances (Fig. 13.11).

Vertigo: Definition and Pathogenesis

Vertigo (dizziness) is a frequent symptom that is often very difficult to relate to a specific diagnosis. This may be due partly to ambiguities in the term “dizziness,” which patients use to describe a multitude of complaints that can range from a vague queasy feeling to “blacking out” due to circulatory problems.

**Definition:** Vertigo is a subjective disturbance of integrity caused by contradictory sensory information processing.

**Pathogenesis:** Based on this definition, there are two different ways in which vertigo can occur:

- **The function of a sensory system is impaired:** The usual interaction of vestibular, visual, and proprioceptive sensory systems is disrupted because of a functional disturbance in a peripheral system (see also p. 273). For example, the unilateral loss of a peripheral vestibular system creates an imbalance between the two vestibular systems, leading to rotary vertigo. With a constant or insidious disorder, the central nervous system is usually able to compensate for the deficit over time, with the result that vertigo disappears or does not occur at all.

- **Central processing is impaired:** The information supplied by the normally functioning sensory systems is incorrectly processed or interpreted. This results in conflicting sensory impressions and a feeling of dizziness. The processing disorder may be caused by diffuse changes such as metabolic or circulatory abnormalities, infection, trauma, or intoxication. Local central nervous system (CNS) changes such as vascular occlusions, tumors, and inflammations may also be causative.

A combination of peripheral and central processing disorders may occur. These multisensory vertigo syndromes are particularly common in elderly patients (Fig. 13.11). Contradictory sensory processing is a concept that can also explain the vestibular and spatial sensory disturbances that occur in normal individuals, such as height vertigo and motion sickness. Height vertigo is caused by a conflict between the unaccustomed visual horizon, which signals “depth” and “danger of falling,” and the vestibular and proprioceptive systems, which perceive normal circumstances and no instability. In motion sickness (see also 13.2, p. 284), cumulative conflicts arise between habituated patterns of interaction and the patterns that are experienced in the mode of travel. The moving vehicle repeatedly generates unaccustomed combinations of vestibular, visual, and proprioceptive stimuli that ultimately lead to motion sickness.
**Differential Diagnosis of Vertigo**

The history is important in the differential diagnosis of vertigo. First, an effort should be made to distinguish between **systematic vertigo** and **nonsystematic dizziness**. Vertigo is classified as systematic if it has a definite directional or rotational component or if the patient can clearly describe the dizzy sensation. Next, questions dealing with otologic and neurologic symptoms can narrow the differential diagnosis to a most likely group of disorders. The algorithms in Fig. 13.12 and Fig. 13.13 outline a simple approach.

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**Fig.** History taking in vertigo patients

**Vertigo / dizziness**
- **Type of vertigo?**

  - **Rotary vertigo**
  - **Reeling vertigo**
  - **Elevation vertigo**
  - **Lateropulsion**

  **Obtundation**
  - **Light-headedness**
  - **Giddiness, spaciness**
  - **Unsteadiness**
  - **Impaired consciousness**

**Systematic vertigo** (with motion components)

**Nonsystematic dizziness** (no motion components)

**Otologic symptoms** such as hearing loss, tinnitus, otalgia, or otorrhea?

**Neurologic symptoms** such as dysarthria, diplopia, focal signs, or impaired consciousness?

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An initial history is taken to advance the differential diagnosis of vertiginous complaints. The algorithm provides clues that are helpful in directing further tests. Especially when a vestibular cause is suspected, the time course of the vertigo and associated symptoms can further narrow the differential diagnosis.

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**Fig.** Differential diagnosis of vestibular vertigo based on duration

**Duration of vertigo?**
- **Seconds**
  - During head movements
  - At rest

- **Minutes**
  - Position-dependent

- **Hours or Days**
  - Intensity diminishing
  - Intensity increasing

**During head movements**
- **Peripheral vestibular disorder** (bilateral vestibular loss)
- **Central vestibular disorder and psychogenic vertigo**

**At rest**
- **Benign paroxysmal positional vertigo** or arterial hypertension

**Position-dependent**
- **Central vestibular disorder** (e.g., drug-induced)

**Position-independent**
- **Peripheral vestibular disorder** (vestibular neuritis)
- **Central vestibular disorder or multisensory vertigo**

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13.4 Vestibular Disorders

Peripheral dysfunction and diseases of the vestibular system are more common than central vestibular disorders. The most common clinical entities based on an isolated peripheral vestibular disorder are unilateral vestibular loss and benign paroxysmal positional vertigo. Less common entities are Ménière’s disease and bilateral vestibular loss. Central vestibular disorders can usually be recognized by their presentation, and most cases are investigated further by a neurologist.

Peripheral Vestibular Disorders

Acute Unilateral Vestibular Loss

Synonyms: vestibular neuritis, vestibular neuritis

Definition: Analogous to sudden sensorineural hearing loss, this disease is characterized by a sudden impairment of peripheral vestibular function on one side.

Epidemiology: Middle-aged individuals are affected predominantly but not exclusively. The disease often appears about two weeks after a viral upper respiratory infection.

Etiopathogenesis: The exact site of the lesion (end organ or vestibular nerve) and the etiology are unknown. Viral (“neuritis”), vascular, metabolic, and mechanical causes have been proposed.

Symptoms: The onset is marked by a sudden, severe attack of rotary vertigo with no apparent cause. Generally, a continuous vertigo develops and persists for several days. Nausea, vomiting, and disequilibrium cause severe subjective distress, and many patients are rushed to the emergency room with suspicion of an acute neurologic disease. There are no other otologic symptoms such as pain, hearing loss, or tinnitus, and there is no impairment of consciousness or other neurologic symptoms.

Diagnosis: A horizontal or rotary spontaneous nystagmus can always be detected in the acute stage and is increased by visual fixation suppression with Frenzel glasses. Caloric stimulation demonstrates non- or hypoexcitability of the affected labyrinth, which is on the side opposite to the direction of the nystagmus. The audiogram is normal.

Differential diagnosis: Differentiation is mainly required from central vestibular disorders such as vascular insufficiency and mass lesions, particularly a cerebellar infarction.

Treatment: Treatment in the acute stage consists of bed rest in a darkened room, corticosteroids, fluid intake (infusion), and antivertiginous drugs (thiethylperazine type) administered i.v. or by suppositories. After the acute phase and nausea have passed, the patient should be mobilized from bed as soon as possible, as this will promote central compensation.

Course: The symptoms subside in a few days with spontaneous recovery of equilibrium (vestibular compensation). Younger patients tend to recover more quickly than older patients. Caloric hypoexcitability of the affected labyrinth often persists, but recovery of vestibular function may occur.

Prognosis: The prognosis is favorable, and most patients recover completely. Latent dysequilibrium with a falling risk may persist in older patients, however, and balance training is recommended in these cases.

Benign Paroxysmal Positional Vertigo (BPPV)

Synonyms: canalolithiasis, cupulolithiasis

Epidemiology: Benign paroxysmal positional vertigo is a frequent cause of sudden attacks of vertigo. Women predominate over men by about a 2:1 ratio.

Etiology: Possible causative factors include head trauma, a prior history of vestibular neuritis, and inner ear disorders (e.g., previous stapes surgery or labyrinthitis).

Pathogenesis: BPPV is a peripheral disorder based on a lesion of the end organ. The vertigo is believed to be incited by particles floating in the endolymph of a semicircular canal (canalolithiasis). Usually, the particles are otoconia that have become separated from the macula. During certain movements, the increased mass of the otoconia causes an unphysiologic deflection of the cupula, which produces the typical symptoms. At rest, the particles are deposited mainly in the posterior semicircular canal.
**Symptoms:** The patient complaints of severe, recurrent attacks of rotary vertigo lasting approximately 1 minute. The attacks are provoked by certain movements and are fatigable. Nausea may occur, but there are no other associated symptoms. Often the symptoms are more intense following rest, and the patient may be awakened at night by vertiginous attacks.

**Diagnosis:** The diagnosis is based on the typical history or is established by the Hallpike maneuver (position change with head hanging, Fig. 13.10, p. 279). Placing the head over the end of the table typically evokes a rotary nystagmus to one side after a latent period of about 10 seconds. The nystagmus increases for about 30 seconds, then diminishes (crescendo–decrescendo nystagmus). When the patient is brought back to an upright position, a similar nystagmus occurs in the opposite direction. This positional nystagmus is fatigable and disappears after several repetitions of the maneuver.

**Differential diagnosis:** Positional nystagmus can also occur in the setting of central or neural vestibular disorders. These cases, however, generally do not manifest the typical clinical complaints and diagnostic findings described above.

**Treatment:** Initial treatment consists of therapeutic exercises—brisk positioning maneuvers or a defined sequence of head positions designed to displace and reposition the canaliths (canalith repositioning). In rare cases, it may be necessary to surgically disrupt the neural connection to the ampulla of the posterior semicircular canal (singular neurotomy) or to block the posterior semicircular canal.

**Course:** As the name indicates, the vertigo is benign and generally resolves spontaneously in a matter of days or weeks. Occasionally the symptoms recur after a variable period of time. It is rare for the complaints to persist for a prolonged period.

**Ménière’s Disease**

**Definition:** Ménière’s disease is generally unilateral. The classic clinical triad consists of
- **Attacks of vertigo** lasting several hours
- **Tinnitus**
- **Fluctuating hearing loss**

Terms such as “Ménière’s complaints” or “atypical Ménière’s disease” should be avoided, as they may refer to any type of rotary vertigo. Ménière’s disease is diagnosed only when the classic triad is present.

**Epidemiology:** Ménière’s disease is rare. It can be diagnosed in only about 5–10% of patients with rotary vertigo and is certainly overdiagnosed. The typical age of onset is between the third and fifth decades, but Ménière’s disease also occurs in younger and older individuals. Bilateral involvement, which is usually monosymptomatic, occurs in approximately 5–10% of cases.

**Etiology:** The etiology of Ménière’s disease is poorly understood. Most cases are idiopathic, and some patients have a positive family history, suggesting that there may be predisposing anatomical factors such as a narrow vestibular aqueduct. In rare cases, Ménière’s symptoms can also result from previous inner ear damage, usually due to trauma (post-traumatic cochlear hydrops).

**Pathogenesis:** Endolymphatic cochlear hydrops has been identified as a morphologic correlate of Ménière’s disease. It involves a relative overproduction of endolymph with distention of the cochlear duct and displacement of the Reissner membrane toward the scala vestibuli (see Fig. 7.6, p. 160). In most cases, the hydrops probably results from an inadequate reabsorption of endolymph in the endolymphatic sac rather than a true overproduction. The hydrops causes displacement of the basilar membrane, creating an unfavorable mechanical environment for the hair cells that can lead to sensorineural hearing loss and probably to tinnitus as well. It has been theorized that attacks of vertigo may be caused by a rupture of the Reissner membrane, which would allow the mixing of perilymph and endolymph leading to potassium intoxication in the perilymphatic space. As yet, however, there is no proof that this type of rupture occurs or is linked to vertiginous attacks.

**Symptoms:** Ménière’s disease is characterized by a typical history with the following symptoms:
- Fluctuating hearing loss and dysacusis
- Tinnitus, which is continuous but of varying intensity and low frequency (roaring)
- Sensation of aural fullness
- Attacks of rotary vertigo, nausea and vomiting, which last for hours. Dys-equilibrium and dizziness may be present shortly before and long after the attacks.

The tinnitus and hearing loss may change before or during the attack. Most commonly, the tinnitus becomes louder while hearing becomes poorer. Hearing generally improves again following the attack. A marked improvement of hearing during the attack itself is called the Lermoyez phenomenon. Drop attack (Tumarkin crisis), in which the patient falls suddenly to the ground without losing consciousness, is somewhat rare and occurs late in the disease process.
13.2 Other vestibular disorders

Motion Sickness
Motion sickness is related to the function of the labyrinth and, as such, is a "physiologic" response. Motion sickness does not occur in patients with nonfunctioning peripheral vestibular organs. It probably results from a shift in accustomed patterns of sensory processing (sensory conflict or mismatch). The major conflicts are between the visual and vestibular systems, bringing about the typical symptoms of malaise, fatigue, yawning, increased salivation, nausea, and finally vomiting. Prevention relies on anticipating and visually perceiving the motion with as much of the visual field as possible. Motion with a limited visual field (porthole, rear seat) predisposes to motion sickness. Training can induce the development of central patterns that reduce susceptibility to motion sickness. The condition can be treated medically with antivertiginous drugs, which should be taken before travel whenever possible (scopolamine transdermal patch, dimenhydrinate, cinnarizine). These drugs may induce fatigue and therefore should not be taken before operating a motor vehicle.

Vestibular Disorders in Children
The most common type of vestibular dysfunction in small children is benign paroxysmal vertigo of childhood. This condition is not related to benign paroxysmal positional vertigo in adults (see p. 282). It is characterized by brief attacks of vertigo lasting seconds or minutes that occur when the child is feeling well and are associated with dysequilibrium and nystagmus. The child recovers completely within a short time. The attacks are most common between 1 and 4 years of age. They occasionally progress to migraine attacks, suggesting a link between this type of vertigo and basilar artery migraines. Generally the prognosis is good and the attacks disappear before puberty.

Psychosomatic Vertigo
Depression, anxiety neuroses, hyperventilation, and panic attacks may be associated with a sensation of subjective dizziness. Patients complain of constant, vague dizziness or of brief, recurring dizzy spells. The history provides the best information, and no evidence of a vestibular disorder can be detected. Treatment is geared toward the underlying disorder. Antivertiginous drugs are of no benefit.

Vestibular Disorders with a Cervical Cause
Proprioceptors located in the deep muscles of the neck and cervical vertebral joints are linked to vestibular and oculomotor centers and can evoke the cervicovestibular reflex (CVR) and cervico-ocular reflex (COR). The relationship between disturbances of these reflexes and vertiginous complaints is uncertain, however, as there is no reliable test for evaluating these reflexes. This makes it difficult to interpret the frequent concomitant complaints involving the nuchal region and the vestibular or cochlear system (tinnitus). These combinations of complaints may occur spontaneously in association with nuchal degenerative changes or may follow a whiplash injury. With whiplash trauma, interpretation is also hampered by a possible brain concussion or direct labyrinthine injury (otoIthcs).

Diagnosis: The diagnosis is based on the typical history described above. Otoscopic findings are normal in idiopathic Ménière’s disease, but late signs of previous inflammation or trauma may be found in symptomat-ic forms.

Pure-tone audiometry demonstrates a sensorineural hearing loss that mainly affects low frequencies in the initial stage of the disease (Fig. 13.14) and later affects all frequencies (pantonal hearing loss). Speech audiometry typically shows a disproportionate hearing loss for speech compared with the pure-tone audiogram. This is attributed to mechanical distortions within the cochlea.

Auditory brainstem response (ABR) testing is normal, and electrocochleography usually shows an elevation of the summation potential. Glycerin therapy (1.5 g glycerin/kg body weight) can improve the hearing threshold or reduce the summation potential in approximately two-thirds of patients (glycerol test). The vestibular findings in Ménière’s disease are variable initially and may reflect hypofunction or hyperfunction of the labyrinth. With progression of the disease, however, peripheral vestibular hypofunction will invariably develop on the affected side. Vestibular nystagmus is detectable during the attacks and generally beats toward the unaffected side. It may also beat toward the affected side, however, and it may even change direction during an attack.

Differential diagnosis: Other vestibulocochlear disorders (e.g., vestibular schwannoma, sudden sensorineural hearing loss) should be excluded as well as other causes of cochlear hydrops such as syphilitic labyrinthitis. Differentiation from basilar artery migraine is also occasionally required.

Treatment:
Acute attack: Treatment consists of bed rest and antivertiginous or antiemetic medications administered orally, rectally, or intravenously (e.g., thiethylperazine, dimenhydrinate, metoclopramide HCl). Patients with profuse vomiting should receive infusions for fluid and electrolyte replacement.

Prevention of attacks: Medical therapy between attacks should be tried initially in patients who experience frequent attacks. Antivertiginous antihistamines (be- tahistine) or centrally acting calcium antagonists (cinnarizine) can be tried between attacks, although the efficacy of this therapy is uncertain. If symptoms do not improve with medical treatment, measures should be taken to destroy portions of the labyrinth. This carries a risk of hearing loss or even deafness, and surgery may rarely lead to facial nerve paralysis. The following options are available:
- Intratympanic gentamicin: The drug is instilled into the middle ear through a myringotomy tube or directly through the tympanic membrane while response is monitored audiometrically and by watching for nystagmus. The primarily vestibulotoxic gentamicin diffuses through the round and oval window niche, causing partial ablation of the labyrinth.

- Vestibular neurotomy: The vestibular nerve is surgically divided through a transtemporal approach (through the middle cranial fossa), translabyrinthine approach (always associated with deafness), sublabyrinthine approach (through the mastoid), or retrosigmoid approach (through the posterior cranial fossa).

- Labyrinthectomy: Since surgical ablation of the labyrinth sacrifices hearing, this procedure should be considered only in patients who are already deaf.

**Course:** A characteristic feature of Ménière’s disease is its unpredictable course. Besides rare abortive forms in which the symptoms cease to occur after several attacks, periods of frequent attacks alternate with periods with few or no vertiginous complaints. The most distressing aspect for patients is the sudden, unpredictable attacks of vertigo. When such attacks are frequent, the patient may be incapacitated by the vertigo itself and by the fear of new attacks. Often, however, periods of frequent attacks are followed by months or years of remission in which the dominant findings are hearing loss and tinnitus. In typical cases, Ménière’s disease runs a progressive course for years with gradually diminishing attacks of vertigo but increasing hearing loss that culminates in deafness (“burned-out Ménière’s disease”).

**Bilateral Vestibular Loss**

**Pathogenesis:** Bilateral vestibular loss is a balance disorder caused by severe, bilateral hypofunction or failure of the peripheral vestibular apparatus. It usually has a systemic cause but may also result from bilateral disease of the vestibular apparatus. Examples of **systemic causes** are ototoxic drugs (aminoglycosides, diuretics, cisplatin), industrial toxins (aromatic hydrocarbons, heavy metals), and endogenous lesions (renal failure). Examples of **local diseases** that may occur bilaterally are bacterial, viral or autoimmune labyrinthitis, congenital or acquired labyrinthine disorders (anomalies, otosclerosis, acute vestibular loss on both sides), and diseases of the vestibular nerves (polyneuropathy, type 2 neurofibromatosis).

**Symptoms:** Bilateral vestibular loss does not produce nystagmus because there is no asymmetry of vestibular response. There is, however, a destabilization of visual fixation with typical oscillopsia (see p. 277). The patient describes a balance disturbance or “drunken” feeling, with an inability to recognize faces or read street signs while walking. The complaints are exacerbated in darkness.

The disorder is commonly associated with a variable degree of bilateral sensorineural hearing loss.

**Diagnosis:** Oscillopsias (see p. 277) are an abnormal finding. Caloric and rotational stimulation of the labyrinths elicit either no response or a very weak response. Auditory testing is also indicated and will often demonstrate at least a high-frequency hearing loss.
**Differential diagnosis:** A central vestibular disorder can produce similar complaints.

**Treatment:** Physical therapy with specific balance training may be helpful, as it can enhance the compensation of vestibular function by nonvestibular systems. Generally, however, patients are left with some degree of residual dysequilibrium.

**Other Causes of Peripheral Vestibular Disorders**

A viral, bacterial or autoimmune labyrinthitis generally leads to vertigo in addition to hearing impairment. Tests disclose the typical signs of peripheral vestibular dysfunction with vestibular nystagmus toward the opposite side.

**Otogenic inflammatory vestibular disorders:** An acute or chronic inflammatory process in the middle ear can lead to vertigo and dysequilibrium. The associated labyrinthitis is described in 12.2 (p. 262). When the cause is *acute otitis media* with an intact tympanic membrane, paracentesis should be performed at once to decompress the middle ear and windows. In patients with *chronic otitis media*, a cholesteatoma with a labyrinthine fistula should be considered whenever vestibular symptoms appear.

**Perilymphatic fistula:** A connection between the perilymphatic space and middle ear, usually through the oval or round window, can lead to vestibular symptoms and hearing disorders. Besides the otogenic inflammatory etiology mentioned above, the fistula may be caused by congenital anomalies or by direct trauma or barotrauma to the middle ear (see 11.4, p. 251).

**Treatment** in acute cases consists of bed rest with the head elevated and the avoidance of effort, nose blowing, and bearing down. If the symptoms persist or increase, the middle ear should be surgically explored. The window niches are examined under local anesthesia, and any fistulas are repaired.

**Tullio phenomenon:** Vestibular symptoms are occasionally induced by acoustic stimulation, usually with sound at low frequencies and high levels. Called the Tullio phenomenon after the author who first described it, this response requires differentiation from a perilymphatic fistula. The cause of the Tullio phenomenon is usually a change in the stapes (hypermobile stapes footplate or adhesion with the saccule), but it can also be associated with a perilymphatic fistula.

**Post-traumatic vestibular disorders:** Vestibular symptoms may be provoked by blunt head trauma with a labyrinthine concussion or contusion, a temporal bone fracture (see 15.2, pp. 302–305), trauma to the cervical spine, or a surgical operation. Most cases involve a peripheral, unilateral, cochleovestibular disorder. Central vestibular disorders can also occur in patients with a brainstem contusion or cerebral contusion. It can be difficult to differentiate among peripheral, central, and combined vestibular disorders. Central compensation of the peripheral disorders may be delayed. Also, a peripheral paroxysmal positional vertigo may develop several weeks after the trauma (see p. 251). Psychogenic factors also play a role in prolonged cases of post-traumatic vertigo.

**Neural and Central Vestibular Disorders**

**Vestibular Disorders with a Neural Cause**

**Cerebellopontine angle tumors:** Cerebellopontine angle tumors are a relatively infrequent cause of vestibular disorders, as they are usually slow-growing and symptoms are easily compensated over time. The symptoms associated with these tumors are described in 15.3 (pp. 306–309).

A cerebellopontine angle tumor should be included in the differential diagnosis of a unilateral cochleovestibular disorder. Vestibular signs consist of various forms of vestibular spontaneous or provoked nystagmus and caloric hyporeactivity of the labyrinth. Large tumors may exert pressure on the brainstem leading to central vestibular signs such as impaired smooth pursuit tracking or altered optokinetic nystagmus.

**Vestibular neuritis:** Isolated neuritis of the vestibular nerve is recognized as one cause of acute unilateral vestibular loss (see p. 282). This form of the disease is described under that heading.

Vestibular nerve disorders may occur in the setting of cranial polyneuritis or polyneuropathy. It is particularly important to detect or exclude treatable causes such as herpes zoster, borreliosis, and toxoplasmosis.

**Central Vestibular Disorders**

A central vestibular disorder may present with the following signs:

- Ataxia with relatively little vertigo
- Saccades during smooth pursuit tracking
- Gaze-evoked nystagmus or irregular nystagmus that beats upward or downward (upbeat, downbeat nystagmus)
- Nystagmus not suppressible by visual fixation
- Disturbances of the optokinetic reflex

The most frequent cause of a central vestibular disorder is vascular insufficiency in the brainstem region. The lesions may consist of very small or larger ischemic areas or hemorrhages. Other possible causes are inflammation (e.g., multiple sclerosis), infection (e.g., viral encephalitis), tumors (e.g., gliomas), metabolic disorders (e.g., Wernicke–Korsakoff syndrome), and trauma (e.g., brainstem contusion). The investigation and treatment of central vestibular disorders fall within the neurologic specialty.
Vertebrobasilar insufficiency: Vertebrobasilar insufficiency is a rare cause of vertiginous complaints. A transient blood-flow disturbance in the vertebrobasilar territory can lead to attacks of rotary vertigo, which usually have a central cause. But since the labyrinth derives its blood supply from the same territory (labyrinthine artery), decreased blood flow to the peripheral vestibular organs cannot be excluded. Vertebrobasilar insufficiency is also accompanied by other neurologic symptoms: visual disturbances, diplopia, drop attacks, loss of consciousness, speech disorders, or paralysis. Direct compression of the vertebral artery by a cervical spinal lesion or extreme head movements is rare, but blood flow may be reduced in a setting of transient ischemic attacks (TIAs) or as a steal effect associated with a general decrease in cerebral blood flow.

The diagnostic workup includes the assessment of cardiovascular risk factors and an ultrasound evaluation of the cranial vessels. Occasionally, vertigo and nystagmus can be provoked by a certain head position. Further investigation and treatment are directed by the findings.

Differentiation is required from positional vertigo and cervical vertigo.

Wallenberg syndrome: The onset of this syndrome is manifested by sudden, severe rotary vertigo, nausea, vomiting, severe ataxia, and additional brainstem signs such as dysphagia and dysphonia. There is ischemia of the lateral medulla oblongata (vertebral artery or posterior inferior cerebellar artery, PICA), leading to crossed signs. The nuclei of cranial nerves V–X are affected ipsilaterally and temperature sensation is affected contralaterally, sparing the face (spinothalamic tract).

Cerebellar infarction (PICA infarction): An infarction in the territory of the posterior inferior cerebellar artery can lead to an acute vestibular disorder that has the features of an acute, unilateral vestibular loss. Differentiation can be difficult based on clinical presentation. Gait and stance ataxia are frequently more pronounced. Magnetic resonance imaging (MRI) can confirm the presence of a cerebellar infarction.

Patients with a cerebellar infarction require particularly close surveillance due to the potential for an intracranial herniation (mass effect) involving the brainstem.

Multiple sclerosis: Vertigo and dysequilibrium occur as the initial manifestations of multiple sclerosis in approximately 5% of cases. The typical signs of a central vestibular disorder are found, often in relatively young patients. A retrocochlear hearing disorder can also be detected in most cases. MRI can demonstrate demyelinated areas in the brainstem, which require further investigation and treatment by a neurologist.

Other causes of central vestibular disorders: Another vascular cause of central vestibular symptoms is basilar artery migraine. A history of typical, paroxysmal headaches suggests the correct diagnosis. Other possible causes of central vestibular symptoms are malformations (basilar impression, Arnold–Chiari malformation), degenerative CNS diseases (Friedreich ataxia, cerebellar degeneration, Parkinson disease, Alzheimer disease), intoxication (alcohol, barbiturates, antiepileptics), and systemic diseases (diabetes mellitus, renal failure, acquired immune deficiency syndrome). Very rarely, vestibular symptoms may be precipitated in a setting of epilepsy (vestibular epilepsy).
Facial Nerve

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14.1 Clinically Relevant Anatomy, Function, and Evaluation of the Facial Nerve

The facial nerve travels an anatomic complex course through the temporal bone, middle ear, and parotid gland. A knowledge of the anatomy, relations, and various physiologic functions of the facial nerve is essential for understanding and diagnosing functional disorders.

Anatomy

The course and various fiber components of the facial nerve are shown schematically in Fig. 14.1. The facial nerve can be divided into six segments:

- **Intracranial:** The frontal branch components of the facial nucleus, unlike the other motor components, are innervated by the left and right corticonuclear tract. Before the facial nerve leaves the brainstem, its motor fibers wind around the abducens nucleus and form the "internal genu" of the nerve. After leaving the brainstem, the facial nerve enters the internal porous acusticus along with the vestibulocochlear nerve.

- **Intrameatal:** Accompanied by cranial nerve VIII, the facial nerve travels through the internal auditory canal to the fundus; there it passes anterosuperiorly through the meatal foramen, leaving the meatus. This is the narrowest point in the bony *fallopian canal* (facial canal) and is the site where the nerve is most likely to become entrapped due to inflammatory swelling.

- **Labyrinthine:** After running a short distance anteriorly, the facial nerve gives off the greater petrosal nerve with its secretory fibers to the lacrimal glands and nasal mucosal glands. The facial nerve turns sharply downward and posteriorly at the geniculate ganglion, forming the first genu.

- **Tympanic:** This segment of the facial nerve runs horizontally through the middle ear, passing above the stapes, to the aditus ad antrum near the lateral semicircular canal. The tympanic nerve segment is covered by a thin bony sheath.

- **Mastoid:** The mastoid segment of the facial nerve forms the second genu by the aditus ad antrum, turning vertically downward at an approximate 90° angle. It courses through the mastoid and leaves its bony canal at the stylomastoid foramen. Just before exiting at this foramen, the facial nerve gives off the *chorda tympani*, which runs back to the middle ear and passes through it. It contains sensory gustatory fibers.

- **Extracranial:** After emerging from the stylomastoid foramen, the facial nerve enters the parotid gland (see 6.1, p. 132), where it branches at the *pes anserinus*. The individual branching pattern within the gland is variable.

- **Extracranial:** After emerging from the stylomastoid foramen, the facial nerve enters the parotid gland (see 6.1, p. 132), where it branches at the *pes anserinus*. The individual branching pattern within the gland is variable.

Function

The cardinal symptom of a facial nerve lesion is paralysis of the mimetic facial muscles, which is the primary focus of the examination. A distinction is drawn between complete paralysis and paresis (incomplete paralysis). A detailed history and clinical examination can often establish the cause of facial nerve paralysis.

Evaluation

**Clinical Examination**

**History:** The onset and course of facial nerve paralysis should be documented. The patient should be questioned about:

- Otologic symptoms and diseases or previous ear surgery
- Trauma
- Neurologic disease
- Tick bites (borreliosis) or evidence of other infections
- Systemic diseases such as diabetes mellitus, cancer, autoimmune diseases, or sarcoidosis

The symptoms of a facial nerve lesion may also include hyperacusis (paralysis of the stapedius muscle), otalgia (irritation of the sensory fibers), gustatory disturbances, and disturbances of lacrimation (dryness, crocodile tears = gustatory lacrimation due to faulty neural regulation).

**Inspection:** A general otolaryngologic status should be obtained, giving particular attention to otologic findings and the function of the other cranial nerves. The presence of other cranial nerve deficits will generally exclude idiopathic facial paralysis. Any changes in the *parotid gland* should be noted. Testing the motor function of the facial nerve should include the following:

- **Frontal branch:** wrinkling the forehead or looking upward. Intact function of the frontal branch compared with the other facial nerve branches indicates a central or supranuclear lesion when paresis is present (see Fig. 14.1).
- **Ophthalmic branch:** rapid blinking (slowed in mild paresis); eyelid closure, spontaneous or against a resistance (weakened in mild paresis); or incom-
complete lid closure (deficit measured in millimeters, seen in severe paresis or paralysis; Fig. 14.2).

- **Oral branch:** baring the teeth, whistling, and inflating the cheeks; air will escape even with mild paresis.

The examiner should also watch for synkinesis—an involuntary associated movement of mimetic muscles accompanying the voluntary movement of other muscles, such as an unintended movement of the oral commissure induced by closing the eyes. This type of synkinesis generally persists as a residual defect following the complete degeneration of nerve fibers (neurotmesis).

**Lacrimal secretion** can be tested by placing strips of filter paper in the lower eyelid and comparing the sides (Schirmer’s test, see 14.1). The amount of tears produced can be helpful in planning corneoprotective therapy.

**Additional tests:** Patients with facial paralysis should undergo laboratory tests to screen for infectious diseases (borreliosis, herpes zoster, syphilis, human immunodeficiency virus [HIV], mononucleosis, toxoplasmosis). **Audimetric testing** (pure-tone, speech and immittance measurements) is necessary due to stapedius muscle involvement and the close proximity of cranial nerve VIII.
14.1 Function tests in facial paralysis

Various function tests can provide limited information on the site of a facial nerve lesion. It is assumed, for example, that if Schirmer’s test shows intact lacrimal gland function, the lesion must be distal to the geniculate ganglion (see Fig. 14.1). Due to the different sensitivities of different-sized nerve fibers and measurement inaccuracies, however, this test is often imprecise for site-of-lesion determination. Imaging studies are much more accurate in this regard. Function tests are considered to have only limited prognostic importance and are seldom performed today. The most important functions are:

- **Stapedial reflex testing** (see 8.4, p.185): If the reflex is absent, the lesion must be proximal to the mastoid segment. The reflex is lost even with mild degrees of nerve damage, as it is mediated by delicate fibers.

- **Schirmer’s test**: Lacrimation is measured with a 5-cm-long strip of litmus paper placed into the conjunctival sac. A 30% reduction in lacrimal secretion relative to the opposite side is considered abnormal.

- **Gustometry**: Taste sensation is evaluated (e.g., by electrogustometry, 4.2, p.76) and compared between the sides. This tests the fibers of the chorda tympani. A right-left discrepancy means that the lesion is proximal to the mastoid segment.

- **Sialometry**: This is a technically demanding test that is rarely practiced. The Wharton duct is catheterized on each side to measure salivary production by the submandibular glands. A discrepancy between the sides means that the lesion is proximal to the origin of the chorda tympani.

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### Site-of-Lesion Determination and Imaging Studies

Today, the best and most widely used topodiagnostic tests are computed tomography (CT) and magnetic resonance imaging (MRI). Inflammatory facial nerve lesions can be demonstrated by MRI after gadolinium contrast administration. Otogenic and traumatic facial paralysis should always be evaluated by thin-slice bone-window CT scanning of the temporal bone. If the site of the lesion is not visualized in one plane, scans should be acquired in a second plane (axial and coronal; temporal bone CT, see p.233).

### Electrical and Magnetic Testing

The importance of electrical and magnetic testing lies in the objective detection and quantification of facial paralysis. Repeated tests are useful for defining the degree of facial nerve damage and assessing the prognosis for recovery. From a pathophysiologic standpoint, three degrees of facial nerve fiber injury are distinguished:

1. **Neurapraxia**: disruption of axonal transport without degeneration.
2. **Axonotmesis**: wallerian degeneration of the myelin sheath with an intact perineurium. There is complete paralysis of the affected fibers, but regeneration of the axon is also complete.
3. **Neurotmesis**: wallerian degeneration and loss of the perineurial sheath with complete paralysis of the affected fibers. Regeneration is unpredictable. The outcome is residual dysfunction with synkinesis and persistent palsy.

#### Electroneurography (ENoG)

This test involves supramaximal stimulation of the nerve trunk and measuring the muscular response with surface electrodes. The degree of degeneration is expressed as a percentage relative to the healthy side ($\approx 100\%$). More than 90% degeneration of the nerve fibers is a poor prognostic sign in terms of complete recovery.

#### Electromyography (EMG)

The electrical potentials of the mimetic muscles are measured with needle electrodes. Recordings are made during spontaneous and voluntary muscular activity for the objective detection of paralysis and reinnervation. EMG is also used for the intraoperative monitoring of facial nerve function during parotid and otologic surgery and intracranial operations.
Magnetic stimulation: The intracranial portion of the facial nerve can be stimulated with a magnetic coil to test function over the entire course of the nerve. If the nerve is responsive to stimulation when facial paralysis is present, there is a good prognosis for recovery. If the nerve is unresponsive, a prognostic assessment cannot be made.

Diagnosis and Management of Facial Paralysis

Motor paralysis is the most important and by far the most common symptom of facial nerve pathology. Generally, it is easily determined from the history whether the paralysis is traumatic or nontraumatic. With nontraumatic paralysis, the function of the frontal branch can be tested to determine whether the lesion is central (i.e., supranuclear) or peripheral. This is followed by various diagnostic steps and considerations that are outlined in Fig. 14.3. It is particularly important to differentiate between complete paralysis and incomplete paralysis (paresis). Before instrumented tests are performed, the cause of facial paralysis can almost always be determined from a detailed history and clinical examination of the external ear, middle ear, hearing, vestibular function, other cranial nerve functions, and the parotid gland. The most frequent causes of peripheral facial paralysis are listed in Table 14.1.
14.2 Clinical Aspects of the Facial Nerve

The most common type of facial nerve paralysis is idiopathic or Bell’s palsy, followed by traumatic and inflammatory otogenic forms. Traumatic causes can usually be established by the history, giving particular attention to possible infectious and inflammatory otogenic causes. Idiopathic facial paralysis is a diagnosis of exclusion.

Inflammatory Changes

Idiopathic Facial Paralysis (Bell’s Palsy)

Synonyms: cryptogenic, rheumatic or ischemic facial paralysis

Definition: Bell’s palsy is a unilateral, peripheral facial paralysis of acute onset that has no discernible cause and does not involve any other cranial nerves.

Occurrence: Bell’s palsy is the most common form of facial paralysis. The incidence is approximately 20 per 100,000 population.

Pathogenesis: Unknown. Theories include infection and inflammation (viral, autoimmune), vascular ischemia, and constitutional factors. Idiopathic facial paralysis is more common in diabetic patients and in pregnancy (third trimester).

Symptoms: Often the initial symptom is retroauricular pain, followed by unilateral peripheral facial paralysis in which the frontal branch is equally affected. The paralysis is partial in 30% of cases and complete in 70% of cases. It develops within a few days (2–5 days) and has no systemic manifestations. The clinical features may include hyperacusis (stapedius muscle paralysis), dysgeusia, and decreased lacrimation.

Diagnosis: The diagnosis is one of exclusion based on the typical clinical course and the absence of an identifiable cause of the paralysis.

Differential diagnosis: Otogenic and infectious causes of peripheral facial paralysis should be excluded (see Table 14.1, p.292). Bell’s palsy requires differentiation from Melkersson-Rosenthal syndrome (peripheral facial paralysis with recurrent swelling of the face and lips and a fissured tongue).

Course and prognosis: Partial paralysis always resolves completely within a few weeks. Recovery from complete paralysis takes longer (months) and is complete in only about 60–70% of cases. Approximately 15% of patients are left with troublesome residual palsy and/or synkinesis.

Complications: The most serious complication is corneal damage due to lagophthalmos, ectropion, and/or decreased lacrimation. The best preventive measures are moisturization and ophthalmologic follow-up.

Treatment: Treatment with corticosteroids, occasionally combined with rheologic or antiviral agents, is generally recommended although definite efficacy has not been confirmed. Surgical decompression of the facial nerve is also of unproven benefit and carries a considerably higher risk. It is important to protect the cornea with a watchglass eye dressing, moisturizing eye drops and ointments, and if necessary by tarsorrhaphy or by placing a gold or titanium weight in the upper eyelid.

Inflammatory Otogenic Facial Paralysis

Definition: This is a type of facial paralysis caused by the spread of an infectious or other inflammatory process from the ear and temporal bone to the facial nerve.

Occurrence: Facial paralysis can occur as a relatively rare complication in all forms of otitis and osteitis/osseomyelitis of the temporal bone, particularly in cases with cholesteatoma, subacute mastoiditis in pediatric patients, and advanced necrotizing otitis externa (see p.221).

Etiology: The functional damage is caused by a direct toxic insult, inflammatory epineurial edema and pressure, and in some cases by osteitis.

Symptoms: Otologic symptoms are usually the dominant findings, and facial paralysis occurs as a complication. A chronic process (cholesteatoma) may have an insidious onset.

Diagnosis: The otoscopic findings suggest the correct diagnosis. Further investigation requires an audiologic examination and CT scans of the temporal bone.

Differential diagnosis: Differentiation is required from infectious diseases, especially herpes zoster oticus, and from tumors of the lateral skull base, temporal bone, and parotid gland.
Complications: The complications depend largely on the underlying disease.

Treatment: Except for facial paralysis in the setting of acute otitis media (antibiotics), treatment consists of prompt surgical exposure of the nerve and appropriate antibiotic therapy. Corticosteroids are also administered to reduce edema.

Course and prognosis: These depend on the degree of paralysis, the underlying disease, and the timing of treatment. The less complete and more acute the paralysis and the earlier treatment is initiated, the better the prognosis.

Facial Paralysis Secondary to Infection

It should always be determined whether facial paralysis has an infectious etiology, since infections are often amenable to specific treatment. The most frequent causes of infectious facial paralysis are herpes zoster oticus and borrelia. A history of tick bite can usually be elicited in patients with borrelia. This infection can be detected by serologic testing and can be treated with antibiotic. Other infections that may be associated with peripheral facial paralysis are meningitis, Guillain-Barré syndrome, poliomyelitis, and HIV infection.

Traumatic Facial Paralysis

Definition: A complete or partial facial nerve injury may result from a traumatic rupture or stretch injury, nerve compression (by hematoma or bone fragments), trauma-induced swelling, or thermal injury (from a drill during otosurgery). The paralysis may be immediate (a complete, instantaneous traumatic lesion) or delayed (progresses over several days following the trauma).

Pathogenesis: Facial paralysis may be caused by a temporal bone fracture, facial trauma (sharp or blunt), surgical trauma (surgery of the petrous bone or parotid gland), or obstetric trauma. Depending on the degree of the trauma, the effects can range from a transient conduction block (neurapraxia) or nerve damage with an intact perineurium (axonotmesis) to a complete nerve transection (neurotmesis).

Symptoms: Traumatic facial paralysis does not occur in isolation but is accompanied by other symptoms relating to the trauma.

Diagnosis: A history of trauma usually suggests the cause of the facial nerve paralysis. The paralysis is difficult to detect only in patients who are comatose. In patients who have sustained a temporal bone fracture, facial nerve function should always be tested and documented so that immediate paralysis can be distinguished from paralysis of delayed onset. Thereafter the course is monitored by electrodiagnostic testing (neurography and EMG).

It is also important to determine the site of the lesion. This requires thin-slice CT scanning, especially in patients with a temporal bone fracture (see p. 300).

Course: The course depends on the extent of the injury. A nerve rupture causes immediate and complete paralysis that does not tend to resolve over time. With milder injuries, the paralysis often increases over a period of several days and then tends to resolve spontaneously. It is important, therefore, to document the course of traumatic facial paralysis clinically and by electrodiagnostic testing.

The indication for surgical decompression of traumatic facial paralysis depends on the course. For this reason, the function of the facial nerve and the course of the paralysis should always be carefully documented in patients with temporal bone fractures.

Complications: Complications may consist of corneal damage, residual palsy, and complications relating to the primary trauma.

Treatment: Every case of immediate paralysis should be surgically explored. The timing depends on clinical circumstances. A severe or badly damaged nerve is either directly reapproximated or microsurgically repaired with an interposed nerve graft (great auricular nerve or sural nerve). Delayed paralysis is treated initially with corticosteroids to reduce edema. If neurography indicates more than 90% degeneration or if CT indicates compression by bone fragments, the nerve is surgically explored. This is also done if other indications for temporal bone surgery exist (cerebrospinal fluid leak, ossicular chain disruption). It is usually sufficient to decompress the nerve.
Lateral Skull Base

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15.1 Anatomy, Evaluation, and Surgery of the Lateral Skull Base

Both in the region of the paranasal sinuses (the anterior skull base) and the mucosa-lined cells of the temporal bone (the lateral skull base), the base of the skull borders on air-filled cavities that communicate with the external environment. While the anterior skull base is discussed in the chapters on the paranasal sinuses (Chapters 1–3), this chapter takes a more detailed look at the lateral skull base. This focuses on the anatomy, symptomatology, evaluation, and treatment of this anatomical region.

Anatomy of the Lateral Skull Base

Topography

The close proximity of the intracranial cerebrospinal fluid (CSF) space to the temporal bone air cells, and thus to the outside world, accounts for the special clinical features and management of diseases and injuries in this region (Fig. 15.1). The complex, variable anatomy of a great many vitally and functionally important structures can lead to serious sequelae and can hamper surgical treatment.

Depending on the pneumatization of the temporal bone, there are places where only a paper-thin layer of bone separates the dura mater from the mucosa of the temporal bone air cells. Venous connections exists between the dura and the mucosa, particularly at the roof of the mastoid ( tegmen ) and in the epitympanum, both of which border on the middle cranial fossa and thus on the temporal lobe of the brain. The posterior surface of the mastoid borders on the posterior cranial fossa and cerebellum.

Neurovascular Connections

In the lateral skull base region, there are many neural and vascular connections between the cranial interior on the one hand and the neck and face on the other (Fig. 15.2). The principal sites at which neurovascular structures pass through the bony skull base are as follows:

- The internal auditory canal with the facial nerve (cranial nerve VII), which reemerges from the temporal bone at the stylomastoid foramen (see 14.1, p. 290), the vestibulocochlear nerve (cranial nerve VIII), and the labyrinthine artery.
- The jugular foramen with the internal jugular vein, where the sigmoid sinus drains in a siphon-like termination to form the jugular bulb. The foramen is traversed anteriorly by the basal cranial nerves, the glossopharyngeal nerve (cranial nerve IX), the vagus nerve (cranial nerve X), the accessory nerve (cranial nerve XI), and the hypoglossal nerve (cranial nerve XII).
- The foramen lacerum with the internal carotid artery, which runs horizontally forward along the anterior border of the tympanic cavity and enters the skull by the sphenoid sinus.
- The foramen ovale with the mandibular nerve ( synonym : third branch of cranial nerve V).
- The foramen spinosum with the middle meningeal artery, a branch of the external carotid artery.
Symptoms and Evaluation

Clinical Examination

The symptoms of injuries and diseases of the lateral skull base are variable and diverse due to the specific anatomy of this region. The dominant features are functional disorders of cranial nerves VII–XII, and therefore the testing of these nerves is an essential part of the clinical examination of the lateral skull base. A complete ear, nose, and throat status should always be established. For example, examination of the oral cavity (see 4.2, p. 76) is useful for assessing the functional status of the hypoglossal and glossopharyngeal nerves, and indirect laryngoscopy (see 17.2, pp. 346–348) can assess the function of the recurrent laryngeal nerve, which arises from cranial nerve X.

The clinical findings provide a basis for identifying various characteristic syndromes, which can vary considerably in their severity.

Cochleovestibular syndrome: The hallmarks of the cochleovestibular syndrome are sensorineural hearing loss, tinnitus, and vestibular symptoms such as dysequilibrium and vertigo. Vestibular nystagmus may be present. It is often difficult to distinguish clinically between a cochlear and retrocochlear disorder, making it necessary to rely on further audiologic testing (see 12.1, General Clinical Aspects of Cochlear and Retrocochlear Disorders, pp. 257–259) and imaging studies.

Jugular foramen syndrome: Cranial nerves IX, X, and XII are affected in various combinations. Vagus nerve involvement, manifested by unilateral vocal cord paralysis and dysphagia, is the dominant clinical finding in many cases. The history often raises suspicion of aspiration. Glossopharyngeal nerve involvement causes velum paralysis with deviation of the soft palate toward the healthy side (“backdrop sign,” see Fig. 4.7b, p. 77). Hypoglossal nerve involvement leads to paralysis of the tongue, which deviates toward the affected side (see Fig. 4.7a, p. 77), in addition to lingual atrophy. Fasciculations of the lingual muscles are also common.

Petros apex syndrome: Gradenigo described the classic triad of purulent otorrhea, stabbing ipsilateral facial pain (trigeminal nerve irritation), and diplopia due to abducens nerve palsy in the setting of a petrous apex abscess (petrous apicula). Today the petrous apex syndrome is very rarely encountered in its classic form.

The combination of trigeminal nerve irritation with paralysis of the abducens nerve and occasionally of the oculomotor nerve should raise suspicion of a petrous apex lesion and requires immediate investigation by computed tomography (CT) or magnetic resonance imaging (MRI).
Most patients also complain of a deep headache, and other symptoms such as cochleovestibular disorders, facial paralysis, or meningitic signs are often present as well. The cause may be a suppurative inflammatory process (e.g., a petrous apex abscess or osteomyelitis), a congenital cholesteatoma of the petrous apex, or infiltrative lesions such as lymphomas or metastases.

**Imaging Studies**

Every combined functional disorder of the laterobasal cranial nerves that cannot be diagnosed clinically requires further investigation by imaging procedures, primarily CT and MRI. Both CT and MRI are often necessary for the precise delineation and differentiation of findings (Fig. 15.3). CT is best for defining the infiltration and destruction of bony structures. CT scans are of major importance in making a differential diagnosis and also in therapeutic and preoperative planning. MRI is often better for defining and differentiating the lesion itself, which is usually a tumor or inflammatory process. Because the disease process may be in very close proximity to major vessels, conventional angiography is also occasionally required. Angiography can also be used therapeutically (e.g., embolization) and therefore should be appropriately planned and coordinated.
Surgery of the Lateral Skull Base

General

Because symptoms are influenced not just by the nature of the disease but to a large extent by its location, and even imaging studies cannot reliably distinguish tumors from inflammatory processes in all cases, often a definitive diagnosis can be made only intraoperatively.

Three main routes of approach are available for surgery of the lateral skull base (Fig. 15.4):

- **Intracranial–intradural**: These are neurosurgical approaches, of which the suboccipital or retrosigmoid approach to the cerebellopontine angle is most commonly used.
- **Intracranial–extradural** (trans-temporal): This is an otosurgical approach for exposing the surface of the petrous pyramid through a temporal craniotomy. The dura is separated from the surface of the petrous pyramid and elevated away from it with the temporal lobe. This approach is used, for example, in the surgical treatment of temporal bone fractures or tumors of the internal auditory canal (intra- and vestibular schwannomas, see p. 308).
- **Extracranial–extradural** (trans-mastoid and infra-temporal): This is an otosurgical approach that is directed through the mastoid and/or other portions of the temporal bone.

Extracranial–Extradural Approaches

**Transmastoid operation**: The mastoid air cells can be completely excavated to gain access to broad portions of the lateral skull base including the sigmoid sinus and the dura of the posterior and middle cranial fossa. The vestibular labyrinth can also be ablated (= trans-mastoid–trans-labyrinthine operation) to access the internal auditory canal and cerebellopontine angle. This approach is used to remove vestibular schwannomas that have already caused deafness.

**Infratemporal operation**: The basis of this operation is a subtotal petroectomy (see 11.2, p. 245). In this approach the bony external auditory canal is removed in addition to the mastoid, and the tympanic cavity is broadly opened along with the mastoid cells. Both the cutaneous ear canal and the eustachian tube are permanently occluded. In most cases, the facial nerve is exposed from the geniculate ganglion to the stylomastoid foramen and transposed anteriorly, providing access to the jugular bulb, the jugular foramen and, further anteriorly, the internal carotid artery. The operation can be extended anteriorly into the deep infratemporal regions, to retromaxillary structures, and to the orbit by removing the zygomatic arch and disarticulating the temporomandibular joint. The postoperative cavity is usually obliterated with abdominal fat.
15.2 Trauma to the Lateral Skull Base

Injuries are a frequent and important cause of lateral basal pathology, along with inflammations and (usually benign) neoplasms (see 15.3, pp. 306–309). Fractures of the lateral skull base consist mainly of indirect burst fractures caused by blunt head trauma. Direct trauma, as from a gunshot injury, is much less common. Lateral basal fractures generally involve the middle ear and less commonly the inner ear (see 12.2, p. 262). They are usually easy to differentiate from isolated middle ear trauma (see 11.4, p. 250) based on the history and clinical findings.

Laterobasal Fractures

Synonym: temporal bone fractures

The primary examination and the initial findings and diagnosis can be very important in dealing with complications that may arise later. The accurate documentation of findings is essential, therefore.

General

The lateral skull base is a frequent site of occurrence of basal skull fractures. The stresses generated by the deformation of the skull can lead to characteristic burst fractures of the petrous pyramids and neighboring bones. The trauma is commonly associated with fractures of the calvaria and also with brain injury. Below we shall review the complications that may arise in all types of laterobasal fractures. Then we shall consider specific fracture types and their associated features, diagnosis, and management.

Cerebrospinal fluid leak:

Pathogenesis: As long as the dura mater is intact and the injuries are confined to bony structures and the mucosa of the pneumatized cavities, the interior of the skull will remain sealed and protected. But because the dura is firmly attached to the bone in some places, it may become torn when injuries are sustained, allowing CSF to leak into the cells of the temporal bone. With extensive injuries to the lateral skull base, both dura and brain tissue may prolapse into the air cells of the temporal bone.

Symptoms: A CSF leak is manifested by the following clinical signs:

- Air in the intracranial cavity (pneumoencephalos)
- CSF dripping as a watery discharge from the ear (CSF otorrhea) or nose (CSF rhinorrhea, see also 3.6, pp. 44–46)
- Infection, which may be acute (meningitis) or chronic (brain abscess)

Diagnosis: The clinical discovery of a CSF leak can be confirmed by laboratory tests, imaging studies, or intraoperatively.

- CSF leak can be diagnosed by sampling the watery discharge and sending it for laboratory analysis. Glucose test strips can be used to make a quick assessment of the sugar content of the fluid. A more accurate test involves the assay of CSF-specific enzymes such as β₂-transferrin or β tracers, which can also be detected by inserting a foam wick into the nose or ear to collect discharge that is not clinically apparent.
- If CT demonstrates pneumoencephalos (intracranial air), this indicates at least a transient CSF leak.
- A profuse CSF leak is easily detected at operation, but this can be difficult with a smaller leak. In these cases the leaking fluid can be clearly visualized endoscopically with a blue filter following the intra- thecal injection of sodium fluorescein.

Treatment: A CSF leak in the lateral skull base is associated with a relatively low risk of acute infection (meningitis) or chronic infection (brain abscess). In many cases, moreover, the leak will stop spontaneously and the fistula will become sealed. Generally, then, a CSF leak requires surgical treatment only if it persists for 5–7 days, or if complications such as meningitis occur. General antibiotic prophylaxis is controversial when a CSF leak is present, as it cannot reliably prevent meningitis and could promote the selection of resistant strains.

Cochleovestibular symptoms: Hearing loss following a laterobasal fracture may be a conductive hearing loss caused by fluid (blood or CSF) in the tympanic cavity or by ossicular dislocation, or it may be a sensorineural hearing loss or deafness caused by a fracture of the labyrinth. The latter is associated with failure of the vestibular organ and its clinical manifestations such as severe vertigo, nausea, and vomiting.

The mechanism of the injury may also cause a labyrinthine concussion or contusion, which usually occurs in association with acute noise trauma (see 12.2, pp. 260–262).

Facial nerve function: With a fracture of the lateral skull base, facial nerve function is commonly affected along with cochleovestibular function (see 14.2, p. 295).
When a temporal bone fracture has occurred, facial nerve function should be tested and carefully documented during the initial days and weeks following the injury.

The treatment of traumatic facial nerve paralysis, and especially the indication for surgical repair, depends mainly on the course and degree of the paralysis. If the facial paralysis is present immediately after the injury (early paralysis), it is likely that the nerve has sustained direct trauma and the prognosis is poor. Surgical treatment is often necessary. Paralysis that occurs 24 hours or more after the trauma (late paralysis) very often resolves spontaneously, especially when the paralysis is incomplete.

Since the detection of early paralysis has major prognostic significance, the initial emergency examination should be conducted with meticulous care.

It is difficult to test facial nerve function in the unconscious trauma patient, but occasionally a grimace can be elicited in response to a painful stimulus. The results of the examination should be documented along with any uncertainties regarding the findings. Facial paralysis that is diagnosed later should be interpreted with particular care (late paralysis or undiagnosed early paralysis).

**Classification of Temporal Bone Fractures**

Typical burst fractures involving the petrous pyramid are known also as *pyramid fractures*. There are also isolated fractures of the temporal squama and various combinations that include other calvarial fractures. The diagnosis and classification of a temporal bone fracture can often be accomplished clinically. Axial and coronal thin-slice CT scans should be obtained when complications arise and prior to a surgical procedure. CT can at least partially define the course of the fracture lines. Several different types of temporal bone burst fracture may be encountered:

- **Squama–mastoid fractures**: The fracture is confined to the temporal squama and mastoid air cells. The auditory canal and tympanic cavity may also be involved.
- **Longitudinal temporal bone fracture**: The fracture runs along the petrous bone and petrous pyramid (see Fig. 15.5, left). The typical fracture line runs along the auditory canal, continues across the mastoid roof and tegmen tympani to the carotid canal, and terminates in the sphenoid sinus.
- **Transverse temporal bone fracture**: The fracture line runs transversely across the petrous bone or petrous pyramid (see Fig. 15.5, right) along the internal auditory canal and/or through the labyrinth.
- **Isolated mental fracture**: This fracture is most often caused by a posterior displacement of the mandibular condyle. The typical mechanism of the injury is a fall onto the chin. The fracture penetrates the posterior wall of the glenoid fossa and the anterior wall of the ear canal and is often associated with a condylar neck fracture.

Besides these more or less typical temporal bone fractures, there are also atypical fracture patterns involving other bony structures of the lateral skull base such as the occipital bone (lateral or clivus) and sphenoid bone. These patterns are particularly common with direct or high-energy trauma.

**Longitudinal Temporal Bone Fracture**

**Definition**: This fracture runs along the external auditory canal and the anterior border of the petrous pyramid (Fig. 15.6).

**Epidemiology**: The longitudinal temporal bone fracture is the most common burst fracture affecting the lateral skull base.

**Pathogenesis**: The fracture is caused by a diffuse, lateral traumatizing force. It may result from a fall in which the head is struck without any other injuries, but it is associated with brain trauma in many cases.

**Symptoms**: Aural discharge is present and may consist of pure blood or blood mixed with CSF (CSF otorrhea). Hearing loss is also present. Occasionally a slightly bloody rhinorrhea may also occur due to involvement of the sphenoid sinus. Approximately 10–20% of cases exhibit facial paralysis, which is usually of delayed onset (see above).
**Diagnosis:** The otoscopic findings usually establish the diagnosis when combined with the history and auditory tests. Otoscopy shows tearing of the meatal skin and tympanic membrane, with bleeding into the ear canal. Generally, the posterosuperior quadrant of the tympanic membrane and superior wall of the ear canal are affected. Blood and cerumen can hamper the examination. Cleansing of the ear canal should be left to a specialist.

The examiner who suspects a temporal bone fracture should not perform any manipulations in the ear canal and should particularly avoid irrigating the ear.

On **clinical auditory testing,** the Weber test is lateralized to the affected ear and the Rinne test is usually but not always negative. Further evaluation consists of detecting or excluding CSF otorrhea (see p. 302), defining the fracture line by **thin-slice CT scanning** of the temporal bone, the objective testing of facial nerve function by **neurography,** and auditory testing by **pure-tone audiometry.** All of these tests are useful in specific cases, but they are mandatory only in cases that require surgical exploration.

**Differential diagnosis:** A longitudinal temporal bone fracture mainly requires differentiation from an isolated fracture of the ear canal when otorrhea is present.

The latter type of injury mainly involves the anterior wall of the ear canal, leaving the superior wall intact. Occasionally, blood from a head wound that has flowed into the ear canal is mistaken for evidence of a temporal bone fracture.

**Complications:** Possible **early complications** are meningitis in the presence of a CSF leak, otitis media in the presence of a tympanic membrane perforation, and facial nerve paralysis. Facial nerve damage or a perforated tympanic membrane rarely leads to **permanent sequelae.** A more frequent complication is conductive hearing loss caused by fracture or dislocation of the auditory ossicles; the incus is most commonly affected. Surgical exploration of the tympanic cavity with an ossiculoplasty is indicated in patients with permanent conductive hearing loss. Rare **late complications** are stenosis of the ear canal due to scarring or a post-traumatic cholesteatoma caused by the ingrowth of meatal skin into the middle ear through a fracture.

**Treatment:** Whenever possible, the ear should be covered with a sterile dressing and left alone. Facial paralysis is treated with corticosteroids. Surgical exploration is necessary in patients with infectious complications or a persistent CSF leak and in selected cases of facial paralysis.
Transverse Temporal Bone Fracture

**Definition:** A transverse temporal bone fracture runs across the petrous pyramid along the internal auditory canal and/or through the labyrinth (Fig. 15.7).

**Epidemiology:** Transverse temporal bone fractures are much less common than longitudinal fractures.

**Pathogenesis:** This burst fracture is caused by a traumatizing force in the frontal plane. It may occur in isolation without associated injuries. The fracture line runs medial to the external auditory canal and tympanic membrane plane, with the result that these structures are not directly affected and there is no blood or CSF discharge from the ear canal.

**Symptoms:** The foremost symptom is acute vestibular dysfunction with severe vertigo, generally accompanied by nausea and vomiting. Severe hearing loss or even deafness occurs acutely on the affected side but often goes unnoticed. Otorrhea is absent, but there may be CSF rhinorrhea through the eustachian tube. Facial paralysis occurs in up to 50% of cases.

**Diagnosis:** On clinical examination, the Weber test is lateralized to the healthy ear and there is spontaneous nystagmus toward the healthy side (due to vestibular loss). Otoscopy typically shows signs of hemotympanum. The fracture line is defined by thin-slice CT scans of the temporal bone in two planes.

**Differential diagnosis:** Autonomic symptoms caused by the vestibular disorder are often misinterpreted as central nervous system signs of cranioencephalic trauma. If CSF rhinorrhea is present, the differential diagnosis should include a frontobasal fracture. Hemotympanum can also occur in this injury because blood from epistaxis may enter the tympanic cavity through the eustachian tube.

**Course and complications:** The risk of meningitis is greater with a transverse temporal bone fracture, and there is less likelihood that the fistula will close spontaneously. It is extremely unlikely that hearing or vestibular function will be recovered. In most patients, the loss of vestibular function is well compensated by central mechanisms over time. Facial paralysis associated with a transverse fracture also has a poorer prognosis than in patients with a longitudinal fracture.

**Treatment:** A CSF leak associated with a transverse fracture is a better indication for surgical closure than a leak caused by a longitudinal fracture—often there is a gaping fracture line, and it is less likely that the leak will close spontaneously. Otherwise treatment is conservative, with emphasis placed on mobilizing the patient and restoring vestibular functions.

**Direct Trauma to the Lateral Skull Base**
Direct injuries to the lateral skull base are less common than burst fractures, although both types may be combined. A depressed fracture of the temporal squama or mastoid generally leads to associated external injuries such as lacerations or injuries to the external ear or a squama-mastoid fracture. Gunshot injuries to the lateral skull base and other direct trauma generally require surgical exploration. The first priority is the repair of injured vascular structures. After that, an effort is made to preserve the function of the cranial nerves, external ear, and middle ear.
15.3 Inflammations and Tumors of the Lateral Skull Base

Inflammations involving the lateral skull base most commonly originate from the middle ear. Tumors of the lateral skull base are generally rare and most are benign. Both inflammations and tumors are often manifested by varied and nonspecific signs relating to the various functional structures of the lateral skull base. As a result, the diagnosis of tumors in particular is often delayed, and the process may spread to neighboring structures. Accordingly, a continuum exists between severe inflammations and tumors of the ear canal, middle ear, intracranial cavity, and lateral skull base. The surgical treatment of these processes is challenging and often requires an interdisciplinary approach with neurosurgical consultation.

Inflammations

Otitis media is usually a tubogenic condition. It is among the most common inflammations and infections that affect the lateral skull base region (see 11.3, pp. 238–242). Otogenic complications with potential extensive involvement of the lateral skull base have become rare in developed countries. The most frequent cause of an inflammatory process arising from the middle ear and spreading to the lateral skull base in these countries is cholesteatoma (see pp. 243–244). Other, less common local or systemic diseases such as Wegener’s granulomatosi also play a role. Infections can also occur as a sequel to injuries. The most hazardous is meningitis, which may be manifested during the initial days or weeks following an injury (early meningitis). In other cases, months or even years may elapse between the trauma and the development of an infectious complication (delayed meningitis).

Tumors of the Temporal Bone

Virtually any structure can serve as a nidus for tumor growth. Some of the more common tumors of the temporal bone are discussed below in greater detail. The diagnosis of temporal bone tumors relies almost exclusively on imaging procedures, especially neuroradiologic studies.

Paraganglioma

Synonyms: glomus tumor, chemodectoma, nonchromaffin paraganglioma

Sites of occurrence: This tumor arises from the paraganglia of the temporal region, most commonly in the area of the jugular bulb and along the neural plexus of the tympanic cavity (tympanic plexus). Paragangliomas may be located in the middle ear (glomus tympanicum) or on the jugular bulb (glomus jugulare). They can also develop outside the temporal bone region from paraganglia at the carotid bifurcation (glomus caroticum) and along the vagus nerve (glomus vagale). The tumors often extend toward the temporal bone region. Once they have reached a certain size, their site of origin can no longer be determined with certainty.

Epidemiology: Paragangliomas are the most common tumors of the middle ear and adjacent lateral skull base. Their overall incidence is low, however.

Symptoms: Paragangliomas in the tympanic cavity are sometimes manifested early by a pulsatile tinnitus and conductive hearing loss. By contrast, paragangliomas of the jugular bulb often become symptomatic at a later stage due to the development of basal cranial nerve deficits.

Diagnosis: MRI and CT can both demonstrate paragangliomas. CT is necessary for the detection of bone destruction in the tympanic cavity or about the jugular foramen (Fig. 15.8).

On audiologic testing, unilateral conductive hearing loss is detected at an early stage in patients with a glomus tympanicum tumor. This is seen later with a glomus jugulare tumor. Sensorineural hearing loss may also occur due to infiltration of the inner ear.

Otoscopy may demonstrate the typical finding of a bluish mass in the lower part of the tympanic cavity. This occurs only if the tumor has reached the mesotympanum.

Angiography (Fig. 15.9) is helpful in determining both the etiology and extent of the lesion. It should be done as an immediate preoperative study and should include embolization.

Treatment:

The preoperative angiographic embolization of a tumor can reduce intraoperative blood loss.

Paragangliomas of the tympanic cavity are relatively easy to remove in their early stage by a middle ear operation.

The surgical treatment of a glomus jugulare tumor is challenging and generally requires a subtotal petrosectomy (see 11.2, p. 245).
Other Tumors of the Lateral Skull Base

The symptoms and findings that are associated with tumors depend on their growth rate, invasive potential, and location. Generally, a tumor can be diagnosed with imaging procedures, but often its etiology can only be determined histologically by biopsy or operative exposure.

Possible benign tumors include primary cholesteatoma or epidermoid, which may occur in the temporal bone and especially at the petrous apex (Fig. 15.3, p. 300), mucosal tumors (such as papilloma and adenoma), tumors of connective tissue and bone (such as meningioma and fibrous dysplasia), and eosinophilic granuloma in the setting of Langhans-cell histiocytosis.

Primary malignant tumors are carcinomas of the mucosa, sarcomas, and lymphomas. The lateral skull base may also be affected by skeletal metastases from various other tumors such as breast cancer and bronchial or renal carcinoma.

Tumors of the Internal Auditory Canal and Cerebellopontine Angle

The most common identifiable cause of a retrocochlear hearing disorder (see 12.1, p. 259) is a tumor of the internal auditory canal or cerebellopontine angle. These lesions can produce a unilateral cochleovertebral syndrome by exerting pressure on cranial nerve VIII and the labyrinthine artery. The most common of these tumors is vestibular schwannoma, known also as acoustic neurinoma (approximately 80% of cases), followed by cerebellopontine angle meningioma.

Vestibular Schwannoma

Synonym: acoustic neurinoma

Definition and forms: Vestibular schwannoma is a generally slow-growing, benign tumor that arises from the Schwann cells of cranial nerve VIII. The vestibular nerve is most commonly affected. Rarely, the tumor may arise from the cochlear nerve or may form within the labyrinth.

Two forms are distinguished clinically: medial and lateral. Medial tumors arise from the intracranial part of cranial nerve VIII—i.e., in the actual cerebellopontine
angle, while lateral tumors are located in the internal auditory canal.

**Epidemiology:** Though it is the most common tumor of the posterior cranial fossa, vestibular schwannoma has a low overall incidence. It seems to become more common with ageing, and there are probably a considerably greater number of vestibular schwannomas that are asymptomatic and therefore undiagnosed. Rarely, these tumors occur in the setting of neurofibromatosis type 2 (15.1).

**Symptoms:** The clinical hallmark of vestibular schwannoma is a unilateral hearing disorder, which may consist of tinnitus, hearing loss, or dysacusis.

Medial tumors may reach a considerable size before becoming symptomatic. Lateral tumors at an early stage evoke hearing disorders.

The hearing disorder may have a sudden onset (sudden sensorineural hearing loss) or gradual onset and may show improvement, which is usually transitory.

Recovery from sudden sensorineural hearing loss does not exclude vestibular schwannoma, therefore. This tumor should always be included in the differential diagnosis of a unilateral hearing disorder.

Vestibular symptoms such as acute vertigo or dysequilibrium are less common. Impairment of facial nerve function is unusual, even with very large tumors. Medial schwannomas can occasionally produce trigeminal nerve symptoms such as facial pain or numbness in the jaw. Large tumors also produce signs of brainstem compression and/or hydrocephalus with ataxia, nausea, and vomiting.

**Diagnosis:** Clinical examination reveals the typical signs of a unilateral cochleovestibular disorder. Spontaneous nystagmus is rarely present. Audiometry usually demonstrates retrocochlear impairment with a lengthening of auditory brainstem response latencies (see 8.4, p. 187).

The diagnosis is established by MRI (Fig. 15.10). Classically the tumor enhances after gadolinium administration, but even without contrast medium it can be detected in thin-slice sequences of the posterior fossa with three-dimensional data acquisition.

**Treatment:** Large tumors that exceed 2.5–3.0 cm in diameter constitute a vital indication for surgical removal. Small tumors less than 1 cm in diameter may be managed by a “wait-and-see” approach with regular follow-ups, or the lesion may be treated by stereotactic radiosurgery or open surgery. The indication depends on the quality of hearing and the age and wishes of the patient. Tumors in the 1.0–2.5 cm range...
should generally be treated by open surgery or radio-
surgery.

Other Tumors of the Cerebellopontine Angle

The second most common cerebellopontine angle tu-
mor is meningioma, which arises either from the pos-
terior surface of the petrous pyramid or from the ten-
torium. Less common lesions include hemangiomas
and lipomas. The clinical presentation and treatment
are basically the same as for vestibular schwannomas.
Postoperative function depends critically on the loca-
tion of the tumor and its relation to the cranial nerves.

15.1 Neurofibromatosis 2

A rare special form of vestibular schwannomas is neurofi-
bromatosis 2 (synonym: central neurofibromatosis), an in-
herited autosomal-dominant disease with a genetic defect
in chromosome 22, characterized by bilateral vestibular
schwannomas and other tumors in the cranial cavity and spi-
nal canal. Differentiation is required from neurofibromato-
sis 1 (synonym: von Recklinghausen disease), which involves
a defect on chromosome 17 and is mainly characterized by
neurofibromas of the skin and café au lait spots.
The management of neurofibromatosis 2 is difficult and must
be tailored to the situation in order to preserve auditory
function for as long as possible. Cases with complete bilat-
eral deafness can sometimes be rehabilitated by direct electri-
cal stimulation of the cochlear nucleus (brainstem implant).
External Neck

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16.1 Anatomy of the External Neck

As the connecting link between the head and trunk, the neck contains a number of vitally important neurovascular structures (arteries, veins, nerves, lymphatics) within a very confined space. Its musculoskeletal system must protect the neck while also allowing maximum mobility of the head and the coordinated processes of ventilation, deglutition, and speech. The neck also contains numerous lymph nodes and a high density of lymphatic vessels. A precise knowledge of the anatomical relations in the neck is an essential prerequisite for diagnosis and surgical treatment.

Topographic Anatomy

The neck is bounded above by the inferior border of the mandible, the tip of the mastoid process, and the external occipital protuberance. The lateral contours of the neck are defined by the palpable sternocleidomastoid muscles and the borders of the trapezius muscles. Palpable medial structures are the hyoid bone, thyroid cartilage, cricoid cartilage and, when enlarged, the thyroid gland (Fig. 16.1).

The neck muscles, the cervical viscera (larynx, trachea, pharynx, esophagus, etc.), and the neurovascular structures in the neck are encased by sheets of connective tissue, the fascial planes. These fasciae are comprised of superficial, middle, and deep layers (see 16.1).

Blood Supply

**Arterial blood supply:** The common carotid artery divides into its two main branches, the internal and external carotid arteries, at the level of the superior border of the thyroid cartilage (roughly the level of the C4 vertebral body). The common carotid artery arises from the aortic arch on the left side and from the brachiocephalic trunk on the right side. The lower part of the neck receives most of its blood supply from the thyrocervical trunk, which arises from the subclavian artery. Branches from the external carotid artery supply the neck and face; the internal carotid artery gives off no branches in the neck.

**Venous drainage** from the head and neck is received by the superficial cutaneous veins that open directly into the subclavian vein (external jugular vein and anterior jugular vein) and particularly by the internal jugular vein, which has a much larger lumen. The vertebral veins and the venous plexuses in the cervical spinal canal normally handle approximately 30% of the cerebral venous return.

Lymphatic Drainage

Tributary areas in tissues are drained by lymphatic channels that lead to regional lymph nodes or groups of lymph nodes. The lymph nodes in the neck are integrated as biological “filtering stations” into this network of lymphatic capillaries and vessels. Of the approximately 1000 lymph nodes in the human body, some 300 are located in the head and neck region. The most important of these are found between the middle and deep layers of cervical fascia. Based on the arrangement of the lymphatic channels, the lymph nodes at the junction of the facial and internal jugular veins (see 16.3) receive drainage from almost all parts of the head and neck region and are a site of predilection for lymphogenous metastases in a large percentage of malignant head and neck tumors (Fig. 16.2).
16.1 Facial compartments in the neck

**Fascial planes**
The fascial planes subdivide the neck into compartments that can easily shift in relation to one another. The superficial cervical fascia (synonym: superior layer of cervical fascia, Fig. yellow) underlies the platysma and subcutaneous fat, invests the entire neck, and encases the sternocleidomastoid and trapezius muscles. This fascial layer is attached to the hyoid bone and stretches superiorly to the mandibular border and inferiorly to the manubrium sterni and clavicle. It is fused at the midline to the middle cervical fascia (synonym: pretracheal layer of cervical fascia; Fig. green). This fascial layer stretches between the hyoid bone, the posterior surface of the manubrium sterni, and the clavicle and extends laterally to the omohyoid muscle and scalpa. It encases the infrahyoid muscles and forms a general anterior boundary for the cervical viscera. The deep cervical fascia (synonym: prevertebral layer of cervical fascia; Fig. blue) arises from the spinous processes of the cervical vertebrae and forms a rigid tube around the deep neck muscles that is adherent posteriorly to the superficial cervical fascia enasring the trapezius muscle. This prevertebral layer is part of a fascial system that extends continuously from the skull base to the lower end of the spinal column (prevertebral gravitation abscesses).

**Neurovascular sheath**
The carotid artery, internal jugular vein, and vagus nerve are invested by their own connective-tissue sheath (Fig. a, red) that is attached to the middle cervical fascia at the tendon between the inferior and superior bellies of the omohyoid muscle, so that contractions of this muscle place tension on the "neurovascular sheath" and especially on the internal jugular vein.

**Neck spaces**
While the space between the superficial and middle layers of the cervical fascia is closed inferiorly (common insertion on the manubrium sterni and clavicle), the visceral compartment of the neck has an open connection with the mediastinum between the middle and deep fascial layers. This allows abscesses in the cervical soft tissues and other disease processes to spread freely into the chest.

16.2 Carotid sinus and carotid body

The bifurcation of the common carotid artery marks the location of the carotid sinus. This dilatation of the carotid wall contains mechanoreceptors that are sensitive to blood pressure increases based on changes in vessel wall tension, and they evoke an antihypertensive response in the reticular formation via the glossopharyngeal nerve. Also located at the carotid bifurcation is the carotid body, composed of nonchromaffin paraganglia and measuring a few millimeters in size. It contains chemoreceptors that can modulate respiration in the reticular formation, also via the glossopharyngeal nerve, in response to changes in the arterial $PO_2$, $PCO_2$, and pH.

16.3 Jugular vein and its junctions with the facial and subclavian veins

The internal jugular vein arises at the confluence of the sigmoid sinus and inferior petrosal sinus and expands at the jugular foramen to form the jugular bulb. Two venous junctions are important because they receive drainage from major groups of lymph nodes: the jugulofacial venous junction, formed by the termination of the facial vein at the internal jugular vein, and the larger jugulo-subclavian venous junction, formed by the union of the internal jugular and subclavian veins at the brachiocephalic vein behind the sternoclavicular joint.
On the left side of the neck, lymph drains into the junction of the left subclavian and internal jugular veins at the termination of the left thoracic duct. Lymph on the right side drains into the junction of the right subclavian and internal jugular veins at the termination of the right thoracic duct. Included among the portals of entry to this system are the lymphatic organs of the nasopharynx and oropharynx (Waldeyer’s ring).

Normal cervical lymph nodes are neither visible nor palpable.

With a normal configuration of the neck, lymph nodes that enlarge to a diameter of 1 cm or more can be palpated.

**Innervation**

The external neck receives a nonmetameric sensory supply from the **cervical plexus** (C1–C4), which emerges at the posterior border of the sternocleidomastoid muscle (Erb’s point). **Motor components** of the cervical plexus (inferior root) unite with motor fibers of the cervical plexus, some of which have hitchhiked with the hypoglossal nerve (superior root), to form the ansa cervicalis and supply...
the infrahyoid muscles and geniohyoid muscle. The hypoglossal nerve exits the skull through the hypoglossal canal and crosses over the branches of the external carotid artery.

The glossopharyngeal nerve, vagus nerve, and accessory nerve pass jointly through the jugular foramen to enter the neck (16.4, Figs. 16.3–16.6).

The cervical part of the sympathetic trunk lies behind the neurovascular sheath between the layers of the deep cervical fascia on the prevertebral muscles. The location of the three associated ganglia is described in Table 16.1.

Table 16.1 Ganglia of the sympathetic trunk

<table>
<thead>
<tr>
<th>Ganglion</th>
<th>Location</th>
<th>Segments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Superior cervical ganglion</td>
<td>Behind the internal carotid artery</td>
<td>C1–4</td>
</tr>
<tr>
<td>Middle cervical ganglion (inconstant)</td>
<td>At the bend of the inferior thyroid artery</td>
<td>C5–6</td>
</tr>
<tr>
<td>Inferior cervical ganglion</td>
<td>Between the transverse process of C7 and the head of the first rib</td>
<td>C7–8; usually fused with the cervicothoracic ganglion (synonym: stellate ganglion)</td>
</tr>
</tbody>
</table>
Fig. 16.4 Course of the vagus nerve (cranial nerve X)

Source: adapted from Whitaker, see p. 410.
Fig. 16.5  Course of the accessory nerve (cranial nerve XI)

Source: adapted from Whitaker, see p. 410.

Fig. 16.6  Course of the hypoglossal nerve (cranial nerve XII)

Source: adapted from Whitaker, see p. 410.
16.2 Symptomatology and Examination of the External Neck

Many lesions in the neck are palpable owing to the topographic anatomy of that region. Palpation of the neck is an essential part of every physical examination, therefore. Modern imaging procedures can significantly narrow the differential diagnosis and direct the planning of treatment. Diagnostic surgical exploration of the neck is still necessary in some cases, however.

Cardinal Symptoms

Patients with diseases of the external neck very often complain of shape changes in the neck due to circumscribed or diffuse swellings or masses. Other common symptoms are pain, which may change with head movements, and limitation of neck motion. These symptoms require differentiation from dysphagic complaints, which are projected into the swallowing tract and are usually caused by disease in that region.

Methods of Examination

History

The history should focus on the duration, presentation, and time course of the symptoms (acute or chronic); the degree of pain or tenderness; constant, increasing, or fluctuant swellings; diseases involving the upper aerodigestive tract; other enlarged lymph nodes elsewhere in the body; and any prior history of similar complaints. The patient should also be questioned about possible contact with animals, travel abroad, and eating habits.

Inspection

The history is followed by visual inspection, which focuses on the contour-defining structures of the neck and any external changes in the overlying skin such as vascular markings, venous stasis, radiodermatitis, and skin tumors. The examiner should also look for fistulous openings that might indicate a branchial fistula, thyroglossal fistula, or actinomycosis and for sites of swelling and induration (lymph nodes, abscesses). Attention is also given to the position and mobility of the head and neck (e.g., a guarded head position due to an abscess or torticollis).

Palpation

The cervical soft tissues are palpated bimanually from the front or preferably from behind in the seated patient, alternating from right to left and comparing the sides. The patient’s head should be tilted slightly forward to relax the soft tissues of the neck. It is best to palpate the various lymph-node groups by following a routine sequence, as illustrated in Fig. 16.7.

The rest of the neck should be palpated as well, giving attention to the number of any palpable masses, their size (in centimeters), tenderness, mobility relative to the skin and underlying tissues, consistency, and especially their relationship to surrounding neck structures.

Both sides of the neck should never be deeply palpated at the same time, especially in older patients, due to the risk of evoking a carotid sinus reflex with vasovagal syncope.

Pulsating masses in the neck (paraganglia, vascular aneurysms) should be evaluated by auscultation as well as palpation.

The examiner should also test the mobility of the cervical spine in all planes and the mobility of the shoulder girdle.

Imaging Studies

Two-dimensional B-mode ultrasonography is considered the standard method for investigating soft-tissue lesions in the neck. The ultrasound examination can supply information useful in establishing the identity of specific lesions based on somatologic criteria.

Ultrasound cannot positively distinguish between benign and malignant masses, however.
Ultrasound scanning of the cervical soft tissues is particularly important for the follow-up of malignant upper aerodigestive lesions to detect a possible recurrence. For physical reasons, however (the strong reflection of sound waves at air and bone interfaces), ultrasound has only limited value in evaluating lesions that are adjacent to bone, the pharynx, or the larynx. Ultrasound, especially Doppler ultrasound, is excellent for evaluating the extracranial cerebral arteries (steno-ses, tumor invasion) and (very) vascular tumors (paraganglioma, hemangioma).

Axial computed tomography (CT) can provide a detailed view of inflammatory, space-occupying, infiltrating, or destructive cervical mass lesions. Scans can be obtained before and after contrast administration if required.

Iodinated contrast media should not be used in patients with a suspected thyroid disease, as they would interfere with subsequent tests.

Within limits, CT can provide a “radiologic tissue diagnosis.” CT is particularly indicated in cases with suspected involvement of cartilaginous or bony structures (larynx, cervical spine, mandible), which are demonstrated less clearly by MRI.

Magnetic resonance imaging (MRI) provides better soft-tissue discrimination than computed tomography and can aid in the differentiation of neoplastic, cicatricial, and inflammatory lesions, aided if necessary by paramagnetic contrast administration. Coronal and sagittal images are excellent for defining the location and relations of soft-tissue lesions.

Plain radiographs in the anteroposterior or lateral projection are rarely obtained in routine evaluations of the cervical soft tissues and have been largely superseded by the imaging modalities described above. Plain radiographs can detect prevertebral soft-tissue swellings, calcifications (e.g., tuberculous lymph nodes), emphysema following an upper airway injury, and radiopaque foreign bodies. They continue to have an important role in evaluating the cervical spine and, in some cases, detecting laryngeal fractures.

Positron emission tomography (PET), which can demonstrate the increased uptake of injected fluorodeoxyglucose (FDG) in metabolically active tissue (tumor, inflammation), is the latest imaging modality and is applied mainly in tumor diagnosis.

Cervical lymphography and lymphoscintigraphy no longer have a significant clinical role in evaluating the cervical lymphatic system.

Cytologic and Histologic Examination

Equivocal and suspicious neck masses should always be sampled so that a histologic tissue diagnosis can be made. With cystic masses, percutaneous needle aspiration can furnish material for cytologic and/or microbiologic examination. This diagnostic puncture is separate from a possible subsequent therapeutic incision, which may done to treat a suppurrative process.

Caution: pulsating neck masses (paraganglioma, vascular aneurysms should not be punctured!)

With solid masses, fine-needle aspiration biopsy can be performed (under ultrasound guidance if needed) to obtain material for cytologic analysis. This diagnostic procedure requires considerable experience in obtaining and evaluating the sample and is definitive only when positive due to the risk of false-negative findings. A negative result should be interpreted with caution.

In a core biopsy, the needle has a larger bore that provides a sample (“tissue core”) for histologic evaluation. As in fine-needle aspiration, however, only a positive result is definitive.

The most reliable method for establishing the identity of a long-standing neck mass unresponsive to conservative therapy is open biopsy, in which part of the tumor or preferably the entire “lump” is excised for histologic scrutiny and possible further analysis. A pre-scare biopsy (synonym: Daniel biopsy) involves the removal of lymph nodes at the jugulosubclavian venous junction, which represents the final collecting point for lymphatic drainage in the body.
16.3 Malformations of the Neck

Malformations of the neck include branchial and thyroglossal duct cysts and fistulas, vascular malformations such as hemangiomas and lymphangiomas, musculoskeletal anomalies, and dysontogenetic tumors.

Definitions: A cyst is an epithelium-lined cavity that does not have an internal or external opening. A sinus is an epithelium-lined cavity that opens either internally (common but incorrect synonym: incomplete internal fistula) or externally (synonym: incomplete external fistula). A fistula is an epithelium-lined tract that has both an internal and external opening (Fig. 16.8).

Branchial Cleft Cysts and Fistulas

Etiology and pathogenesis: The basic embryology of branchial cleft cysts and fistulas is reviewed in 16.5. The classic theory holds that branchial cleft cysts and fistulas result from a persistence or incomplete regression of the cervical sinus (cervical sinus theory). Sinuses that open internally (incomplete internal fistulas) are viewed as a remnant of the second branchial pouch, and those that open externally (incomplete external fistulas) are a remnant of the second branchial cleft, each having a connection with the cervical sinus. (Complete) fistulas can develop only if both the endoderm of the branchial pouch and the ectoderm of the branchial cleft communicate with the cervical sinus, rupturing the pharyngeal membrane.

Topography: Cervical fistulas and cysts occur predominantly at the border of the second branchial arch, because it is the largest of the arches and persists for the longest time during embryonic development. The cervical fistulas and cysts that develop there are very closely related to the carotid bifurcation. An internal sinus in this area opens into the supraventricular fossa. The sac of a branchial cleft cyst is always located lateral to the internal jugular vein and caudal to the posterior belly of the digastric muscle (Fig. 16.9).

Malformations of the third and fourth branchial arches (see 16.6 for location) are considerably less common.

Epidemiology: Branchial cleft cysts and sinuses are generally manifested between 15 and 25 years of age, whereas branchial fistulas are usually noted immediately after birth. Both sexes are affected equally. Branchial cleft cysts are approximately four times more prevalent than branchial fistulas, hence they are relatively common and one of the main causes of neck swelling in children. It is extremely rare for branchial cleft cysts to undergo malignant transformation.

Symptoms and diagnosis: Branchial cleft cysts generally have a short history. Patients notice a painless, tense swelling in the carotid triangle between the hyoid bone and sternoclavicular muscle. It is not unusual for an acute inflammation, which can be difficult to distinguish from a cervical abscess, to direct attention to a branchial cleft cyst.
16.5 Embryology and malformations of the branchial apparatus

Embryology

The branchial apparatus is composed of six mesodermal branchial arches (pharyngeal arches; the fifth and sixth arches are rudimentary in humans), five entodermal branchial pouches (pharyngeal pouches), and four ectodermal branchial clefts (branchial grooves). In mammals, entodermal and ectodermal duplications of epithelium that are devoid of mesenchyme, called the pharyngeal membranes, separate the internal branchial pouches from the external branchial clefts at the borders of the branchial arches. In fish, these membranes rupture to form the gills.) Each branchial arch has an artery, cartilage bar, muscular element, and nerve.

Theories of pathogenesis

The two main theories on the pathogenesis of branchial anomalies are presented below:

Classic theory: According to the classic theory, the second branchial arch enlarges disproportionately during the fifth week of development and overlaps the third and fourth arches, forming an ectodermal cavity called the cervical sinus (of His), which is surrounded by the operculum (Fig. a). Afterward the cervical sinus becomes progressively smaller, finally closing to form an ectodermal inclusion in the mesoderm, the cervical vesicle, which disappears with further development (Fig. b). But this classic theory, which regards branchial cleft cysts and fistulas as a failure of regression of the cervical sinus and vesicle, has certain deficiencies that have been addressed in the current model proposed by Otto.

Otto theory: This theory disputes the notion that the cervical sinus closes to form the cervical vesicle, followed by complete resorption of the enclosed epithelium (viewing this instead as a sectioning or preparation artifact), claiming that all of the ectodermal epithelium in the cervical sinus migrates back to the surface. According to this theory, the pharyngeal membrane stage (separating the branchial clefts and pouches; Fig. c) is followed developmentally by a stage of internal and external connecting lamellae, with the external and internal lamellae eventually migrating back respectively to the external and internal body surface (Fig. d). Otto theorizes that a local interepithelial adhesion (LIAD), represented by desmosomes and interdigitations, forms at the site of a pharyngeal membrane or connecting lamella, where a gap exists in the mesenchyme (Fig. d). If mesodermal fusion fails to take place at this LIAD at the proper time, epithelial retention and proliferation can occur as a redundancy anomaly.

Pathogenesis of branchial cleft cysts and fistulas: When this LIAD theory, which is valid for embryonic development in general, is applied to branchial cleft cysts and fistulas, it means that a persistent LIAD is formed in the area of the second, third, or fourth pharyngeal membrane. Further growth leads to a displacement of the ectodermal or entodermal epithelial elements in which the LIAD becomes elongated and is “dragged along” to form a fistulous tract (Fig. e).

If the LIAD persists in its entirety, a (complete) fistula develops (Fig. f). If the fistulous tract is torn from one of the two surface aggregates of epithelial cells, an internal or external sinus is formed. And if the fistulous tract is separated from both epithelia, a branchial cleft cyst will form.

The pathogenesis of thyroglossal duct cysts and thyroid anomalies is reviewed in 16.7.

(Fig. c-f adapted from Otto, see p. 410)
Fig. Branchial cleft cyst

16.6 Malformations of the third and fourth branchial arches

Malformations of the third branchial arch open internally in the lateral wall of the piriform recess anterior to the plica of the superior laryngeal nerve at the level of the cricoid cartilage. The very rare malformations of the fourth branchial arch open at the tip of the piriform recess, the fistulous tract imitating the course of the recurrent nerves.

Besides palpation, B-mode ultrasound is the diagnostic method of choice, demonstrating branchial cleft cysts as homogeneous, echo-free masses with smooth margins located at a typical site. Acutely infected cysts may contain a viscous, purulent material that appears echogenic.

Sinuses (“incomplete fistulas”) that open internally are more common than complete fistulas and can produce the same symptoms as branchial cleft cysts. Branchial fistulas always open externally at the anterior border of the sternocleidomastoid muscle (usually one to two fingerwidths above the sternocleidomastoid joint) and may be marked by a clear, amber-colored discharge. Acute infections with the typical signs of pain, redness, and purulent drainage are common. Accessory cartilage may occur at the fistula opening. The tract itself can be defined by contrast radiography. An effort should be made to locate the pharyngeal opening of the fistula, which may be in the suprathyroid fossa, faucial pillar, lateral pharyngeal wall, or piriform recess.

Differential diagnosis: The differential diagnosis of cervical cysts includes all other cervical masses. With fistulas, the possible presence of lymph-node tuberculosis or actinomycosis should be considered.

Treatment: Surgical treatment is generally indicated.

To prevent a recurrence, no residual epithelium should be left behind.

Fistulas are identified intraoperatively with methylene blue and excised. The close relationship of the fistulous tracts to the cervical vessels and nerves, as well as inflammatory adhesions, often make the surgery difficult.

Even in the excision of presumed cysts, a careful search should be made for a tract to the suprathyroid fossa. When present, the tract should be traced and excised to prevent a recurrence. For this reason, the operation should always include a tonsillectomy. Peri-tonsillar abscesses following tonsillectomy generally result from the presence of entodermal duct remnants (the “duct of His”) in the suprathyroid fossa. Acutely inflamed cysts or fistulas should first be treated conservatively with antibiotics to facilitate subsequent surgery.

Puncture or incision of branchial cleft cysts is indicated only in exceptional cases with severe pain.

Thyroglossal Duct Cysts and Fistulas

Etiology and pathogenesis: The pathogenesis of thyroglossal duct cysts and fistulas is closely linked to the embryonic development of the thyroid gland (16.7).

Epidemiology: Approximately 75% of thyroglossal duct cysts are manifested before 5 years of age, and most are diagnosed before 12 months. The malignant transformation of thyroglossal duct cysts is rare.

Symptoms: Parents often notice a tense, firm swelling in the midline of the neck between the chin and thyroid gland (or very rarely at the suprasternal level). Not infrequently, an inflammatory process may first draw attention to a thyroglossal duct cyst. The swelling may be intermittent or continuous. Swallowing difficulties are uncommon.
16.7 Embryology and malformations of the thyroid gland

Embryology (Fig. a)

On day 24 of embryonic development, a median epithelial thickening appears in the floor of the ectodermal pharyngeal gut, dorsal to the future tuberculum impar. It develops into the thyroglossal duct, on which the thyroid primordium descends into the neck. The thyroid gland reaches its pretracheal position by the end of the seventh week, and the thyroglossal duct is obliterated or resorbed. The foramen cecum in the midline of the tongue and the pyramidal lobe of the thyroid gland are viewed as the remnants (ends) of the thyroglossal duct. The hyoid bone develops later and has a variable relationship to the thyroglossal duct.

Pathogenesis

Thyroglossal duct cysts and malformations of the thyroid gland

Classic theory: The classic theory attributes the formation of thyroglossal duct cysts to incomplete obliteration or resorption of the thyroglossal duct.

Otto theory: More recent studies call into question the descent of the thyroid gland, instead suggesting that the thyroid primordium develops near the heart with the subsequent formation of a connecting tract between the thyroid gland and the epithelium of the oral floor (thyroglossal duct); this results in an ascent of the head, taking the thyroid gland with it. According to this theory, the thyroglossal duct contains two types of epithelium: oral floor epithelium superiorly (pars epithelialis) and thyroid epithelium inferiorly (pars thyroidea). The differentiating boundary between the two epithelial types (local interepithelial adhesion, LIAD; see also 16.5) ruptures by the middle of the sixth week (Fig. a).

All malformations of the thyroid gland can be explained on the basis of this theory:

If the thyroglossal duct does not rupture at the level of the differentiating boundary but at a lower level (in the pars thyroidea), thyroid tissue will remain at the cranial end of the thyroglossal duct during the subsequent ascent of the head. Ectopic thyroid tissue or a lingual goiter may develop as a result of this mechanism (Fig. b). If the thyroglossal duct ruptures at a higher level (in the pars epithelialis), non-thyroid epithelial tissue will persist between the base of the tongue and thyroid gland and may form a nidus for the development of an epithelial cyst (= thyroglossal duct cyst). These epithelial cysts may also contain thyroid tissue (Fig. c). The prevalence of thyroid anomalies is approximately 7%, the most frequent anomaly being a persistent pyramidal lobe.

Thyroglossal duct fistulas, which are more aptly called sinuses, are not primary lesions; they are secondary to the (inflammatory) external perforation of a thyroglossal duct cyst or to iatrogenic measures (e.g., needle aspiration).

Ultrasound may support the presumptive diagnosis by showing a well-circumscribed, elliptical, hypoechoic or echo-free mass with distal acoustic enhancement (Fig. 16.11 b).

It should always be confirmed that a normally developed thyroid gland is present.

Thyroglossal fistulas are usually recognized by their external opening (typically located at the level of the thyroid notch) and the associated discharge. Inflammations with retention of secretions and abscess formation may occur.

Diagnosis: The diagnosis is suggested by inspection and palpation (Figs. 16.11 a, 16.12). A thyroglossal duct cyst will invariably move when the patient swallows.
The swelling in the neck appears sonographically (b, transverse B-mode image) as a sharply circumscribed mass with homogeneous contents and distal acoustic enhancement.

Surgical specimen of a thyroglossal duct cyst with attached midportion of hyoid bone.

disposes to a recurrence (recurrent cyst or fistula). For the same reason, the resection should always include the body of the hyoid bone since epithelial remnants are frequently present in or on the bone. Failure to resect the midportion of the hyoid bone is associated with up to a 50% recurrence rate, which otherwise is less than 5%. Resecting the body of the hyoid bone has no functional sequelae. It will not impair swallowing because the functionally significant hyoid muscles are attached to the cornua of the hyoid bone.

Thyroglossal fistulas are outlined with an elliptical excision and removed in toto. This surgery is aided by intraoperative staining of the fistulous tract with blue dye.

**Vascular Malformations**

**Lymphangioma**

Synonym: cystic hygroma

Lymphangiomas develop from sequestered portions of the primordial lymphatics that begin to sprout in the sixth week of embryonic development. These sequestra do not establish connections with the venous system, and they degenerate to form cystic cavities. They tend to grow by expansion and infiltration. Ninety percent of lymphangioma are manifested during the first two years of life. Many are already present at birth and may cause obstruction of labor. Lymphangiomas are typically located in the lateral part of the neck (Fig. 16.13).

The tumor is palpable as a soft, doughy mass. Generally, the true extent of the mass is underestimated by palpation due to its infiltrative growth. B-mode ultrasound provides a more accurate assessment. Local expansile growth can cause symptoms such as dyspnea and stridor.
**Treatment:** Unlike hemangiomas, lymphangiomas very rarely undergo spontaneous regression. Therefore a wait-and-see approach is generally not justified and surgical treatment is required. The local injection of a crystalline corticosteroid solution may be tried as an alternative to surgery.

**Hemangioma**

Hemangiomas are growing vascular lesions (ectasias) that belong to the group of hamartomas (= dysontogenic growth of normally formed tissue).

**Epidemiology:** Hemangiomas have a reported incidence of 10% during the first year of life. Premature infants are predominantly affected. Superficial hemangiomas are more common in females.

**Symptoms:** Hemangiomas can occur anywhere on the body, appearing clinically as a soft, reddish-purple swelling (Fig. 16.14). Large hemangiomas may be hemodynamically significant.

**Course:** Unlike lymphangiomas, hemangiomas regress spontaneously in up to 80% of cases. The natural history proceeds in phases—a two-part proliferative phase is followed at about 1 year of age by a stable plateau in which the lesion ceases to grow. The involutional phase then occurs during the first or second year of life or may be delayed until puberty.

**Diagnosis:** Besides B-mode ultrasound, which can define the overall extent of the lesion, color duplex ultrasound has proved to be a particularly useful diagnostic tool.

The patient should always be screened for additional hemangiomas of the internal organs (lung, liver, etc.).

**Treatment:** The high rate of spontaneous involution justifies a wait-and-see approach, unless the hemangioma is growing rapidly or is causing symptoms due to its location or extent. Treatment options depend on the individual case and consist of systemic corticosteroid therapy (anti-inflammatory and angiostatic effect), local sclerotherapy or embolization, magnesium spiking, laser treatment—dye laser, argon laser, neodymium: yttrium—aluminum garnet (Nd:YAG)—and surgical excision. Radiotherapy is used only in highly selected cases due to the risk of late sequelae.

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**Fig. Lymphangioma in an infant**

**Fig. Hemangioma**

**16.8 Special types of hemangioma**

The Kasabach–Merritt sequence (synonym: thrombopenia–hemangioma syndrome) is characterized by the presence of typically large (“giant”) hemangiomas in which thrombotic processes lead to disseminated intravascular coagulation with consumption coagulopathy. The syndrome affects small infants almost exclusively. Another rare entity is blue rubber bleb nevus syndrome, which has an autosomal-dominant mode of inheritance. Multiple, deep blue hemangiomas with a rubbery consistency form on the integument and persist without involution. Gastrointestinal manifestations can lead to recurrent episodes of bleeding.
Paraganglioma

Synonyms: chemodectoma, glomus tumor

Nonchromaffin (16.9) paraganglia, which function as chemoreceptors, are located at various sites in the neck: the carotid bifurcation (carotid paraganglion), the vagus nerve (vagal paraganglion), the internal jugular vein (jugular paraganglion), and the larynx (laryngeal paraganglion). Neoplasms that are derived from the chemoreceptor tissue in paraganglia are called paragangliomas. “Glomus tumor” is an archaic term that should no longer be used.

Epidemiology: Paragangliomas are mostly solitary and benign. Malignant transformation occurs in up to 10–20% of cases. Paragangliomas are believed to occur more frequently in persons living at higher altitudes.

16.9 Meaning of the term “nonchromaffin”

The term “nonchromaffin” stems from the historical observation that the cells of the paraganglia do not take up the chromaffin stain that is used in the detection of catecholamines.

Carotid Paraganglioma

Symptoms: This tumor presents as a painless, sometimes pulsatile mass in the neck. It may be accompanied by a dry cough (due to vagus nerve irritation), hoarseness (vagus or recurrent nerve lesion), and Horner syndrome (sympathetic trunk irritation). A painful sensation is sometimes present.

Diagnosis: Palpation reveals a soft or tense, sometimes pulsatile mass at the level of the carotid bifurcation. In typical cases the mass is mobile in the lateromedial direction but not cranio-caudally. A bruit is usually heard at auscultation. When imaged by B-mode ultrasound, the paraganglioma appears as a hypoechoic mass splaying the carotid bifurcation.

Paragangliomas cannot always be distinguished from branchial cleft cysts or enlarged lymph nodes. This differentiation is aided by color duplex sonography, which can clearly demonstrate the rich vascularity of paragangliomas (Fig. 16.15). Magnetic resonance imaging and somatostatin receptor scintigraphy can generally furnish a definitive diagnosis. Any surgical measures should be preceded by angiography.

Needle aspiration is contraindicated due to the rich vascularity of the tumor.

Treatment: The treatment of choice for paragangliomas of the carotid bifurcation is surgical removal. Most tumors infiltrate the adventitia of the carotid artery, making removal difficult. Generally, the surgery requires autologous blood donation and/or the intra-operative use of a cell saver.

Torticollis

Synonym: wryneck

Torticollis may be congenital or acquired. The possible causes are listed in Table 16.2. Diagnosis and treatment are usually handled by a neonatologist or orthopedist.

Dysontogenetic Tumors

Embryonal tumors are derived from immature, primitive tissue, and malignant forms are common due to the pluripotency of the cells. A particularly common representative in the neck is embryonal rhabdomyosarcoma, which is derived from the primordia of striated skeletal muscle. Treatment follows established protocols for soft-tissue sarcomas in pediatric oncology. Surgical treatment should be provided in collaboration with a pediatric oncologic center.

Teratomas are characterized by the presence of cellular elements from all three germ layers (ectoderm, mesoderm, endoderm) and show varying degrees of differentiation from immature embryonic tissue to fully formed elements (e.g., cartilage, bone, teeth). Cervical teratomas account for approximately 5% of congenital teratomas and are usually manifested before 12 months of age. Differential diagnosis is aided by plain radiographs of the neck, which often show inclusions of calcific density. Given their potential for malignant transformation, teratomas should be surgically removed. This particularly applies to teratomas in adults, which show a strong propensity for malignant degeneration (“teratocarcinoma”).

Dermoids, unlike teratomas, are composed mainly of ectodermal elements with a complete absence of endodermal elements. Dermoids are formed by the encapsulation of ectodermal epithelial elements during development and thus represent dystopic epithelium within the mesenchyme. Given the ectodermal origin of the tissue, histologic examination often reveals dermal appendages (hair follicles, sebaceous glands, sweat glands) and desquamated amorphous material.

Treatment: consists of surgical removal.

Hammartomas result from the circumscribed, neoplastic overgrowth of tissue that is normally differentiated in most cases and is normally present in that part of the body. The most common example of a hamartoma in the neck is hemangioma (see above).
16.10 Vagal paraganglioma

The paraganglia of the vagus nerve are usually directly adjacent to the ganglia of this cranial nerve that lie below the jugular foramen. Tumors of the vagal paraganglion are relatively rare (approximately 3% of paragangliomas). The **cardinal symptom** is vagus nerve palsy with hoarseness and aspiration. Extensive tumors may cause a jugular foramen syndrome (synonym: Avellis syndrome). Intracranial extension (dumbbell tumor) is particularly common with paragangliomas close to the superior vagal ganglion. As with carotid paragangliomas, the diagnosis is established by B-mode ultrasound, color Doppler ultrasound, computed tomography, magnetic resonance imaging, and in some cases angiography. The **differential diagnosis** consists mainly of vagus neurinoma. **Surgical removal** is the treatment of choice and generally requires sacrificing the affected vagus nerve. Extensive tumors will require a combined transcervical-transmastoid approach.

**Table 16.2 Causes of torticollis**

<table>
<thead>
<tr>
<th>Congenital</th>
<th>Acquired</th>
</tr>
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<tbody>
<tr>
<td><strong>Muscular</strong></td>
<td>Radical neck dissection with removal of the sternocleidomastoid muscle</td>
</tr>
<tr>
<td>- Common: obstetric trauma with tearing and hematoma formation in the sternocleidomastoid muscle</td>
<td>- Trauma to the cervical spine</td>
</tr>
<tr>
<td>- Less common: congenital fibrous transformation of muscle tissue</td>
<td>- Functional disorders of the ocular muscles (ocular torticollis)</td>
</tr>
<tr>
<td>- Muscular dystrophies</td>
<td>- Analgesic guarding due to cervical abscess or pronounced lymphadenitis</td>
</tr>
<tr>
<td><strong>Bony</strong></td>
<td>- Klippel–Feil syndrome with synostoses in the cervical spine</td>
</tr>
<tr>
<td>- Goldenhar syndrome with fusion and/or absence of cervical vertebrae</td>
<td>- Accessory nerve palsy</td>
</tr>
<tr>
<td></td>
<td>- Atlantoaxial torticollis following inflammatory disease, surgery, or radiotherapy to the nasopharynx (synonym: Grisel syndrome)</td>
</tr>
</tbody>
</table>

**Fig. Carotid paraganglioma**

a Doppler ultrasound scan of an extensive paraganglioma that has grown into the internal jugular vein. The color flow signals clearly demonstrate the vascularity of the tumor.

b Paraganglioma in the left side of the neck above the carotid bifurcation (arrows). The tumor does not involve the internal jugular vein.
16.4 Inflammations of the Neck

For differential diagnostic and therapeutic reasons, it is necessary to distinguish inflammatory reactions of the cervical lymph nodes (lymphadenitis) from "deep" inflammations of the cervical soft tissues, which may be circumscribed (abscess) or diffuse (cellulitis). This topic does not include cutaneous inflammations.

Inflammations of the Cervical Lymph Nodes

The cervical lymph nodes are affected by (infectious) inflammations with remarkable frequency. One reason for this is that the topographic anatomy of the neck makes it easy to detect even small masses; another is that the upper aerodigestive tract is a frequent portal of entry for infectious micro-organisms.

The cardinal symptom is almost always a palpable neck mass (Fig. 16.16). "Healthy" lymph nodes are neither palpable nor detectable by ultrasound. Constant pain or tenderness are typical signs of a (usually acute) inflammation but may be absent. Additional symptoms and findings such as neck pain, dental pain, otalgia, fever, nausea, swollen salivary glands, skin changes, etc. can help to narrow the differential diagnosis.

Possible methods of classification are shown in Table 16.3.

Fig. Enlarged cervical lymph nodes

Important considerations in differential diagnosis are the rate of enlargement, consistency, and mobility of the mass and the patient's general state of health.

Acute Cervical Lymphadenitis

Etiology and pathogenesis: Acute lymphadenitis in the neck is usually a reactive lymphadenopathy that develops in response to an infection of the upper respiratory tract (viral or bacterial rhinitis, sinusitis, pharyngitis, tonsillitis), the teeth and periodontal structures, the salivary glands, or the facial and neck skin (e.g., erysipelas, impetigo contagiosa). Thus, it presents few difficulties in terms of differential diagnosis.

The most frequent cause of acute cervical lymphadenitis in children is a streptococcal infection of the palatine tonsils. Other potential causes are the rubella virus and cytomegalovirus (CMV), and mycobacteria have increasingly been implicated in recent years. Infectious mononucleosis is an acute disease of the lymphatic system caused by the Epstein–Barr virus, a DNA virus that belongs to the Herpesviridae family.

Symptoms: Patients typically experience a systemic inflammatory reaction with fever and malaise. The cervical masses have a soft consistency and are usually painful. Multiple lymph nodes are typically affected and may be unilateral or bilateral, depending on the site of the underlying organic infection.

Infectious mononucleosis is characterized by tonsillitis and a usually pronounced (voluminous), painful cervical lymphadenitis (see 5.4, pp. 116–118).

In a primary infection with the human immunodeficiency virus (HIV), an incubation period of 1–3 weeks is followed by an "acute" stage marked by flu-like symptoms, an itchy skin rash, and generalized lymphadenitis.

16.11 Kawasaki syndrome

Kawasaki syndrome (synonym: mucocutaneous lymph node syndrome) is a diffuse vasculitis that occurs in children and presents with fever, mucocutaneous changes, and swollen cervical lymph nodes. A feared complication, occurring in one-fourth of cases, is cardiac involvement in the form of myocarditis, coronary arteritis, or both. A bacterial etiology is presumed, but it has not been possible to isolate a causative organism. Whenever the syndrome is suspected, a pediatric examination should be scheduled without delay.
Table 16.3  Classification of cervical lymphadenitis

<table>
<thead>
<tr>
<th>Classification</th>
<th>Example or features</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>by time course</strong></td>
<td></td>
</tr>
<tr>
<td>Acute</td>
<td>Duration of symptoms &lt;4 weeks</td>
</tr>
<tr>
<td>Chronic</td>
<td>Duration of symptoms &gt;4 weeks</td>
</tr>
<tr>
<td><strong>by etiology</strong></td>
<td></td>
</tr>
<tr>
<td>Infectious</td>
<td>Viral</td>
</tr>
<tr>
<td></td>
<td>Bacterial, nonspecific</td>
</tr>
<tr>
<td></td>
<td>Bacterial, specific</td>
</tr>
<tr>
<td></td>
<td>Fungal</td>
</tr>
<tr>
<td></td>
<td>Parasitic</td>
</tr>
<tr>
<td>Noninfectious lymphadenitis and lymphadenitis of unknown cause</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>by microbiologic or histologic features</strong></td>
<td></td>
</tr>
<tr>
<td>Serologic tests</td>
<td>Mononucleosis</td>
</tr>
<tr>
<td></td>
<td>Toxoplasmosis</td>
</tr>
<tr>
<td></td>
<td>Brucellosis</td>
</tr>
<tr>
<td></td>
<td>Tularemia</td>
</tr>
<tr>
<td></td>
<td>Syphilis</td>
</tr>
<tr>
<td></td>
<td>Cyto megalovirus</td>
</tr>
<tr>
<td></td>
<td>HIV infection</td>
</tr>
<tr>
<td></td>
<td>Cat-scratch disease</td>
</tr>
</tbody>
</table>

Culturing or identifying the organism

| | |
| Tuberculosis | Culture, DNA detection by PCR |
| Listeriosis | Culture |
| Cyto megalovirus | Isolation of the virus |

Histologic examination

| | |
| Nonspecific lymphadenitis | The causative agent cannot be determined from histologic findings |
| Specific lymphadenitis | The histologic findings provide clues to the causative agent |
| | |
| • Abscess-forming reticular lymphadenitis (pseudotuberculosis-type granulomas) | Yersinia pseudotuberculosis, Y. enterocolitica, Francisella tularensis, Afipia felis, Bartonella henselae |
| • Epitheliod cell lymphadenitis | Toxoplasmosis, sarcoidosis, tuberculosis, atypical mycobacteriosis, sarcoid-like reaction |
| • Necrotizing histiocytic lymphadenitis | Kikuchi lymphadenitis |
| • Sinus histiocytosis with hemophagocytosis | Rosai–Dorfman syndrome |
| • Angiofollicular lymph-node hyperplasia | Castleman lymphoma |

CMV: cytomegalovirus; EA: early antigen; EBNA: Epstein–Barr nuclear antigen; EBV: Epstein–Barr virus; ELISA: enzyme-linked immunosorbent assay; FTA-Abs: fluorescence Treponema antibody absorption test; HIV: human immunodeficiency virus; IgG: immunoglobulin G; IgM: immunoglobulin M; PCR: polymerase chain reaction; TPHA: Treponema pallidum hemagglutination; VCA: virus capsid antigen.

**Diagnosis:** When acute cervical lymphadenitis is present, it is first necessary to look for the precipitating cause, i.e., an organic infection (mirror examination by an otolaryngologist, dental consultation if needed). B-mode ultrasound examination of the neck is helpful in detecting or excluding lymphadenitis with abscess formation (central echo = liquefaction) and can better quantify the extent of lymphadenitis in relation to palpable findings (for follow-up and monitoring therapeutic response).
Differential diagnosis: An acutely inflamed thyroglossal duct cyst or branchial cleft cyst should be considered when acute lymphadenitis is present, as these lesions are usually associated with an accompanying (reactive) lymphadenitis.

With repeated episodes of cervical lymphadenitis, it is not uncommon for palpable lymph nodes to persist beyond the stage of the acute inflammation. This is attributable to inflammatory fibrosis and may lead to problems of differential diagnosis.

Treatment: Treatment is directed mainly toward the underlying organic infection. For example, penicillin is the agent of choice for a streptococcal infection. The parenteral administration of antibiotics should be considered in patients with systemic symptoms or suppurative lymphadenitis. The latter condition requires close-interval follow-up and may require an incision for abscess drainage.

Chronic Cervical Lymphadenitis

Etiology: Table 16.3 lists diseases that may be associated with lymphadenitis. Various medications may also cause lymphadenopathy, including antiepileptic drugs, tuberculosis, heparin, phenacetin, salicylates and other nonsteroidal anti-inflammatory drugs (NSAIDs), allopurinol, antibiotics, gold, and methyldopa.

Diagnosis and treatment:

Chronic cervical lymph-node enlargement (present for more than 4 weeks) often leads to diagnostic problems because it requires differentiation from a malignant disease such as malignant lymphoma and cervical lymph-node metastasis.

Besides growth kinetics, other factors to be considered in selecting a diagnostic procedure for cervical lymphadenopathy are the age of the patient, the presence of risk factors for malignancy (e.g., nicotine abuse), and any associated symptoms (e.g., "B symptoms" such as fever, night sweats, and undesired weight loss; see also 16.5, p. 333).

History: In planning the diagnostic algorithm, it is important to ask the patient about risk factors for the causes listed in Table 16.3:

- House pets: dogs, cats, rodents, etc.
- Occupational contact with animals or animal products: slaughterhouse worker, butcher, meat seller, cheese production and sales, farm workers, gardeners
- Eating habits: consumption of raw meat or sheep cheese
- Trips abroad
- Institutional settings: medical staff, nursing home workers, kindergarten teachers

Palpation (consistency, mobility relative to skin and underlying tissues) and B-mode ultrasound can provide initial clues in making a differential diagnosis.

Antibiotic trial: Since it is relatively common for cervical lymph-node enlargement to have a bacterial cause, a broad-spectrum antibiotic (e.g., doxycycline, macrolide) may be tried before proceeding with serologic tests or invasive diagnostic procedures in patients who have no obvious signs of a malignant underlying disease.

Make certain, however, that the patient understands the importance of follow-ups.

Serology: The selection of serologic tests should be patient-specific due to the long list of possible causative organisms (detailed history; see also Table 16.3).

Histology: As a general rule, a surgically removed lymph node will provide the most accurate diagnosis. A histologic work-up should be done in patients with a known immune deficiency (e.g., HIV infection) and in patients on immunosuppressive therapy due to the great many possible (and atypical) causative organisms and the increased incidence of malignant tumors in these subgroups.

Owing to the topographic anatomy of the neck, this type of surgery can usually be done under local anesthesia with relatively low risk. Because the histological architecture of the lymph node provides the best or only indicator of certain diseases such as malignant non-Hodgkin lymphoma, the most reliable biopsy technique is to excise a complete, representative lymph node rather than perform a partial excision or obtain a core- or fine-needle specimen.

Special forms and differential diagnoses:

Actinomycosis: This disease is caused by the gram-positive anaerobic bacterium Actinomyces israeli. The cervicofacial form of actinomycosis causes a "boardlike" infiltration of the subcutaneous tissue with firm nodules, fistulas, and ulcerations. Treatment consists of a prolonged course of antibiotic therapy, which may be supplemented if necessary by surgical measures.

Rosai-Dorfman syndrome (synonym: sinus histiocytosis with massive lymphadenopathy) is an essentially benign disease occurring in children and young adults and characterized by a (grotesque) enlargement of the cervical lymph nodes. The causative agent of the disease, which is unknown but may be infectious, incites an excessive phagocytosis of hematolymphatic cells, mostly lymphocytes. The disease usually runs a self-limiting course of several weeks, but aggressive forms may occur.

Castleman lymphoma (synonym: angiofollicular lymph-node hyperplasia) is a histologically defined entity that occurs in a localized form characteristic by a benign course and in a more aggressive multicentric
form; the latter is more likely to affect the cervical lymph nodes. The pathogenesis appears to be based on an abnormal regulation of interleukin-6 (IL-6) production, justifying a trial of corticosteroid therapy.

**Deep Neck Infections**

**Synonym:** parapharyngeal abscess

**Etiology:** Cervical abscesses can develop from various diseases:
- Tonsillar infections, peritonsillar abscess (see 5.4, pp. 113–116)
- Suppurative lymphadenitis
- Mastoiditis: Bezold’s mastoiditis (see Fig. 11.18, p. 248)
- Dentogenic infections: cellulitis of the oral floor (Ludwig’s angina)
- Lesions of the pharyngeal mucosa (retropharyngeal abscess, foreign bodies)

**Symptoms:** The characteristic features of cervical abscesses are high fever, tenderness to pressure and, in some cases, holding the neck and head in a guarded position. The symptoms of the causative underlying disease may also be present.

**Diagnosis:** The diagnosis is established by the typical clinical findings, markedly elevated inflammatory parameters—whole blood cell count (WBC), erythrocyte sedimentation rate (ESR), C-reactive protein (CRP)—and the detection of an abscess by ultrasound or CT.

**Treatment:** Treatment consists of antibiotic therapy specific for the underlying disease and surgical abscess drainage.

**Cervical Cellulitis**

Synonym: cervical phlegmon

Unlike circumscribed cervical abscesses, cervical cellulitis involves a diffuse inflammation of the cervical soft tissues.

Given the anatomical relationships of the cervical soft tissues and the communication of the parapharyngeal space with the mediastinum, cervical cellulitis is a life-threatening condition (see 16.1, p. 313).

**Etiology:** Cervical cellulitis may occur as a primary condition in patients with a predisposing underlying disease, or it may complicate an infection of the upper aerodigestive tract.

**Symptoms:** Most patients present with high fever and severe malaise. The cervical soft tissues are diffusely tender to pressure; even imaging studies cannot demonstrate discrete inflammatory foci. The cervical soft tissues appear markedly rarefied, and occasionally there is loss of normal delineation of anatomic structures.

**Treatment:** Treatment with broad-spectrum antibiotics (first- or second-generation cephalosporins, clindamycin, aminopenicillins + β-lactamase inhibitor, possibly combined with aminoglycosides) should be instituted without delay. All of the compartments in the neck should be broadly opened, and mediastinal drainage may be necessary (see Fig. 16.17).
16.5 Tumors of the Neck

Benign tumors of the neck are relatively rare. Except for the paragangliomas covered in 16.3 (p.326), the most common benign tumors of the neck are lipomas. A special type of cervical lipoma is Madelung disease. Very rare disorders are desmoid fibromatosis, fibrosing cervicitis, and rhabdomyoma, which are beyond our present scope. The presence of a malignant disease should always be considered in the differential diagnosis of neck masses. The diagnostic workup should be planned with this in mind: histologic confirmation should be obtained in patients with a suspicious history or clinical findings and for masses unresponsive to a trial of antibiotic therapy.

Benign Tumors

Lipoma

Lipomas present clinically as soft, circumscribed, painless masses. They have a characteristic ultrasound appearance: superficial (subcutaneous) lipomas are usually encapsulated, while deeper lipomas tend to show more aggressive, infiltrative growth. Surgical removal should be considered only for cosmetically objectionable masses.

Madelung Disease

Synonyms: benign symmetrical cervical lipomatosis, horse collar, Launois-Bensaude syndrome
This special form of lipoma occurs predominantly in middle-aged men who often have a history of excessive alcohol consumption (60–90% of patients). Hyperuricemia, diabetes mellitus, disorders of fat metabolism, and obstructive sleep apnea syndrome are common associated features.
Histologic examination shows a diffuse, unencapsulated, nonseptate proliferation of univacuolar adipocytes with tongue-like extensions into the surrounding tissue, which may involve a neoplasia of the brown fat. The changes are poorly demarcated from surrounding tissues.
Course: The most common "horse collar" form of the disease is characterized by the intermittent, symmetrical proliferation of fatty tissue about the neck (Fig. 16.18), nuchal area (“buffalo hump”), and/or the upper arms (“puffed sleeves”).
Treatment consists of surgical removal of the excess fatty deposits. Recurrences are not uncommon. Any metabolic diseases that are diagnosed should be referred for appropriate therapy.

The risk profile associated with Madelung disease (increased cancer risk!) requires surveillance by an otolaryngologist.

Neurinoma

Neurinomas arise from the Schwann cells of the fibrous nerve sheath and are also referred to as schwannomas. Most of these tumors are solitary, circumscribed, and encapsulated.
The most frequent sites of occurrence in the neck are the cervical plexus, brachial plexus, and vagus nerve. Neurinoma may also arise from the sympathetic trunk, glossopharyngeal nerve, accessory nerve, or hypoglossal nerve.

Symptoms: Occasionally the tumor causes local pain that is aggravated by palpation. There may also be symptoms relating to dysfunction of the affected nerve.

Diagnosis: A neurinoma should be suspected when imaging (ultrasound or CT/MRI) demonstrates a fusiform mass.

Treatment consists of surgical resection, which may be followed by a nerve reconstruction.

Differential diagnosis: Neurinomas require definitive histologic differentiation from neurofibromas. The latter are particularly common in Recklinghausen disease, which has an autosomal-dominant mode of inheritance. In contrast to neurinomas, it is not unusual for neurofibromas to undergo malignant transformation.

Fig. Madelung disease
Symmetrical lipomatosis of the neck in a 33-year-old man with a history of alcoholism. The affected areas feel soft on palpation (e.g., like a fatty abdominal wall).
Malignant Tumors of the Cervical Lymph Nodes

Malignant tumors in the neck are manifested predominantly in the cervical lymph nodes. Malignancies such as Hodgkin disease and non-Hodgkin lymphoma require differentiation from metastatic tumors with a different histologic origin. Soft-tissue tumors (sarcomas) that arise from connective tissue, muscle, or vessels are relatively rare.

Malignant Lymphomas

Certain subtypes of malignant lymphoma show a predilection for the head and neck region. Solitary or multiple lymph nodes or nodal stations may be involved, depending on the subtype and stage of the disease. Extracranial sites of involvement such as the spleen, liver, lung, skeleton, mucosa—mucosa-associated lymphatic tissue (MALT) lymphoma—Waldeyer’s ring (especially with B-cell lymphomas), and skin (especially with T-cell lymphomas) are not uncommon.

Diagnosis: The history and clinical findings can suggest the presence of a malignant lymphoma. Given the topographic anatomy of the neck, enlarged cervical lymph nodes become conspicuous at a relatively early stage. B symptoms (unexplained weight loss of > 10% within 6 months and/or unexplained fever > 38°C and/or night sweats), while nonspecific, may be suggestive of malignant lymphoma in patients with enlarged cervical lymph nodes and should prompt an immediate histologic examination. This is the only way to establish a diagnosis and also identify the tumor subgroup, which usually has a significant bearing on treatment and prognosis. Immunohistochemical and/or molecular biological techniques are generally applied. The best sites for finding additional, extracranial manifestations of lymphoma are the mucous membranes of the upper aerodigestive tract and particularly the lymphatic structures of Waldeyer’s ring.

Further management: Once the diagnosis of malignant lymphoma has been confirmed histologically, the patient should be referred at once to a hematologic oncology center so that the disease can undergo Ann Arbor staging (see textbooks of internal medicine) and appropriate management.

Lymph-Node Metastases

Synonym: malignant lymphadenopathy

Etiology: Lymph-node metastases are predominantly of epithelial origin; the lymphatic metastasis of mesenchymal tumors (sarcomas) is relatively rare. The most common primary tumors are carcinomas of tributary mucosal areas of the upper aerodigestive tract, salivary glands, and thyroid gland (Table 16.4). But tumors of any other organs may also metastasize to the cervical lymph nodes. It is not uncommon for cancers of the lung, breast, stomach, kidneys, cervix, and prostate to metastasize to cervical nodes. The ”sentinel node” (signal node, Virchow node) in the left supraclavicular fossa is considered a fairly reliable indicator of gastric carcinoma.

Owing to the anatomy of the lymphatic drainage system, the site of a cervical lymph-node metastasis shows some degree of correlation with the site of the primary tumor (see Fig. 16.2, p. 314).

Symptoms: Cervical lymph-node metastases generally present as a painless, more or less fast-growing swelling on one or both sides of the neck (Fig.16.19). The underlying primary tumor may be asymptomatic, or it may be possible to elicit symptoms of a primary tumor by careful questioning. Information from the patient on the duration of symptoms may be unreliable.

Diagnosis: During the clinical examination, attention is given to the number and size of enlarged lymph nodes and their mobility relative to their surroundings. B-mode ultrasound can provide an accurate size determination and can also help define the relation of the masses to the large vessels and muscles in the neck.

<table>
<thead>
<tr>
<th>Table 16.4 Incidence of cervical lymph-node metastases associated with carcinomas of the upper aerodigestive tract</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tumor location</td>
</tr>
<tr>
<td>Oral cavity</td>
</tr>
<tr>
<td>Oropharynx</td>
</tr>
<tr>
<td>Nasopharynx</td>
</tr>
<tr>
<td>Hypopharynx</td>
</tr>
<tr>
<td>Supraglottis</td>
</tr>
<tr>
<td>Glottis</td>
</tr>
<tr>
<td>Nasal cavity and paranasal sinuses</td>
</tr>
<tr>
<td>Salivary glands</td>
</tr>
<tr>
<td>Thyroid gland</td>
</tr>
</tbody>
</table>

* Depends on age and histologic subtype
Doppler ultrasound is helpful for investigating vascular compression and infiltration.

**Search for a primary tumor:** Whenever a cervical lymph node is suspicious for metastasis, the upper aerodigestive tract should be screened for a primary tumor by ear, nose, and throat mirror examination, endoscopy, and B-mode ultrasound.

- If a lesion is found that is suspicious for a primary tumor, it should be investigated further as described on pp. 76–77, 91 and 104–107.
- If a primary tumor is not detected on clinical examination, it is advisable to remove a suspicious lymph node for histologic examination and use this result to direct further evaluation. If squamous cell carcinoma is found, indirect examination of the upper aerodigestive tract should be supplemented by detailed *endoscopy* of the nasopharynx, pharynx, larynx, trachea, bronchial system, and esophagus. Since the base of the tongue, tonsils, and nasopharynx may harbor a submucous carcinoma that is not accessible to direct visual detection, these sites should be carefully *probed* and also routinely evaluated by *deep biopsies, tonsillectomy,* and also by *curetage* in the nasopharynx. Ultrasound, MRI, and PET can also be used to locate a primary tumor. Because even primaries located outside the upper aerodigestive tract may metastasize to the cervical nodes, the search should be extended to other primary sites if a tumor is not detected in that region.

- If a diligent search (see above) fails to disclose a primary tumor (this occurs in approximately 5–10% of cases and usually involves squamous cell carcinoma), then the metastasis is classified as a “carcinoma of unknown primary” (CUP syndrome; see also 16.12). The most common site of involvement in CUP syndrome is a solitary cervical lymph node.

**Staging:** Table 16.5 reviews the TNM classification system for regional lymph-node involvement by tumors of the upper aerodigestive tract.

**Treatment of the cervical lymphatics in patients with head and neck tumors:** Most cancers of the upper aerodigestive tract have already metastasized to the cervical lymph nodes at the time of diagnosis (Table 16.4). It is necessary, therefore, to include the cervical lymphatics in primary treatment planning.

**Neck dissection:** In most cases, a neck dissection is performed concurrently with surgical treatment of the primary tumor. The indication for a neck dissection depends on the location and spread of the primary tumor and on the clinical status of the cervical lymph nodes. A *therapeutic* neck dissection is done in patients with clinically positive cervical nodes (N+), while an *elective* neck dissection is done in patients with clinically negative nodes (N0). The need for an elective neck dissection is dictated by the high rate of occult cervical lymph-node metastasis that is associated with many primary tumor sites (even tumors at a low T category).

- A modified radical ("functional") neck dissection involves the removal of all cervical lymph nodes on one side (Fig. 16.20 a). It was developed as a modification of the radical neck dissection described below.
16.12 Carcinoma of unknown primary (CUP) syndrome

The *etiopathogenesis* of CUP syndrome is uncertain. Four main theories have been advanced:

1. A primary tumor initially present in the mucosa of the upper aerodigestive tract induces regional metastasis and then regresses completely while the regional metastases proliferate (*disappeared primary*).

2. The primary tumor has a weak local growth potential but a strong propensity for metastasis. The primary tumor is not detected by ordinary clinical methods because of its small size, but theoretically it would be detectable if more refined methods were used (*hidden primary*).

3. An ectopic mass of epithelial tissue within a lymph node undergoes malignant degeneration.

4. The lesion is not a lymph-node metastasis but a primary branchiogenic carcinoma.

*Treatment* for CUP syndrome consists of surgery (neck dissection), radiotherapy and, if necessary, chemotherapy. The *5-year survival rate* is approximately 50% or more. Attention is given to possible metachronous manifestations of the primary tumor during oncologic follow-up.

- *A radical neck dissection* additionally removes the sternocleidomastoid muscle, internal jugular vein, and accessory nerve, regardless of whether or not they are involved by tumor (Fig. 16.20b). This operation causes limitation of head and shoulder motion due to resection of the accessory nerve and sternocleidomastoid muscle. The resection of one internal jugular vein generally has no adverse sequelae owing to increased compensatory drainage by the vertebral veins, paravertebral plexus, and the internal jugular vein on the opposite side.

Because these compensatory mechanisms take time to develop, both internal jugular veins should never be resected at the same time, as the resulting acute bilateral decrease in venous outflow would lead to raised intracranial pressure (Monroe–Kellie doctrine), with a high mortality.

Within a few weeks after unilateral resection of the jugular vein, the extrajugular venous channels have increased so much in capacity that the contralateral internal jugular vein can also be resected with very little risk.

- The neck dissection may be limited to selected regions (*selective neck dissection*), depending on the location and extent of the primary tumor.

*Radiotherapy*: Postoperative (adjuvant) radiotherapy may be indicated, depending on the extent of the primary tumor and of cervical lymph-node metastasis. With very extensive (inoperable) tumor growth or smaller tumors with no clinical evidence of neck metastasis, the cervical soft tissues can be treated by radiotherapy alone.
## Larynx and Trachea

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### 17.1 Embryology, Anatomy, and Physiology of the Larynx and Trachea

The complex functions of the larynx rely on the precise, coordinated interaction of intricate anatomic structures, which much carry out the diametrically opposed functions of respiration and phonation. The development of the larynx (17.1) is the basis for understanding many diseases of this organ, which in humans is endowed with the unique function of producing speech. Malformations of the larynx in particular cannot be properly understood without a knowledge of embryology. The trachea transports the inspiratory air to the lungs and channels the expiratory air from the lungs to the mouth and nose. Although the trachea consists of metameric subdivisions, many tracheal diseases show typical sites of predilection.

## Anatomy of the Larynx

**Cartilaginous Skeleton, Ligaments, and Muscles**

The skeleton of the larynx (Fig. 17.1) is composed of the hyaline **thyroid cartilage**, **cricoid cartilage**, and **arytenoid cartilages**, as well as the fibroelastic cartilage of the epiglottis and the tips of the functionally insignificant accessory cartilages located above the arytenoids (the corniculate and cuneiform cartilages). In males, the thyroid cartilage forms an externally visible prominence also known as the Adam’s apple. The laryngeal cartilages begin to ossify at about 20 years of age.

Each of the inferior horns of the thyroid cartilage articulates with the cricoid cartilage, forming the hinged cricothyroid joint, which permits tilting movements in the sagittal plane (Fig. 17.2). Each of the arytenoid cartilages has an anterior vocal process, which attaches to the posterior end of the corresponding vocal cord, and a posterolateral muscular process. The base of the arytenoid cartilage articulates with the superior border of the cricoid cartilage, forming a cricoarytenoid joint of variable shape, which permits rotation and gliding movements. The muscles that attach to the muscular process are particularly active in rotating the arytenoid cartilage about its longitudinal axis. Changing the position of the vocal cords alters the shape and size of the opening (glottis) between the two vocal folds.

---

**Fig. Laryngeal cartilages**

<table>
<thead>
<tr>
<th>a</th>
<th>b</th>
<th>c</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Anterior view of the larynx projected onto the neck" /></td>
<td><img src="image" alt="Body of hyoid bone" /></td>
<td><img src="image" alt="Epiglottis" /></td>
</tr>
<tr>
<td><strong>Sternocleidomastoid muscle</strong></td>
<td><strong>Lesser horn</strong></td>
<td><strong>Thyroepiglottic ligament</strong></td>
</tr>
<tr>
<td><strong>Thyroid gland</strong></td>
<td><strong>Greater horn</strong></td>
<td><strong>Lateral thyrohyoid ligament</strong></td>
</tr>
<tr>
<td><strong>Clavicle</strong></td>
<td><strong>Foramina for superior laryngeal artery and superior laryngeal nerve</strong></td>
<td><strong>Corniculate cartilage</strong></td>
</tr>
<tr>
<td><strong>Sternum</strong></td>
<td><strong>Superior horn</strong></td>
<td><strong>Arytenoid cartilage</strong></td>
</tr>
<tr>
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<td><strong>Thyrohyoid membrane</strong></td>
<td><strong>Cricoid cartilage lamina</strong></td>
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<tr>
<td><strong>Medial cricothyroid ligament</strong></td>
<td><strong>Inferior horn</strong></td>
<td><strong>Tracheal cartilages</strong></td>
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<tr>
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<td><strong>Cricothyroid joint</strong></td>
<td><strong>Membranous posterior wall</strong></td>
</tr>
<tr>
<td><strong>Arch of cricoid cartilage</strong></td>
<td><strong>Cricotracheal ligament</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Tracheal cartilages</strong></td>
<td><strong>Annular ligaments of trachea</strong></td>
<td></td>
</tr>
</tbody>
</table>

*a Anterior view of the larynx projected onto the neck (after Lumley). b, c Anatomy of the laryngeal skeleton (after Tillmann, see p.410).*
The paired vocal cords (vocal folds) stretch between the vocal process of the arytenoid cartilage and the inner surface of the thyroid cartilage at the junction of its lower and middle thirds. The stalk (petiole) of the epiglottis is attached to the posterior surface of the thyroid cartilage by the thyroepiglottic ligament, which may also contain contractile muscle fibers (thyroepiglottic muscle). The thyrohyoid membrane extends from the hyoid bone to the thyroid cartilage. This membrane is traversed by the superior laryngeal artery and vein and by the internal branch of the superior laryngeal nerve, which supplies sensory innervation to the mucosa lining the upper portions of the larynx. The cricothyroid membrane (synonym: cricothyroid ligament) connects the cricoid cartilage to the thyroid cartilage and is divided in a cricothyrotomy (coniotomy; see Fig. 17.3, p. 379). Thus, extrinsic ligaments and connective-tissue membranes serve to anchor the larynx to its surroundings while numerous joints, ligaments, and membranes between the cartilaginous elements interact with the various muscles to ensure the coordinated functional movements of laryngeal structures. Intrinsic laryngeal muscles and one extrinsic muscle (Table 17.1, Fig. 17.3) open and close the glottis and tense the vocal cords. The muscles that close the glottic plane predominate over the single muscle that opens it (posterior cricoarytenoid muscle) by a force ratio of approximately 3:1.

As noted earlier, the larynx is the narrowest point in the upper respiratory tract, making it particularly susceptible to obstructions. The cricoid cartilage encircles the subglottis like a ring, imparting a mechanical stability that helps to prevent collapse of the laryngeal
17.1 Embryology of the larynx and trachea

The epithelium of the respiratory tract (larynx, tracheobronchial system, lungs) develops as an outpouching of the ventral wall of the foregut (laryngotracheal groove, day 26 of gestation) and is therefore of entodermal origin. The tracheoesophageal septum forms during the fourth week, completely separating the primordia of the esophagus and the larynx/trachea/lung (laryngotracheal tube) from one another (Fig. a, b, c). The mesoderm that surrounds the primitive respiratory tract differentiates into cartilaginous, muscular and connective tissue so that the entodermal respiratory tract is finally encased by mesodermal tissue. The paired arytenoid eminences and the unpaired epiglottic swelling impart a T-shaped appearance to the laryngeal inlet (Fig. d).

With further chondrogenesis, epithelial proliferation causes a temporary closure of the laryngeal lumen (7th–10th weeks). This is followed by recanalization of the larynx and the development of the mucosal folds (ventricular and vocal folds, starting in the 10th week; Fig. e).

The hyoid bone is derived from the second and third branchial arches, and the laryngeal cartilages except for the epiglottis, arytenoid cartilages, and accessory cartilages are derived from the fourth and sixth branchial arches. The epiglottis, arytenoid cartilages, and accessory cartilages develop secondary from the mesenchyma ("secondary cartilage"). The cells that give rise to the laryngeal muscles migrate to the larynx from the myotomes of the cranial somites. The associated branchial arch nerves are branches of the vagus nerve (fourth branchial arch: superior laryngeal nerve, sixth branchial arch: recurrent laryngeal nerve).

During embryonic and fetal life, the epiglottis is still located behind the soft palate and extends to the nasopharynx. By the time of birth, the epiglottis has already migrated caudally so that its tip is level with the axis dens. The laryngeal skeleton continues to descend with ageing, after puberty reaching its definitive position at about the level of the C5 vertebral body. It is no longer believed that the larynx develops from two separate parts that fuse together during development (the buccopharyngeal and tracheobronchial buds), with separation of the supra- and subglottic lymphatic drainage systems (compartamentalization of the larynx). Nevertheless, the lymphatic drainage of the larynx is subdivided anatomically into supra- and subglottic, glottic, and sublaryngeal pathways.

Fig. Source: Sadler, Moore and Persaud, see p. 410.

Nerve Supply

The larynx and the trachea derive their motor and sensory innervation from the superior laryngeal nerve and recurrent laryngeal nerve, both of which arise from the vagus nerve. The superior laryngeal nerve supplies motor innervation to the extrinsic laryngeal muscle (anterior cricothyroid) with its external branch, while its internal sensory branch supplies the mucosa of the upper larynx including the glottic plane. The recurrent laryngeal nerve supplies sensory fibers to the laryngeal mucosa below the glottis and to the tracheal mucosa. The recurrent laryngeal nerve also supplies all the intrinsic muscles of the larynx. This means that the muscles that open and close the glottic

skeleton (Table 17.2). It also ensures that any mucosal swelling at that level will be directed toward the lumen.

This is particularly important in newborns and infants, because a circumferential swelling of the glottic and subglottic mucosa by just 1 mm can cause more than a 60% reduction in total luminal cross section.

It should also be recalled that the airway resistance in a stenotic segment is directly proportional to the length of the stenosis and inversely proportional to the fourth power of the radius (Hagen–Poiseuille law, Fig. 17.4).
Table 17.1 Laryngeal muscles

<table>
<thead>
<tr>
<th>Function</th>
<th>Muscle</th>
<th>Origin, insertion, nerve supply, function</th>
</tr>
</thead>
</table>
| Opening the glottis       | Posterior cricoarytenoid muscle (a variable slip from the cricoid cartilage plate to the inferior horn of the thyroid cartilage is termed the ceratocicoid muscle.) | O: Lamina of cricoid cartilage (posterior surface)  
I: Muscular process of arytenoid cartilage  
N: Recurrent laryngeal nerve  
F: Rotates the arytenoid cartilage outward about a vertical axis, tilts it laterally |
| Closing the glottis       | Lateral cricoarytenoid muscle (lateralis muscle)                       | O: Arch of cricoid cartilage  
I: Muscular process of arytenoid cartilage  
N: Recurrent laryngeal nerve  
F: Rotates the arytenoid cartilage inward about a vertical axis |
|                           | Interarytenoid muscle (transverse and oblique interarytenoid muscles) | O: Arytenoid cartilage (posterior surface)  
I: Contralateral arytenoid cartilage  
N: Recurrent laryngeal nerve  
F: Narrows the intercartilaginous part of the vocal fold by adducting the arytenoid cartilages |
|                           | Thyroarytenoid muscle, lateral part                                   | O: Thyroid cartilage (inner surface, laterally adjacent to the vocalis muscle)  
I: Arytenoid cartilage  
N: Recurrent laryngeal nerve  
F: Rotates the arytenoid cartilage inward about a vertical axis |
| Tightening the vocal cords| Cricothyroid muscle                                                    | O: Arch of cricoid cartilage (outer surface)  
I: Thyroid cartilage  
N: External branch of superior laryngeal nerve  
F: Tilts the cricoid cartilage relative to the thyroid cartilage on a transverse axis |
|                           | Thyroarytenoid muscle, medial part (vocalis muscle)                   | O: Thyroid cartilage (inner surface)  
I: Vocal process and oblong fovea of arytenoid cartilage  
N: Recurrent laryngeal nerve  
F: Controls the tension on the vocal fold by isometric contraction |

plane receive their motor innervation from the same nerve.
The course of the recurrent laryngeal nerve differs between the left and right sides. While the larger left recurrent nerve curves around the aortic arch, the right recurrent nerve winds around the subclavian artery. On each side, the nerve then passes between the trachea and esophagus and enters the larynx behind the inferior horn of the thyroid cartilage (Fig. 17.5). The difference in length between the two nerves is a few centimeters. The different calibers of the nerve fibers offset this difference, enabling the impulses to reach the muscles at the same time.

**Vascular Supply**
The glottic plane divides the blood supply to the larynx into two territories. The blood supply at the supraglottic and glottic levels is derived from the superior laryngeal artery arising from the external carotid, while the subglottic areas are supplied by the inferior
laryngeal artery arising from the subclavian artery and thyrocervical trunk. **Venous drainage** of the larynx is handled by the superior thyroid vein, which drains into the internal jugular vein, and by the inferior thyroid vein, which drains into the brachiocephalic vein (Fig. 17.6).

Except for the glottic region, the lymphatic vessels in the larynx are well developed and are more numerous above the glottis than below (see above). The very dense lymphatic capillary network of the supraglottis drains into the vertical cervical lymphatic chains (deep cervical lymph nodes) and especially into the lymph nodes at the junction of the facial and internal jugular veins (junctional nodes). The ipsilateral lymphatic drainage is complemented by significant con-
Epithelial Lining

Like the trachea, most of the endolarynx is lined by respiratory epithelium—a stratified ciliated epithelium with interspersed goblet cells. A stratified, nonkeratinized and sometimes keratinized squamous epithelium is found with great regularity in the laryngeal epiglottis, ventricular folds, and vocal cords and occasionally in other areas of the larynx. The presence of this squamous epithelium, and its expansion with aging, are often interpreted as responses to increased mechanical stresses.

Vocal Cord Histology

The histologic structure of the vocal cord is shown in Fig. 17.8. Reinke’s space is a subepithelial plane in the vocal cord that contains no glands or lymphatic capillaries.

Relations of the Larynx

The larynx is bounded cranially by the free margins of the epiglottis, the aryepiglottic fold, and the interarytenoid eminence. Caudally, the inferior border of the cricoid cartilage marks the junction of the larynx with the trachea. The larynx forms the narrowest point in the respiratory tract between the nasopharynx and trachea.

The laryngeal cavity is divided into three parts in relation to the glottis (Fig. 17.9):

- **Supraglottis**: laryngeal inlet to the sinus of Morgagni
- **Glottis**: plane of the vocal cords plus approximately 1 cm of their subglottic flank
- **Subglottis**: extending to the lower border of the cricoid cartilage

The vocal cord consists of the vocal ligament, vocalis muscle, and associated mucosal cover.
The glottis (rima glottidis) is the opening between the vocal cords. It has a membranous part backed by the vocal ligament and a cartilaginous part formed by the vocal process of the arytenoid cartilage. The transglottic space occupies the region from the ventricular folds, or false vocal cords, to the glottis (see Fig. 17.9).

Physiology of the Larynx

The larynx serves as the organ of phonation (vocal cords closed = phonation position) and as an airway (vocal cords open = respiratory position). It keeps the foodway and airway separate during food ingestion.

Protective Mechanisms

The most important protective mechanism is the immediate and complete reflex closure of the vocal cords in response to the pharyngeal phase of swallowing (see also Fig. 5.4, p. 102). Simultaneous contraction of the suprahypoid and infrayoid muscles elevates the entire laryngeal skeleton by 2–3 cm, while the base of the tongue bulges over the larynx and presses the epiglottis downward, directing the food bolus past and behind the larynx to the esophageal inlet. The epiglottis itself has no essential importance either in closing off the larynx or preventing aspiration. If food material does transgress the glottic plane, the cough reflex activates another important mechanism that protects the lower airways. After a deep reflex inspiration, the glottis closes tightly, allowing a rise in intrathoracic pressure. The glottis then opens widely and rapidly to allow forceful expulsion of the aspirated material.

Manipulations within the larynx (e.g., intubation) can stimulate vagovagal reflex pathways that may incite a cardiac arrhythmia, bradycarden, or even cardiac arrest.

Anatomy of the Trachea

The trachea is suspended from the cricoid cartilage—the narrowest stiff-walled element in the upper respiratory tract—by the cricoth caveal ligament. In many cases, however, the upper ring of the trachea is also fused to the cricoid cartilage. The trachea has a length of 10–13 cm in adults, extending from the level of the C6–C7 vertebral body to the T4–T5 vertebra, replicating the curvature of the spine and bifurcating at the T4–5 level into the right and left main bronchi (Fig. 17.10). The trachea is very superficial in its upper portion, but as it descends it passes behind the sternum to a depth of 4–5 cm. In other important relations, the anterior and posterior walls of the trachea are closely related to the isthmus and lobes of the thyroid gland and to the recurrent laryngeal nerves.

The lumen of the trachea is maintained by 12–20 hyaline cartilage “rings” that are horseshoe-shaped (open posteriorly) and are interconnected by strong collagenous and elastic connective-tissue fibers, the annular ligaments. The cervical part of the trachea is 6–7 cm long and contains six to eight cartilaginous rings. The membranous posterior wall of the trachea is related to
the anterior wall of the esophagus (Fig. 17.10b). Foreign bodies and tumors of the esophagus may impinge upon the posterior surface of the trachea, constricting the airway lumen.

The transverse diameter of the trachea averages 13–20 mm (13–16 mm in women, 16–20 mm in men). The trachea is lined by two rows of ciliated epithelium with goblet cells. The mucociliary clearance mechanism in the trachea is directed toward the larynx and contributes to the cleaning, warming, and humidification of the inspired air.

The blood supply to the trachea is provided mainly by the inferior thyroid artery (from the thyrocervical trunk) and to a lesser degree by the superior thyroid artery (see also Fig. 17.10c). As in the subglottic region, the lymphatic drainage of the trachea is handled by the vertical lymphatic chain in the neck (deep cervical lymph nodes) and by the paratracheal and mediastinal groups of lymph nodes.

The trachea receives its nerve supply from the vagus nerve and sympathetic trunk.

For anterior aspect, see also Fig. 17.1, p. 338.
Source: Tillmann, see p. 410.
17.2 Symptomatology and Examination of the Larynx and Trachea

The cardinal symptoms of laryngeal disorders are based on the functions of the larynx as the “mediator” between the airway and foodway and as the organ of phonation. Besides indirect examination with a mirror or endoscope, which yields both morphologic and functional information, diseases of the larynx can be investigated by sectional imaging modalities and function tests (e.g., electromyography).

Cardinal Symptoms

Larynx

The main symptoms of laryngeal disorders are inspiratory stridor, dyspnea, phonation problems (e.g., hoarseness), and eating difficulties. These symptoms may occur separately or in combination.

Trachea

The cardinal symptoms of diseases involving the trachea are cough, sputum production, and respiratory distress due to narrowing of the tracheal lumen. Tracheal stenosis—whether due to inflammation, scarring, or neoplasia—is commonly manifested by an inspiratory and expiratory stridor. When the history is taken, the patient should be questioned about blood in the sputum, exercise-dependence or position-dependence of respiratory problems, exposure to potentially harmful inhalants, and any concomitant diseases of the upper aerodigestive tract.

Methods of Examination

Larynx: Inspection and Palpation

Inspection of the neck provides initial information on the configuration of the larynx and possible inspiratory jugular retraction due to laryngotracheal obstruction. Also, the mobility of the larynx should be observed during swallowing to exclude inflammatory or neoplastic fixation and any extrinsic mass effects on the laryngeal skeleton (e.g., from tumors in the neck). Bimanual palpation of the laryngeal skeleton is done from the front or from behind to check for contour irregularities and sites of tenderness. It may include the thyroid gland, and the cervical soft tissues should be palpated if malignancy is suspected.

Indirect Laryngoscopy

Indirect examination of the larynx can be performed with a laryngeal mirror or with a rigid or flexible endoscope. Classic indirect laryngoscopy requires a laryngeal mirror of suitable size, a light source, a head mirror, and a gauze sponge (Fig. 17.11a). Normally, the examination is done with the patient sitting upright, after any dentures have been removed. The (right-handed) examiner grasps the protruded tongue with a gauze sponge in the left hand, placing the thumb on top of the tongue and the middle finger under the tongue.

When the tongue is pulled forward, care is taken not to injure the frenulum on the lower incisors. The index finger of the left hand is braced on the upper teeth and is used to retract the upper lip. The light from the head mirror is directed toward the uvula. The reflective surface of the laryngeal mirror should be gently warmed with the hand or wetted with alcohol solution to prevent fogging by the patient’s breath and body heat. The examiner holds the laryngeal mirror in the right hand like a pencil and advances it beneath the palate to the uvula. Touching the base of the tongue or the posterior wall of the pharynx may evoke a gag reflex. The uvula is now lifted on the back surface of the mirror, which is angled approximately 45°, and gently pushed backward and upward so that the larynx can be brought into view (Fig. 17.11a, b).

The larynx should always be examined in the respiratory position (tell the patient to take a deep breath) and in the phonatory position (have the patient say “Hee”) in order to assess laryngeal function (Fig. 17.11c, d). Besides the larynx, the examiner can evaluate the base of the tongue, the oropharynx and hypopharynx, and even the anterior wall of the cervical trachea by viewing through the open glottis in the respiratory position. The posterior portions of the larynx (the posterior commissure and subglottic space) are best examined with the examiner sitting and the patient standing with the head tilted forward (Fig. 17.11e). The anterior portions of the larynx (the laryngeal epiglottic surface,
petiole, and anterior commissure) are best examined with the examiner standing and the patient sitting with the head tilted backward (Fig. 17.11f).

Keep in mind that this examination provides an inverted “mirror image” of the larynx (see Fig. 17.11b).

In recent years, routine mirror examination of the larynx has been increasingly superseded by telescopic laryngoscopy using a rigid endoscope with a 90° wide-angle view that can brightly illuminate and also magnify the area being examined (Fig. 17.12a). The telescopic examination is performed in the same way as the mirror examination and yields the same findings, except that the examined areas are not inverted but are displayed in their natural position (Fig. 17.12b, c). This true orientation, plus the option for magnification, also makes it easier to perform biopsies and tissue ablation in the conscious patient.

Flexible endoscopy (Fig. 17.13): In a small number of patients, the larynx cannot be inspected by mirror or telescopic laryngoscopy due to a powerful gag reflex, despite the use of topical anesthesia. The laryngeal structures can be inspected in these patients by using a flexible nasopharyngeal laryngoscope. Another indication for flexible laryngeal endoscopy, in which the endoscope is passed through the nose into the nasopharynx and then down the oropharynx into the larynx, exists in cases where it is necessary to combine laryngeal inspection with tracheobronchoscopy. The flexible endoscopes used for this purpose have an outer diameter of 2.7–6.0 mm but deliver poorer image quality than the rigid telescopes described above.

Direct Laryngoscopy

Direct laryngoscopy, which provides a direct view into the larynx, is most commonly performed under general anesthesia, using either intubation anesthesia or injector ventilation without an endotracheal tube. With the head fully extended, an illuminated rigid tube (laryngoscope) is advanced straight through the mouth to the plane of the larynx (Fig. 17.14). The laryngoscope may be held in place by a lever arm supported on the patient’s chest or a tray suspended above the patient’s chest (“suspension laryngoscopy” or “suspension autoscopy”). Direct laryngoscopy is usually done as a microscope-assisted procedure (microlaryngoscopy or microlaryngoendoscopy, MLE), permitting the detailed scrutiny of laryngeal abnormalities. Other instruments such as laser delivery systems (Fig. 17.14b) can also be used for many therapeutic applications.
Computed tomography and magnetic resonance imaging yield more diagnostic information than plain radiographs and conventional tomography. They are useful in defining the precise extent of laryngeal and tracheal masses.

Ultrasoundography can be used to evaluate prelaryngeal and paralaryngeal soft tissues and determine whether intralaryngeal masses have eroded through the laryngeal skeleton and spread contiguously to extralaryngeal structures. Given the physical principles of sound propagation (“air is the enemy of ultrasound”), intralaryngeal structures cannot be clearly visualized. The same applies to ultrasound evaluation of the trachea.

Laryngography, in which the inner surface of the larynx is wetted with a radiographic contrast medium, no longer has any practical importance today.

Function Tests

Electromyography of the laryngeal muscles is described in 17.9 (see p. 380). Phoniatric studies (stroboscopy, frequency/intensity profile) are covered in 18.1 (see pp. 386–389).

Methods of Examining the Trachea

During inspection, attention is first given to any cervical masses and their relationship to the trachea and respiratory excursions. The trachea is palpated with the head flexed forward (to relax the airway), and tracheal mobility is assessed in relation to the surrounding structures. Because diseases of the thyroid gland may directly affect the upper respiratory tract (dyspnea caused by a hemorrhagic cyst of the thyroid gland, tracheomalacia due to thyroid enlargement, vocal cord paralysis due to a malignancy), the thyroid gland should also be palpated. The upper part of the trachea can often be viewed through the glottis during indirect laryngoscopy. But in cases where tracheal pathology is strongly suspected, endoscopic inspection of the entire trachea is essential for definitive confirmation. Both flexible and rigid optical systems are available for this purpose.

Flexible trache(broncho)scopy can, in principle, be performed under topical anesthesia in the conscious patient. The endoscope (2.7–6.0mm outer diameter) is introduced transnasally or transorally and advanced through the glottis into the trachea (Fig. 17.13). This technique not only permits a detailed evaluation of the trachea, but if necessary the entire bronchial tree can be inspected out to the level of the subsegmental bronchi. Flexible endoscopes with a working channel and suction-irrigation system can be used for diagnostic procedures (specimen retrieval) and also for therapeutic manipulations (fiberoptic intubation, foreign-

Imaging of the Larynx and Trachea

Plain radiographs in the anteroposterior and lateral projections can demonstrate the skeletal framework of the larynx. They are particularly useful in patients with laryngeal fractures or a suspected foreign body. Stenoses of the trachea and larynx, especially when located at the laryngotracheal junction, can also be demonstrated by plain radiography and by motion-controlled tomography.
body extraction even in the peripheral bronchial tree, stent insertion, laser tumor ablation).

Endoscopic examination of the trachea with rigid, illuminated systems (rigid tracheobronchoscopy) is generally performed under general anesthesia, and ventilators can be adapted for this purpose. With the patient’s neck fully extended, the endoscope barrel is advanced through the larynx into the trachea. The limited view of peripheral portions of the bronchial system can be somewhat improved by using an oblique scope. Either a flexible or rigid system can be used for foreign-body removal in the tracheobronchial system, depending on the nature, configuration, and location of the foreign body. More extensive manipulations in the tracheobronchial system that require a clear view, due particularly to an increased risk of bleeding, are still within the domain of rigid endoscopy.

**Plain radiographs** of the upper airways in the posteroanterior (PA) and lateral projections can demonstrate the tracheal air column along with any intraluminal lesions or masses that are causing extrinsic tracheal compression. One PA radiograph taken after forced inspiration and another taken during a subsequent Valsalva maneuver will show evidence of any abnormal weak spots in the tracheal wall (tracheomalacia). The functional impact of stenotic lesions of the larynx and trachea can be objectively evaluated by **pulmonary function testing** and **body plethysmography** (details can be found in textbooks of internal medicine).

The capabilities of direct laryngoscopy (a) can be expanded by using a CO2 laser with a micromanipulator (b). This allows for precise tissue resection under microscopic control.
17.3 Malformations of the Larynx and Trachea

Laryngeal anomalies have a reported incidence of one in 10,000 to one in 50,000 newborns. The severity of the malformations ranges from the extremely rare and almost always fatal laryngeal atresia to the very common and usually harmless laryngomalacia. Besides the cardinal symptoms of inspiratory stridor and dyspnea, other possible features are dysphonia and dysphagia. The anomalies may be manifested at the supraglottic, glottic, or subglottic level. As a rule, the clinical suspicion of a laryngeal malformation can be confirmed only by endoscopic examination.

Malformations of the Larynx

Laryngomalacia

Epidemiology: Laryngomalacia is the most frequent cause of congenital stridor, accounting for 60–75% of cases.

Etiopathogenesis: The supraglottic structures (epiglottis, arytenoid cartilages) in particular are abnormally soft and pliable, causing them to collapse inward during inspiration. Neurologic abnormalities and infectious processes may also have causal significance.

Symptoms: Typically, a low-pitched inspiratory stridor is audible from birth. It may be constant or intermittent and may change with the position of the child, becoming louder in the supine position and quieter in the prone position. Life-threatening airway compromise is extremely rare, but feeding difficulties are occasionally seen.

Diagnosis: Laryngoscopy is helpful in making the diagnosis. In typical cases of laryngomalacia, the aryepiglottic folds are shortened and the arytenoid cartilages are bowed anteriorly and toward each other. Another typical finding is a soft, pale, “omega-shaped” epiglottis with its lateral edges curled inward. In extreme cases the epiglottis completely covers the laryngeal inlet.

Differential diagnosis: Rare cleft anomalies of the larynx are distinguished from laryngomalacia by the additional presence of dysphagia with recurrent episodes of aspiration. Other causes of stridor such as congenital cysts and laryngoceles can be excluded by laryngoscopy.

Treatment: The stridor should resolve without treatment during the first 2 years of life as the laryngeal skeleton becomes more rigid. Very rare cases may require a temporary tracheotomy, however. It is particularly important to explain the condition to the anxious parents and reassure them that it is usually harmless.
### Table 17.3 Malformations of the trachea

<table>
<thead>
<tr>
<th>Condition</th>
<th>Symptoms and findings</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tracheal agenesis or atresia</td>
<td>Ineffictful breathing efforts by the newborn with no lung ventilation; if a tracheoesophageal fistula is also present, limited ventilation may occur</td>
<td>Generally untreable</td>
</tr>
<tr>
<td>Tracheal web</td>
<td>Stridor, depending on the degree of airway obstruction; discrete luminal narrowing by fibrous tissue</td>
<td>Tracheal expansion by perforation and bougie dilation</td>
</tr>
<tr>
<td>Congenital tracheal stenosis</td>
<td>Stridor, depending on the degree of airway obstruction; luminal narrowing involves a longer segment and deeper layers of the trachea</td>
<td>Bougie dilation is often unsuccessful; patient may require a temporary tracheotomy and resection of the stenotic segment</td>
</tr>
<tr>
<td>Stenosis with or without a tracheoesophageal fistula (Fig. 17.15)</td>
<td>Stridor, depending on the degree of luminal narrowing; airway obstruction during feeding, with cyanosis and signs of aspiration; open connection between the tracheal lumen and esophagus</td>
<td>Surgical correction immediately after birth</td>
</tr>
<tr>
<td>Tracheomalacia (occurs independently of laryngomalacia)</td>
<td>Biphasic stridor due to luminal narrowing; difficulty in coughing up secretions</td>
<td>Usually self-limiting as the tracheal framework becomes more stable at age 12–18 months; conservative measures may include humidifying the inspired air</td>
</tr>
<tr>
<td>Tracheal deformities in the setting of vascular malformations</td>
<td>Compression of the trachea by anomalies of the aortic arch or its major branches, causing airway obstruction; concurrent compression of the esophagus leads to dysphagia</td>
<td>Need for surgical intervention depends on the severity of symptoms</td>
</tr>
<tr>
<td>Malformations of individual tracheal cartilages</td>
<td>Circumscribed narrowing of the trachea, especially on expiration, due to partial absence of supporting structures; expiratory stridor</td>
<td>Conservative, expectant approach; some cases may require a temporary tracheotomy and/or reconstruction</td>
</tr>
<tr>
<td>Tracheal cysts and tracheoceles</td>
<td>Outpouchings of the tracheal wall; symptoms depend on the degree of luminal narrowing</td>
<td>Surgical excision may be required</td>
</tr>
</tbody>
</table>

![Fig. Types of tracheoesophageal malformation](source)

Source: Sadler, see p. 411.
Congenital Laryngeal Web and Glottic Atresia

Perhaps the best-known laryngeal anomaly, if not the most common, is the laryngeal web or glottic web.

Etiopathogenesis: This membrane-like stenosis of the glottic plane results from incomplete recanalization of the glottic lumen during embryogenesis (see 17.1, p. 340). If there is a complete failure of recanalization, the result is glottic atresia.

Symptoms: The main symptoms are inspiratory stridor, which is present at rest only with a large web, and a breathy, aphonic voice. In most cases the web is not diagnosed in infancy but is discovered later when the patient is intubated for general anesthesia or during a routine examination of the larynx (Fig. 17.16). A child with glottic atresia is cyanotic at birth and makes scant, ineffectual breathing movements with no respiratory sounds. Only an immediate, emergency tracheotomy can save the life of a newborn with this rare anomaly. In exceptional cases, laryngeal atresia coexists with a tracheoesophageal fistula that permits adequate respiration.

Diagnosis: The web is usually visible at laryngoscopy, stretching across the anterior commissure and leaving a residual posterior glottic airway sufficient for respiration.

Treatment: In most cases the web can be endoscopically divided during microlaryngoscopy, although there is a risk of recurrent synchia formation in the anterior commissure. To prevent this, it is occasionally necessary to place a temporary stent (keel) between the vocal folds through an extralaryngeal approach.

Congenital Subglottic Stenosis

Epidemiology: Congenital subglottic stenosis is the most common stenosing anomaly of the larynx and, after laryngomalacia, is the second most frequent cause of congenital stridor.

Etiopathogenesis: Considered an incomplete form of subglottic atresia, this anomaly usually presents as a ring-shaped narrowing of the larynx approximately 2–3 mm below the glottic plane. A soft stenosis, composed of thickened fibrous tissue, is distinguished from a hard stenosis caused by malformation of the cricoid cartilage.

Symptoms: The inspiratory stridor of congenital subglottic stenosis is unaffected by position (“fixed stridor”) and is variable in its degree, depending on the size of the residual lumen. An upper respiratory infection can cause additional mucosal swelling that exacerbates the symptoms.

Diagnosis: Laryngoscopy demonstrates narrowing of the laryngeal lumen below the glottic plane.

Treatment: The first step after endoscopic confirmation is to establish a secure airway. Although congenital subglottic stenosis tends to resolve as the child grows, the classic wait-and-see approach with a temporary tracheotomy is being superseded more and more by a surgical procedure to enlarge the larynx, which can also be done in infants and small children.

Neurogenic Disorders:

Congenital Vocal Cord Paralysis

Etiopathogenesis: Congenital vocal cord paralysis caused by a lesion of the vagus nerve or its laryngeal branches may be unilateral or bilateral. The causes of vocal cord paralysis include diseases of the brain, myelomeningocele with or without hydrocephalus, Arnold–Chiari syndrome, obstetric head trauma, mediastinal tumors, anomalies of the heart and great vessels (Ortner syndrome), and anomalies of the lung and esophagus. Hereditary bilateral recurrent laryngeal nerve palsy has also been described. In many cases, however, a specific cause cannot be identified for the neural impairment of laryngeal function (idiopathic recurrent laryngeal nerve palsy).

Symptoms: A weak cry in a normally breathing infant is the hallmark of a unilateral recurrent nerve palsy that does not require treatment. The cardinal symptom of bilateral congenital vocal cord paralysis is respiratory distress with stridor and cyanosis. Swallowing may also be impaired due to loss of pharyngeal sensation (superior laryngeal nerve).
**Diagnosis and differential diagnosis:** Laryngoscopy reveals unilateral or bilateral vocal cord fixation. If both vocal cords are immobile, the differential diagnosis should include congenital ankylosis of the cricoarytenoid joints. The two conditions can be differentiated by electromyographic testing of the vocalis muscle.

**Treatment:** In cases with bilateral paralysis, the airway can be secured by tracheotomy or by early laterofixation of one vocal cord, depending on the symptoms and cause.

**Laryngeal Subglottic Hemangioma**

**Occurrence:** Cutaneous hemangiomas are a common finding and are often associated with similar lesions in other organs. Hemangiomas are occasionally observed in the subglottic space of newborns. Many cases become symptomatic during the first months of life as the hemangioma enlarges.

**Etiopathogenesis:** Hemangiomas are growing vascular lesions (ectasias) that belong to the category of hamartomas (= dysontogenetic growth of normally formed tissue).

**Symptoms:** When luminal narrowing is present, the symptoms may resemble those of congenital subglottic stenosis: respiratory distress and inspiratory stridor. Typically the stridor increases during crying due to increased engorgement of the lesion with blood. Hoarseness is not a typical feature.

**Treatment:** The fact that these vascular neoplasms may regress and disappear spontaneously during the first 2–4 years of life justifies an initial wait-and-see period, during which local and systemic corticosteroids may be administered. Depending on the size of the lesion, it may be possible to remove the hemangioma with the CO₂ laser and avoid tracheotomy. If the lesions do not regress spontaneously during the first years of life, they should be surgically removed.

**Malformations of the Trachea**

The symptoms of tracheal malformations that develop starting in weeks 4–6 of embryonic development are similar to the clinical manifestations of laryngeal anomalies: airway obstruction with **inspiratory and expiratory stridor**, wheezing, cyanosis, and gasping for breath. The voice is unchanged, however. Most tracheal malformations are accompanied by anomalies of the esophagus, the most common being a tracheoesophageal fistula. Isolated tracheal malformations are rare and usually involve the cervical part of the trachea. These very rare disorders are listed in Table 17.4.

<table>
<thead>
<tr>
<th>Table 17.4 Clinical classification of laryngotracheal malformations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laryngotracheal malformations can be classified as follows based on clinical criteria:</td>
</tr>
<tr>
<td>• Malformations that are (largely) incompatible with life:</td>
</tr>
<tr>
<td>– Aplasia</td>
</tr>
<tr>
<td>– Atresia</td>
</tr>
<tr>
<td>• Malformations that cause respiratory impairment (congenital stridor):</td>
</tr>
<tr>
<td>– Laryngomalacia, tracheomalacia</td>
</tr>
<tr>
<td>– Webs, stenosis</td>
</tr>
<tr>
<td>– Congenital recurrent laryngeal nerve palsy</td>
</tr>
<tr>
<td>– Laryngoceles, tracheoceles, cysts</td>
</tr>
<tr>
<td>– Subglottic hemangiomas</td>
</tr>
<tr>
<td>– Cartilaginous anomalies</td>
</tr>
<tr>
<td>• Malformations of the major vessels with tracheo-laryngeal compromise:</td>
</tr>
<tr>
<td>– Cleft anomalies</td>
</tr>
<tr>
<td>– Tracheoesophageal fistulas</td>
</tr>
<tr>
<td>• Malformations that have no clinical significance:</td>
</tr>
<tr>
<td>– Epiglottic malformations</td>
</tr>
</tbody>
</table>

Source: modified from Schultz-Coulon, see p. 411.

**Diagnosis:** As with laryngeal anomalies, endoscopic examination is of key importance. Certain clinical problems may additionally require radiographic contrast examination of the esophagus and/or bronchial system (e.g., for a suspected tracheoesophageal fistula) or angiography (for suspected vascular malformations).
17.4 Infectious Diseases of the Larynx and Trachea in Children

Airway infections are the most frequent cause of illness and death in infants and children. The airway dimensions in the pediatric upper respiratory tract are small, and even a slight obstruction can lead to critical luminal narrowing (see Fig. 17.4, p. 339). From 1 to 3 years of age, children come into increasing contact with a multitude of viruses and bacteria, and various portions of the upper and middle respiratory tracts may be affected. Initial challenges from these organisms often incite much more pronounced disease manifestations than subsequent infections that are acquired in later childhood.

Diseases Associated with Croup Symptoms

Croup syndrome refers to the inspiratory stridor that is caused by inflammatory laryngeal or subglottic stenosis. It is usually associated with respiratory distress, cough, and hoarseness. A croup syndrome may be caused by various diseases that have different prognostic implications. True croup is the term applied to specific laryngitis in the setting of diphtheria. Pseudocroup is a collective term for viral, bacterial, and spastic forms of subglottic laryngitis. These entities should be strictly distinguished from acute (usually bacterial) epiglottitis, although the latter is associated with croup-like symptoms. Table 17.5 reviews the most common diseases that may be associated with croup symptoms.

Diphtheria

“True croup” resulting from membrane formation and airway stenosis in diphtheria is very rarely seen today. Diphtheritic laryngitis with typical pseudomembranes occurs as an isolated condition in 25% of cases and is combined with nasopharyngeal diphtheria in 75% of cases. The major symptoms are hoarseness, a barking cough, and inspiratory stridor.

Mucosal swelling or the separation of pseudomembranes may incite an acute attack of respiratory distress that requires immediate intervention (airway maintenance, tracheotomy if required).

The disease is described more fully on p. 115.

Acute Subglottic Laryngitis

Synonym: acute laryngotracheobronchitis

Epidemiology: This viral disease, known also as pseudocroup, is by far the most common (90%) form of the croup syndrome. It occurs predominantly in infants and small children from 6 months to 3 years of age. Acute subglottic laryngitis has a seasonal peak incidence in the spring and fall, is frequently epidemic, and affects up to 5% of the population in this age group.

Etiopathogenesis: The disease develops gradually over 1–3 days during the course of an upper respiratory viral disease transmitted by droplet infection—parainfluenza I–III, influenza, respiratory syncytial (RS) viruses. Measles, varicella-zoster and rubella viruses can produce similar symptoms.

Symptoms: The voice is hoarse due to vocal cord involvement. Patients typically develop a dry, harsh, barking cough and stridor during the evening hours or at night after a few hours’ sleep. The stridor is loudest during inspiration and may progress to severe respiratory distress with cyanosis, depending on the degree of airway compromise. Generally, however, subglottic laryngitis runs a milder course than acute epiglottitis (see below).

Diagnosis: The body temperature is only moderately elevated or normal. Leukocytosis is usually absent. Laryngoscopy demonstrates an inflammatory swelling below the vocal cords and in the upper part of the cervical trachea. As the disease progresses, further luminal narrowing occurs due to crusting.

Treatment: Emphasis is placed on airway humidification and an adequate fluid intake. Generally, there is no need for antibiotics. The efficacy of rectally or orally administered steroids has not been definitely proven but may be beneficial in patients with increasing stridor. In cases with pronounced stridor and dyspnea, inhalation therapy with epinephrine derivatives will promote the regression of inflammatory swelling. Intubation is necessary only in exceptional cases. Most cases of acute subglottic laryngitis run a mild course, showing a complete resolution of symptoms and complaints in 3–5 days. Acute subglottic laryngitis may recur, however.
Table 17.5  Differential diagnosis of the most common infectious diseases associated with croup symptoms in infants and small children

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Epiglottitis</th>
<th>Subglottic laryngitis (pseudocroup)</th>
<th>Bacterial (laryngo)tracheitis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Causative organism</td>
<td><em>Haemophilus influenzae</em> b</td>
<td>Viruses</td>
<td>Viruses with bacterial superinfection</td>
</tr>
<tr>
<td>Age</td>
<td>2–8 years</td>
<td>6 months-3 years</td>
<td>Infants, small children</td>
</tr>
<tr>
<td>Onset</td>
<td>Sudden</td>
<td>Gradual</td>
<td>Gradual</td>
</tr>
<tr>
<td>Stridor</td>
<td>Inspiratory</td>
<td>Inspiratory and expiratory</td>
<td>Inspiratory and expiratory</td>
</tr>
<tr>
<td>Cough</td>
<td>–</td>
<td>Barking, dry</td>
<td>Productive</td>
</tr>
<tr>
<td>Voice</td>
<td>Muffled, soft, strained</td>
<td>Harsh, hoarse to aphonie</td>
<td></td>
</tr>
<tr>
<td>Swallowing</td>
<td>Difficult, painful</td>
<td>Unaffected</td>
<td>Usually difficult, painful</td>
</tr>
<tr>
<td>Dysphagia</td>
<td>+, drooling</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Fever</td>
<td>High</td>
<td>Usually subfebrile</td>
<td>Moderate</td>
</tr>
<tr>
<td>Leukocytosis</td>
<td>++</td>
<td>–</td>
<td>+</td>
</tr>
</tbody>
</table>

**Bacterial (Laryngo)tracheitis**

**Epidemiology:** This disease occurs sporadically and without seasonal incidence in infants and small children.

**Etiopathogenesis:** Bacterial (laryngo)tracheitis has a primary viral etiology (parainfluenza, influenza) with subsequent bacterial superinfection by staphylococci, pneumococci, streptococci, or *Haemophilus influenzae*.

**Symptoms and diagnosis:** The gradual onset with rhinitis and pharyngitis initially resembles the features of a mild subglottic laryngitis, but soon additional symptoms appear that do not occur in isolated laryngitis: expiratory and inspiratory stridor, numerous rales over the lungs, and other signs of a pulmonary complication (pneumonia, atelectasis).

Besides inflammatory airway obstruction, there is also a risk of lower airway obstruction by viscus mucus.

Stenotic symptoms develop relatively slowly. The temperature is moderately elevated, and a left shift is often seen in the blood count.

On clinical examination, mucosal redness is noted throughout the upper and lower respiratory tract. The vocal cords appear just as red and swollen as the subglottic and tracheal mucosae.

**Treatment and prognosis:** Antibiotic therapy should be supplemented by treatment with mucolytic agents (e.g., acetylcysteine, ambroxol), airway humidification, and adequate fluid intake. Intubation is indicated in severe cases for tracheobronchial toilet.

The prognosis is good in cases that receive prompt treatment.

**Acute Spasmodic Laryngitis**

**Synonym:** spasmodic croup

**Epidemiology:** Male infants and small children from 1 to 3 years of age are predominantly affected.

**Etiopathogenesis:** Acute spasmodic laryngitis is a rare disease who etiology is not yet fully understood. Some authors have proposed an allergic etiology analogous to bronchial hyperreactivity as well as gastroesophageal reflux. The symptoms are based on laryngeal spasms.

**Symptoms:** A child with no previous signs of infection wakes up at night in an extremely anxious state with coughing, stridor, and dyspnea. In contrast to bacterial laryngotraheitis, the symptoms subside completely within a few hours. Similar episodes may recur on subsequent nights.

**Treatment and prognosis:** The symptoms can be quickly relieved by administering ipecac syrup to induce vomiting. Additionally, the room air should be cool and humidified. Severe cases may require hospital observation. Intubation is rarely necessary. The disease, though recurrent, is self-limiting and has a favorable prognosis.
**Acute Epiglottitis**

**Epidemiology:** Epiglottitis occurs sporadically with no seasonal predilection and mainly affects children 2–8 years of age. The overall incidence has declined since the introduction of the *Haemophilus influenzae* type B vaccine.

**Etiopathogenesis:** Acute epiglottitis is a bacterial inflammation of the pharynx and laryngeal inlet. The most common pathogen is *Haemophilus influenzae* (type B). Other causative organisms are pneumococci and beta-hemolytic streptococci.

**Symptoms:** The disease has a swift onset marked by high fever, a loud inspiratory stridor, and severe respiratory distress. Hoarseness and cough may be absent. Most children complain of very painful swallowing. A muffled “hot potato” voice is a characteristic feature. Progression of the disease is marked by increasing inspiratory retraction in the jugular, sternal and intercostal areas. The dyspnea may progress rapidly due to increasing inflammatory obstruction of the laryngeal inlet, and death from asphyxia may occur within minutes to a few hours after onset.

The mortality rate is 5–10%.

**Diagnosis:** On examination of the oral cavity and pharynx, the posterior pharyngeal wall appears bright red and the epiglottis is swollen and erythematous (“cherry red epiglottis”). Abscess formation may occur (Fig. 17.17).

This inspection should be done with extreme care. Never press on the base of the tongue with a tongue blade, as this might reflexly evoke a laryngeal spasm causing total airway obstruction.

More than 50% of affected children develop bacteremia (blood culture). The blood count indicates leukocytosis with a left shift.

**Treatment:** Since intubation is usually unavoidable in children with acute epiglottitis, it should be performed as early as possible, but under controlled conditions (most deaths occur in prehospital settings). Given the short natural history of the disease, a tracheotomy is generally unnecessary. Besides sedation, treatment should include antibiotic therapy with adequate coverage for the spectrum of causative organisms (antibiotics of choice: third-generation cephalosporins). With prompt and adequate treatment, most patients can be extubated after the regression of inflammatory stenosis, usually 1–3 days after intubation. The recurrence of acute epiglottitis is very rare.

**Prophylaxis:** The best protection against this potentially life-threatening disease is the universal vaccination of infants and small children against *Haemophilus influenzae* after 2 months of age.

**Laryngeal Involvement by Infectious Diseases**

Inflammatory involvement of the larynx may occur in the setting of measles, scarlet fever, varicella, or pertussis. It is treated symptomatically.

**Syphilitic Laryngitis**

Involvement of the larynx by syphilis has become rare as a result of modern treatment. Although manifestations of syphilis are more common in the mouth and pharynx than in the larynx, laryngeal manifestations may be observed in all three stages of acquired syphilis.
17.5 Inflammatory Diseases of the Larynx and Trachea in Adults

Viruses, noxious agents, and extralaryngeal factors are the principal causes of inflammatory laryngotracheal diseases in adults. The cardinal symptom is hoarseness. Less common symptoms are stridor and respiratory distress.

Acute Laryngitis

Epidemiology and etiopathogenesis: Acute laryngitis often occurs in the setting of upper respiratory tract diseases that descend to involve the larynx. It mainly has a viral etiology, and bacterial superinfection may occur.

Symptoms: The hallmark of simple laryngitis is hoarseness, occasionally accompanied by a dry, nonproductive cough with no signs of airway obstruction. Dyspnea may occur in rare cases that have unusually severe mucosal swelling.

Diagnosis: Inspection of the larynx shows redness and possible thickening or edema of the vocal cords, which are coated with viscous mucus. Besides the glottis, the rest of the intralaryngeal mucosa may also be involved.

A severe form is fibrinous or interstitial laryngitis, marked by homogeneous redness of the vocal cords and possible whitish coatings due to fibrin exudation.

Treatment: The treatment of laryngitis consists of voice rest, inhalation therapy, mucolytic agents, and anti-inflammatory agents as required. Antibiotics are prescribed in patients with a concomitant bacterial infection. The systemic infection is an important factor to be considered in treatment planning.

Exogenous irritants (air pollutants, cigarette smoke, climatic influences) should be eliminated and avoided as much as possible.

Acute Epiglottitis

Synonym: adult supraglottitis

Acute epiglottitis is a bacterial inflammation, the main causative agents for which are Haemophilus influenzae, Streptococcus pneumoniae, and β-hemolytic streptococci. The clinical presentation resembles that described in 17.4 (see p. 356 and Fig. 17.17). Given the risk of airway obstruction, the examination should be performed carefully and with necessary emergency instruments on hand. The patient should be hospitalized for observation and treatment. Early initiation of treatment with intravenous antibiotics (first choice: third generation cephalosporins) and anti-inflammatory agents (corticosteroids systemically and by inhalation, epinephrine derivatives by inhalation) can frequently avoid the need for intubation.

Angioneurotic Laryngeal Edema, Acute Laryngeal Edema

Epidemiology: This rare, paroxysmal disease predominantly affects adolescents and adults. Occurrence in infants and small children is unusual.

Etiopathogenesis: Hereditary angioneurotic edema is caused by a congenital deficiency of C1 esterase inhibitor. An associated rise in esterase levels, often triggered by an infection or other disease process, leads to paroxysmal laryngeal edema.

Symptoms: Rapid edematous swelling of the larynx causes a typical inspiratory stridor. Similar edematous changes in the lip, tongue, palate, uvula, throat, or facial skin may accompany the laryngeal edema or may precede it by a variable period of time.

Diagnosis: Indirect laryngoscopy reveals edematous changes involving the entire larynx. Serologic testing can confirm the C1 esterase inhibitor deficiency. The differential diagnosis should include allergic reactions (e.g., insect bite, anaphylaxis, etc.) and angioedema induced by an angiotensin-converting enzyme (ACE) inhibitor.

Treatment: All forms of acute laryngeal swelling benefit from parenteral treatment with corticosteroids and antihistamines and the subcutaneous or local administration of epinephrine (by spray or inhalation). C1 esterase inhibitor products are available for the replacement therapy of C1 esterase inhibitor deficiency. Intubation is occasionally necessary in patients with increasing airway obstruction unresponsive to these measures.
17.3 Rare inflammatory diseases in adults

Laryngeal cellulitis and laryngeal abscess

**Etiopathogenesis:** Bacterial infections of the larynx may progress, leading to cellulitis or laryngeal abscess.

**Symptoms:** Laryngeal cellulitis and abscess usually have a sudden onset, often marked by initial chills. Cellulitis in particular is characterized by extremely severe odynophagia, which radiates to the ears. Dyspnea and hoarseness may develop, depending on the location and degree of swelling. Externally, the larynx and hyoid bone may be swollen and tender to pressure. The disease usually reaches its climax in about 3–5 days. Not infrequently, an abscess will point and rupture spontaneously and resolve quickly after draining. While an inflammation with abscess formation usually runs a harmless course, diffuse cellulitis poses a risk of descending infection with mediastinal involvement, inciting a life-threatening mediastinitis.

**Diagnosis:** Telescopic laryngoscopy shows an abscess as a bright, circumscribed swelling of the laryngeal mucosa with a central, yellowish zone of liquefaction. It is typically located on the epiglottic surface facing the tongue and less commonly within the larynx.

With *cellulitis*, the entire larynx appears markedly reddened. Cellulitic inflammation may not be clearly distinguishable from an abscess.

**Treatment** consists of administering a broad-spectrum antibiotic (third-generation cephalosporin) and securing the airway. Intubation may be unavoidable in some cases and particularly in patients with cellulitic laryngitis.

If an abscess is noted during laryngeal inspection, it should be surgically opened and drained under general anesthesia.

Laryngeal perichondritis

**Etiopathogenesis:** This inflammatory disease of the larynx has both a primary and a secondary form. Primary perichondritis begins in the perichondrium and spreads from there to the rest of the laryngeal skeleton. In former years, it occurred metastatically in acute infectious diseases such as typhus, spotted fever, and variola. This primary form of laryngeal perichondritis has become rare today.

In secondary perichondritis, the cartilage inflammation is preceded by a mucosal disease (e.g., laryngitis, laryngeal abscess, laryngeal cellulitis). It may also be caused by various kinds of trauma (external laryngeal injury, surgery) or by radiotherapy.

**Symptoms:** Besides respiratory distress, voice change, and odynophagia with referred earache, the affected portion of the larynx is usually tender to pressure. Acute perichondritis may be associated with marked systemic symptoms (chills, fever).

**Diagnosis:** Laryngeal perichondritis is easily diagnosed in cases where an external swelling or abscess has already formed. At first, the perichondritis is virtually indistinguishable from a simple edematous process by laryngoscopic examination. Suspicion is raised by more extensive infiltration with limited motion, ulceration, a protracted course, and severe subjective complaints. The formation of sequestered bone fragments confirms the diagnosis. These bony changes are consistently detectable by magnetic resonance imaging (MRI) and are occasionally visible on plain radiographs.

**Treatment:** Antibiotic treatment is of primary importance. Airway obstruction may necessitate a tracheotomy. If signs of liquefaction are seen, the abscess should be surgically drained and any sequestered bone fragments should be removed. Laryngectomy may be necessary in very rare cases. The resolution of perichondritis may leave the patient with significant laryngeal stenosis requiring surgical treatment. The incidence of this complication depends on the location of the inflammatory process in the laryngeal skeleton.

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**Fig. Causes of chronic laryngitis**

Vocal overuse, inhaled irritants (e.g., smoking)
Climate
Heat
Humidity
Dust

**Sinusitis**
**Rhinitis**
**Deviated septum**
**Tonsillitis**
**Gastroesophageal reflux**

**Chronic laryngitis**
**Tracheitis**
**Bronchitis**
**Bronchiectasis**

Chronic laryngitis has a multifactorial etiology and is often exacerbated by intercurrent viral and bacterial infections.

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**Chronic Nonspecific Laryngitis**

**Epidemiology:** Chronic laryngitis is most prevalent in smokers and in persons who abuse their voice. Most patients are men between 50 and 60 years of age.

**Etiopathogenesis:** The principal causes of chronic laryngitis are shown in Fig. 17.18. The epithelium responds to the irritants by thickening (hyperplasia and hyperkeratosis) and by developing submucous edema with inflammatory infiltrates and increased mucous glands (chronic hyperplastic laryngitis).

This condition is distinguished histologically from the less common chronic atrophic laryngitis (“laryngitis sicca”), which is characterized by a loss of laryngeal mucous glands.

**Symptoms:** Typical symptoms are hoarseness, rapid vocal fatigue, frequent throat clearing, and dry cough. The voice becomes low-pitched.
Diagnosis: Telescopic laryngoscopy yields nonspecific findings, often showing a reddened, hyperemic mucosa with a smooth surface. It is not uncommon to find a cobblestone appearance with leukoplakia and exophytic (whitish) keratosis.

Chronic laryngitis often cannot be positively distinguished clinically from an early stage of laryngeal carcinoma.

Treatment: The first priority is to identify and eliminate the causal irritants. With chronic laryngitis due to an ascending or descending inflammation (e.g., chronic sinusitis, bronchiectasis), an effort should be made to eradicate the primary focus (e.g., septoplasty, see p. 30, paranasal sinus surgery, see p. 55 and **3.16**, p. 57). Inhalation therapy and mucolytics are also beneficial. Acute exacerbations should be treated with antibiotics and, if necessary, with corticosteroid inhalant therapy.

The prognosis of chronic laryngitis depends largely on the ability to eliminate exposure to noxious exogenous agents.

Any intractable form of “laryngitis” requires histologic investigation and long-term follow-up.

Reinke’s Edema

Epidemiology: Men and women are affected equally, with a peak incidence between 40 and 60 years of age.

Etiopathogenesis: A subepithelial fluid collection forms between the glottic epithelium and the vocal ligament (Reinke’s space), presumably due to a local disturbance of lymphatic drainage from that space. The main etiologic factors are nicotine abuse and vocal abuse. Reinke’s edema is also viewed as a characteristic form of chronic hyperplastic laryngitis, although there is no reported tendency for Reinke’s edema to undergo malignant transformation.

Symptoms: The typical symptoms are hoarseness, frequent throat clearing, and a decrease in habitual voice pitch with rapid vocal fatigue. Dyspnea is not typically present.

Diagnosis: Laryngoscopy shows a glassy, edematous swelling at the level of the vocal cords (Fig. 17.19).

Treatment consists of microsurgical removal of the edema while preserving the vocal ligament, with circumscribed resection of the mucosa. Bilateral Reinke’s edema is operated in two stages (6 weeks apart) to prevent the formation of anterior synechiae. Surgical treatment is reinforced by having the patient quit smoking and by referring the patient for voice therapy as needed.

17.4 Fungal infections of the larynx and trachea

Involvement of the larynx and trachea by fungal diseases is rare. Most cases involve the secondary spread of infection from the oral cavity and pharynx. Primary fungal infections of the larynx and/or trachea are unusual. Predisposing factors are generally present and include wasting diseases, cachexia, vitamin deficiencies, congenital immune defects, acquired immunosuppression by treatment with corticosteroids or cytostatics, steroid inhalants for lung disease, antibiotic therapy, human immunodeficiency virus (HIV) infection, etc.

Candidiasis

This disease is caused by yeast fungi of the genus *Candida*. The most important representative, *Candida albicans*, is a normal intestinal saprophyte and becomes virulent only when predisposing host factors are present. In cases with extensive involvement of the oral mucosa, focal or membrane-like whitish plaques may also develop at the laryngeal inlet and especially on the epiglottis. A primary infection confined to the larynx is uncommon. Swallowing difficulties and hoarseness may provide subjective evidence of laryngeal involvement.

Treatment: Anti fungal agents may be applied topically to affected sites (with a brush under local anesthesia) or systemically (e.g., fluconazole).

Blastomycosis, aspergillosis, sporotrichosis

Laryngeal or tracheal involvement is rare. All of these fungal infections are marked by the development of superficial, granulomatous, ulcerating mucosal lesions. Differentiation from one another and from tuberculosis, syphilis, and neoplasia is occasionally difficult and requires identification of the causative organism. Treatment relies on antifungal agents specific for the infecting organism. Patients should also be screened for a predisposing underlying disease.

Posterior Laryngitis

Synonym: gastroesophageal reflux laryngitis (GERL)

Posterior or reflux laryngitis is a special form of chronic laryngitis attributed to the (reflux) reflux of gastric contents into the esophagus and pharynx. The cardinal symptoms are intractable hoarseness and a laryngopharyngeal foreign-body or globus sensation. As 24-hour esophageal pH measurements have shown, laryngeal acid reflux may occur in patients who have no other typical complaints of reflux disease (see textbooks of internal medicine). The disease, then, results from injury to the laryngeal and pharyngeal mucosa caused by chronic exposure to refluxed gastric acid. Given the anatomical relationship of the esophagus and larynx, the most severe damage occurs to the posterior portions of the larynx (arytenoid area) and to the postcricoid area. Inspection of these areas demonstrates redness, edema, and tissue proliferation, occasionally causing the interarytenoid area to have a “garden fence” appearance.
Thus, laryngitis in which the morphologic changes are predominantly posterior should always suggest a diagnosis of reflux laryngopathy. A gastroenterologic work-up (esophagogastroduodenoscopy, 24-hour pHmetry) is advised. Treatment follows established medical guidelines for reflux diseases. Proton-pump inhibitors are particularly effective.

**Laryngitis in Chronic Infectious Diseases**

**Tuberculous Laryngitis**

**Epidemiology:** As the incidence and prevalence of pulmonary tuberculosis have declined, tuberculous laryngitis has also become less common, although a certain rising trend has been documented in recent years.

**Etiology and pathogenesis:** Tuberculous changes in the larynx are almost always secondary to active pulmonary tuberculosis. Bacteria-laden secretions that are coughed up from the bronchi may infect the larynx, showing a special predilection for the posterior larynx, the interarytenoid area, and the laryngeal surface of the epiglottis. Laryngeal involvement usually parallels acute flare-ups of pulmonary tuberculosis. The extent and course of laryngeal tuberculosis, which may also be acquired hematogenously, depend on the efficacy of local host defenses and the virulence of the infecting organism. Laryngeal perichondritis and foci of liquefaction may develop as the disease progresses, and a secondary infection develops in most cases.

**Symptoms:** Tuberculous laryngitis has a variety of subjective symptoms ranging from mild, frequent throat clearing to severe hoarseness or aphony. Odynophagia signifies an advanced process with involvement of deeper tissues.
**Diagnosis:** Laryngoscopic inspection shows the characteristic appearance of tuberculous laryngitis: redness and thickening of one vocal cord (monochorditis), occasionally with small ulcerations.

Only histologic examination can differentiate tuberculous laryngitis from glottic carcinoma.

Reddish-brown, partially confluent submucous nodules are additionally found in the interarytenoid area and supraglottis.

**Treatment:** The mucosal changes will heal in response to tuberculous therapy, with no significant functional sequelae. Sites of cartilage destruction in the larynx will leave residual damage, but the prognosis is still favorable.

**Intubation Granuloma**

See 17.6, p. 365.

**Contact Ulcer**

**Epidemiology:** Men are predominantly affected. Contact ulcers are rare in children.

**Etiopathogenesis:** These lesions appear to be caused by chronic vocal abuse. Overuse of the voice leads to the repetitive, forcible adduction of the vocal processes of both arytenoid cartilages. The added presence of gastroesophageal reflux further contributes to the pathogenesis of this condition.

**Symptoms:** Typical complaints are hoarseness, a foreign-body sensation, and throat pain.

**Diagnosis:** Laryngoscopy initially shows a superficial mucosal lesion that later gives way to a contact granuloma with unilateral epithelial thickening and a rounded mucosal ulceration on the opposite vocal cord (“hammer-and-anvil” effect; Fig. 17.20). A phoniatric examination is indicated.

**Treatment:** Antireflux therapy and voice therapy are the cornerstones of treatment. Surgery (microlaryngoscopic ablation) is necessary only if conservative measures are unsuccessful. Contact granulomas have a strong tendency to recur.
17.6 Foreign-body Aspiration and Injuries to the Larynx and Trachea

Foreign-body aspiration occurs in children as well as adults. It is common for aspirated foreign bodies to enter the lower airways in children, whereas rapid glottic closure in adults is usually sufficient to prevent lower level impaction. For children under 6 years of age, aspirated foreign bodies are the leading cause of death in the home. External injuries to the larynx or trachea are rare occurrences, as this soft-tissue region is protected by the mandible, sternum, and spinal column. Also, the head is often flexed forward as a reflex response to external trauma.

Internal injuries to the larynx and trachea are more common and can have more serious ramifications. Internal injuries include intubation trauma and also the mucosal lesions and scarring that may be caused by many harmful agents. Laryngeal and tracheal injuries are almost always internal in children under 12 years of age; external injuries become more common with ageing.

Foreign-Body Aspiration

**Epidemiology:** Foreign-body aspiration is most prevalent in children under 3 years of age, with an approximately 2:1 preponderance of boys over girls.

**Etiology and pathogenesis:** Oropharyngeal swallowing abnormalities predispose to foreign-body aspiration. For anatomical reasons, aspirated foreign bodies are four times more often in the right main bronchus than in the left main bronchus. They are rarely found in the larynx and trachea.

Radiolucent foreign bodies are considerably more common than radiopaque objects.

Food items are aspirated with particular frequency, especially peanuts (Fig. 17.21; caution: may swell within the airway) and watermelon seeds. Tablets are common (laryngeal) foreign bodies in adults.

**Symptoms:** The initial symptoms of foreign-body aspiration depend on the size, shape, and composition of the foreign body, its location, and the age of the patient. Usually the object provokes an immediate coughing fit with or without cyanosis, accompanied by dyspnea, stridor, and pain. Larger foreign bodies impacted in the larynx may cause death from asphyxiation, whereas smaller objects lead to hoarseness and cough. The clinical picture of a complete obstruction is that of a cyanotic, aephonic patient with spasmodic breathing movements that do not ventilate the lungs. **Bolus death** refers to acute cardiac arrest caused by a vasovagal reflex evoked by obstruction of the upper airways. Foreign bodies in the trachea cause far greater complaints than objects in a bronchus, ranging from a slight cough to fatal asphyxia. If a foreign body in the trachea moves with respirations, it produces the tell-tale signs of a small palpable impact, an audible click, and movement of the trachea. A foreign body close to the vocal cords may be manifested by whistling sounds and stridor.

A foreign body lodged in the cervical esophagus may compromise the upper airway by compression, with associated symptoms. Complete obstruction of the esophagus also poses a risk of **overflow aspiration**.

**Diagnosis and treatment:** Besides the clinical examination with inspection and auscultation, radiographs are of key importance in determining the location of an aspirated foreign body. Chest radiographs at end-inspiration and end-expiration and lateral soft-tissue views of the neck are also helpful in locating nonradiopaque foreign bodies. If there is the least suspicion of an aspirated foreign body, generally there is no substitute for endoscopy of the upper aerodigestive tract for diagnostic and therapeutic purposes. Especially with soft, fragile foreign bodies, rigid endoscopy is definitely superior to flexible endoscopy for mobilizing and removing the foreign material. Special instruments are available for snaring and retrieving peanuts and other foreign bodies without breaking them into smaller fragments that could slip down into smaller airways and become lost in peripheral bronchi. Rigid endoscopy also provides the access needed to break sharp-edged or impacted objects into smaller pieces before definitive extraction. The rigid scope eliminates the risks of mucosal injury and perforation that are associated with a flexible endoscope.

With a foreign body completely obstructing the larynx, the Heimlich maneuver can be used to expel the object from the airway (see Fig. 17.22).

If the larynx is only partially obstructed, allowing some degree of respiration, the recommended treatment is endoscopic extraction of the foreign body under controlled conditions.
External Injuries to the Larynx and Extrathoracic Trachea

**Epidemiology and etiopathogenesis:** The pediatric larynx is less susceptible to external trauma because it is still very mobile, cartilaginous, and less prominent than in adults. Fractures of the thyroid and/or cricoid cartilages are almost unknown in children. The most frequent causes of laryngotracheal injury are deliberate attacks (strangulation, “karate chops”) and other anterior violence that compresses the tissue against the spinal column (e.g., handlebar impact in a bicycle accident, automobile dashboard injury). A life-threatening airway obstruction may occur immediately or with some delay after this kind of trauma.

Ruptures of the extrathoracic trachea are rare. The mechanism is an anterior force acting on the hyperextended neck, causing an outward bending of the posterior ends of the tracheal cartilages with rupture of the membranous posterior wall. The most serious and life-threatening form of blunt neck trauma is a complete avulsion of the larynx from the trachea.

Attempting intubation in this case could rupture the final connective-tissue attachments between the larynx and trachea, completely disrupting the airway and preventing further ventilation.

**Symptoms:** Signs indicating a laryngeal or tracheal injury are palpable discontinuities or crepitation in the larynx or tracheal tract, pain, a change in voice, swallowing difficulty, hemoptysis, cutaneous emphysema, and respiratory distress. An airway obstruction may be caused by edema, hemorrhage, injury to the vocal cord or recurrent laryngeal nerve, or mass effect from a tracheal hematoma. Tears in the mucosa and wall of the larynx and/or trachea lead to cutaneous emphysema of the head and neck area, with or without a pneumomediastinum and pneumothorax. A tracheal injury can also cause the creation of a tracheoesophageal fistula.

**Diagnosis and treatment:** Besides the clinical and endoscopic examination, computed tomography also supplies useful information on the location and extent of injuries.

The first priority in the management of external laryngotracheal injuries is to establish a secure airway.

If spontaneous respiration is satisfactory, it is unwise to perform overly aggressive endoscopic intubation in an emergency setting.

Whenever possible, endoscopy of the larynx and trachea should be performed under controlled conditions. Then an endotracheal tube can be introduced, with facilities available for performing a tracheostomy if needed.
Relatively minor injuries generally do not require further measures. Severe laryngeal and tracheal injuries should be treated operatively (e.g., cartilage repair) to prevent later stenosis. Avulsion injuries should be referred for reconstructive surgery without delay.

**Internal Injuries to the Larynx and Trachea**

Internal injuries to the larynx and trachea may be caused by thermal injuries (scalds and burns), chemical agents (caustic ingestion), or mechanical injuries (intubation, endoscopy).

**Thermal Injuries**

Thermal injuries to the laryngeal and tracheal mucosa may result from aspiration, burns due to hot foods or liquids, generalized burns, or surgical laser use.

**Chemical Agents**

Mucosal injuries by chemical agents most commonly affect the pharynx and supraglottic larynx. The sphincter function of the glottis usually protects the lower airways, although some 50% of pediatric victims will suffer respiratory tract injury due to aspiration, retching, or vomiting. While most lesions involve the supraglottic region, it is also possible for other portions of the endolarynx, the trachea, and the bronchial system to be affected.

**Etiopathogenesis:** Caustic injuries are caused mainly by household cleansers, which are typically ingested by children under 4 years of age. In adults, suicide attempts by acid or alkali ingestion are not altogether uncommon. These agents primarily damage the mucosal surface but may also cause deeper injuries. While acids tend to cause superficial coagulation necrosis, alcalis cause deeper colliquative necrosis, which has a poorer prognosis.

**Symptoms:** Besides acute airway obstruction due to mucosal swelling, patients are also threatened by the systemic effects of the absorbed chemical compounds.

**Diagnosis and treatment:** The main priority in acute cases is to establish a secure airway and provide systemic therapy for the caustic ingestion, which includes adequate parenteral fluid replacement.

An endoscopic survey should be done only during the first 24 hours due to the risk of perforation.

Primary management should also include immediate high doses of corticosteroids. The benefits of continuing this therapy at a maintenance dosage are uncertain. The causative agent should be identified and secured for therapeutic reasons (possible antidote) and to help in making a prognosis. Second-look endoscopy should be performed no earlier than 10–14 days after the injury. A broad-spectrum antibiotic should be administered due to the risk of bacterial infection. After the acute lesions have healed, there may be chronic residual changes in the laryngotracheal system marked by stricture formation and laryngeal dysfunction (incomplete glottic closure with recurrent aspiration). Corrective surgical treatments often prove difficult in these cases.

**Mechanical Injuries**

Internal mechanical injuries to the larynx and trachea are almost always a result of medical procedures such as intubation, endoscopy, and foreign-body extraction. Even suctioning the trachea with a catheter can lead to mucosal injuries. Acute injuries to the mucosal lining usually heal rapidly without treatment. The more deeply the wound extends toward the submucosa, the greater the tendency for granuloma formation (intubation granulomas, 17.6) and the greater the infection risk to cartilage and articular structures. Table 17.6 shows the Eckert scheme for grading the severity of intubation injuries. It is rare for intubation to cause the fracture or fracture-dislocation of an arytenoid cartilage.

**Effects of (Prolonged) Intubation**

The prolonged presence of an endotracheal tube can lead to chronic pathologic changes. There are various primary factors in prolonged intubation that can lead to permanent functional deficits such as stenosis, vocal dysfunction, and dysphagia.

**Etiology and pathogenesis** (Fig. 17.23): Besides the duration of intubation, a major pathogenic factor is a possible disproportion between the caliber of the endotracheal tube and the internal diameter of the cricoid cartilage. The cricoid cartilage encircles the narrowest point in the upper respiratory tract, and it has very little elastic compliance because of its annular geometry. Friction between the tube and larynx—which may occur in restless patients, for example, and especially when ventilator movements are transmitted to the tube—tends to erode through the extremely thin pad of submucosa lining the cricoid cartilage. The exposed cartilage is susceptible to infection, and the resulting perichondritis heals with the formation of granulations and scar tissue. This can result in subglottic stenosis, ankylosis of the arytenoid joints, and scar adhesions between the arytenoid cartilages causing vocal cord fixation in the midline.
17.6 Intubation granulomas

Oorotracheal or nasotracheal intubation may incite circumscribed inflammatory changes that in time can lead to the development of potentially large granulomas. The vocal process of the arytenoid cartilage is a site of predilection for intubation granulomas as a result of tissue damage from the particularly high tube pressure in the posterior glottis. Typically, intubation granulomas do not reach a symptomatic size until several weeks after extubation. Examination of the larynx for increasing hoarseness reveals a smooth, tense, pea-sized to cherry-sized polyoid mass with inflammatory redness (Fig.). Initially large intubation granulomas may regress spontaneously in 2–3 months. Microsurgical removal is indicated if the changes persist, but recurrences are common. Source of Fig.: Rusanowski, see p. 411.

Overinflating the tube cuff, or inflating it for too long a period, can also damage the mucosa. Intubation can also have late sequelae that are caused not only by local injury mechanisms but also by a weakening of host defenses that often accompanies the underlying disease and by intervals of hypotension with microcirculatory insufficiency. Patient age is also a significant factor in upper airway lesions following intubation. Because the cricoid cartilage is still soft and compliant in newborns, this is the age group in which intubation is best tolerated. While there have been isolated reports of subglottic stenosis occurring in infants after only one day of intubation, many other infants have been intubated for up to two months with no adverse effects. Generally, the risk of intubation-related complications is higher after orotracheal intubation than after nasotracheal intubation, because in the latter technique there is less frictional trauma to mucosal surfaces from the tube.

**Manifestation:** The chronic changes that can result from intubation (stenoses and intubation granulomas) often do not appear until 2–6 weeks after removal of the endotracheal tube.

**Treatment:** Generally, these changes, especially subglottic stenoses, can be corrected only by a major surgical procedure (17.7).

**Prophylaxis:** The prevention of these surgically challenging intubation injuries is based on selecting the proper tube size (17.8) and promptly recognizing the need for a tracheotomy where indicated. While tracheotomy should be considered in adult patients when the length of intubation is expected to exceed 3–6 days, infants and small children can tolerate 2–3 weeks of intubation owing to their compliant laryngeal anatomy. Tracheotomy is recommended in older children and adolescents when it is expected that intubation will exceed 1–2 weeks. Tracheotomy can prevent the aforementioned lesions that can develop in the larynx as a result of prolonged endotracheal intubation.

The rare complications of a correctly performed tracheotomy include postinflammatory cicatricial stenosis of the trachea above the stoma (suprastomal stenosis, Fig. 17.24), below the stoma (infrastomal stenosis), or at the level of the stoma (stomal stenosis). Tracheal stenosis can also result from mucosal lesions caused by cuff pressure from the tracheostomy tube (Fig. 17.24). This type of tracheal stenosis is rare, however, compared with the tracheal lesions caused by orotracheal or nasotracheal intubation.

Table 17.6 Classification of acute intubation injuries to the larynx

<table>
<thead>
<tr>
<th>Grade</th>
<th>Symptoms and findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Hyperemia, edema, discoloration of the mucosa.</td>
</tr>
<tr>
<td>II</td>
<td>Ulceration and necrosis of the mucosa and lamina propria.</td>
</tr>
<tr>
<td>III</td>
<td>Deep ulceration and necrosis extending to the cartilage.</td>
</tr>
</tbody>
</table>

Source: Eckerbom, see p. 411.

**Fig.** Causes of intubation injuries

- **Disease**
  - Length of intubation
  - Secondary infection

- **Physician**
  - Tube and tube location
  - Intubation process
  - Cuff pressure
  - Tracheal toilet
  - Sedation

- **Patient**
  - Laryngeal anatomy
  - Host defenses

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17.7 Treatment of subglottic and tracheal stenosis

For best results, the surgical treatment of subglottic and tracheal cicatricial stenosis should be deferred until the site is free of inflammation. Generally, there has been sufficient scarring to stabilize the site by 4–6 months after the causative event.

A subglottic stenosis (Fig. a) can be expanded simply by splitting the cricoid cartilage and interposing an anterior or posterior autologous (costal) cartilage implant. Increasingly, however, surgeons are favoring a partial resection of the cricoid cartilage with separation of the larynx and trachea and an end-to-end anastomosis of the cervical trachea to the thyroid cartilage (thyrotracheopexy, Fig. b). The inherent tension of the trachea requires that the neck be immobilized in an ante-flexed position for 1–4 weeks to relieve tension on the anastomosis.

When the operative site is exposed, care is taken to preserve the recurrent laryngeal nerves running between the trachea and esophagus. Ordinarily, a transverse resection cannot be performed if the segment to be resected is more than 5–6 cm long, and a different plastic reconstructive technique should be used (e.g., reconstructing the tracheal wall with cartilage grafts). A permanent tracheostomy or permanent intraluminal stent insertion may be unavoidable in some cases. A long-segment cicatricial stenosis should not be treated by repeated bougie dilation or by partial laser resection of the stenosis. These treatments are appropriate only for discrete webs.
17.8 Tube sizes recommended for endotracheal intubation and tracheostomy

Based on the relationships that have been described and the complications that may arise, the size of an endotracheal tube should be appropriate for the age of the patient. A good rule of thumb, especially in children, is that the tube diameter should not exceed the diameter of the patient’s small finger. The figures listed below are based on data from the anesthesiologic literature (source: Larsen, see p. 411). Usually, a somewhat larger ventilation tube can be used in a tracheostomy, because the trachea has a larger diameter in that region than at the subglottic level.

<table>
<thead>
<tr>
<th>Age group</th>
<th>Tube circumference</th>
<th>Inside diameter of tube</th>
</tr>
</thead>
<tbody>
<tr>
<td>Premature infants</td>
<td>Ch 10–12</td>
<td>2.5 mm</td>
</tr>
<tr>
<td>Newborns</td>
<td>Ch 12–14</td>
<td>3 mm</td>
</tr>
<tr>
<td>1–6 months</td>
<td>Ch 16</td>
<td>3.5 mm</td>
</tr>
<tr>
<td>6–12 months</td>
<td>Ch 18</td>
<td>4.0 mm</td>
</tr>
<tr>
<td>1–2 years</td>
<td>Ch 18–20</td>
<td>4.0–4.5 mm</td>
</tr>
<tr>
<td>2–3 years</td>
<td>Ch 20–22</td>
<td>4.5–5.0 mm</td>
</tr>
<tr>
<td>3–4 years</td>
<td>Ch 22–24</td>
<td>5.0–5.5 mm</td>
</tr>
<tr>
<td>4–5 years</td>
<td>Ch 24–26</td>
<td>5.5–6.0 mm</td>
</tr>
<tr>
<td>5–7 years</td>
<td>Ch 26–28</td>
<td>6.0–6.5 mm</td>
</tr>
<tr>
<td>7–9 years</td>
<td>Ch 28</td>
<td>6.5 mm</td>
</tr>
<tr>
<td>10–11 years</td>
<td>Ch 28–30</td>
<td>6.5–7.0 mm</td>
</tr>
<tr>
<td>12–13 years</td>
<td>Ch 32</td>
<td>7.5 mm</td>
</tr>
<tr>
<td>14–16 years</td>
<td>Ch 34</td>
<td>8.0 mm</td>
</tr>
<tr>
<td>Adults (women)</td>
<td>Ch 30–34</td>
<td>7.0–8 mm</td>
</tr>
<tr>
<td>Adults (men)</td>
<td>Ch 34–36</td>
<td>8.0–9 mm</td>
</tr>
</tbody>
</table>

17.9 Tracheomalacia

Cicatricial stenosis of the trachea requires differentiation from tracheomalacia, in which the tracheal walls are flaccid and tend to collapse causing functional stenosis. This condition is attributed to cartilage damage due, for example, to a pressure-induced deficiency of blood flow (e.g., goiter). Treatment options are tracheostomy (sutting the trachea to adjacent structures) or intraluminal stent insertion.
17.7 Tumors of the Larynx and Trachea

The cardinal symptom of all neoplasms of the larynx, whether benign or malignant, is persistent hoarseness. Any hoarseness that lasts longer than two weeks should be investigated by laryngoscopy. Making risk patients (smokers) more aware of hoarseness as a potential warning sign can aid in the early detection and more effective treatment of glottic carcinoma, which is the most common malignant tumor of the head and neck.

Benign Neoplasms of the Larynx

Benign tumors constitute the majority of laryngeal neoplasms in both children and adults. They generally present clinically as a mechanical obstruction of the upper airways with coughing, hoarseness, wheezing, and dyspnea.

Vocal Cord Polyps

**Epidemiology:** Adults in speaking professions are mainly affected, with a preponderance of males.

**Etiopathogenesis:** The most frequent cause is a mechanical alteration of the vocal cords caused by vocal overuse (phonotrauma) and chronic inflammation. Histologic examination reveals a polypoid mucosal hyperplasia with an inflammatory component. Most vocal polyps are unilateral (90%) and are located on the free edge of the anterior two-thirds of the vocal cord.

**Symptoms:** The cardinal symptom is hoarseness. Floating polyps may cause diplophonia.

**Diagnosis:** Telescopc laryngoscopy demonstrates a grayish-red sessile or pedunculated mass on the vocal cord (Fig. 17.25).

**Treatment:** Treatment consists of microsurgical removal, which may be followed by voice therapy.

Cysts and Mucoceles

**Epidemiology:** These lesions are most common in older patients and are rarely encountered in children.

**Etiopathogenesis:** Cystic lesions originate in the small mucosal glands of the laryngeal mucosa and form in the area of the ventricular fold, sinus of Morgagni, or subepithelial vocal fold. They are usually lined by squamous or columnar epithelium. Mucus-filled retention mucoceles and extravasation mucoceles may occur anywhere in the larynx and trachea where mucous glands are present. They are lined by respiratory epithelium.

**Symptoms:** Clinical manifestations depend on the size and location of the lesions and may consist of hoarseness, globus sensation, and rarely dyspnea.

**Diagnosis:** Telescopc laryngoscopy demonstrates smooth, epithelium-covered masses of varying size.

**Treatment:** The recommended treatment is removal by endolaryngeal microsurgery.

Papillomas and Laryngeal Papillomatosis

**Epidemiology:** Papillomas are the most common benign laryngeal tumors in children. Juvenile papillomas are most prevalent between the second and fourth years of life. While papillomas in adults (second to fourth decades) are usually solitary, juvenile laryngeal papillomatosis is characterized by multiple lesions that spread to the trachea and bronchial system. Multiple lesions may also be found in adults, however.

**Etiopathogenesis:** Histologically, the papillomas are neoplasms and not a reaction to a chronic inflammatory stimulus. The causal agents are human papillomaviruses (HPVs), most notably HPV 6 and HPV 11. Malignant transformation is rare in juvenile papillomatosis but is more common in the adult form.

**Symptoms:** The initial symptoms are hoarseness and an inspiratory stridor that develops with increasing obstruction. Occasional aggressive growth may cause life-threatening luminal obstruction of the larynx or trachea.

**Diagnosis:** The typical endoscopic appearance is that of multiple soft, reddish-pink, villous, raspberry-like lesions covering a large area of the glottis and supraglot-
tis (Fig. 17.26). Generally, the lesions first appear on
the vocal cords.

Regular histologic follow-ups are essential due to
the potential for malignant transformation.

Treatment: The treatment of choice is CO₂ laser sur-
gery, which can remove the frequently recurring le-
sions with very little bleeding. A potential complica-
tion of repeated excisions is glottic webbing. Other
treatment modalities (beta interferon, virostatics,
photodynamic therapy) have been used with varying
success.

Prognosis: The papillomas may resolve spontaneously
in rare cases, or may recur after an asymptomatic in-
terval.

Vocal Nodules

Epidemiology: Vocal nodules can occur in children
(screamer’s nodules), singers (singer’s nodules), and
in patients who speak professionally, especially
younger women.

Etiopathogenesis: The underlying cause is harmful vo-
cal habits (vocal abuse) causing bilateral nodules to
form at opposing sites at the junction of the anterior
and middle thirds of the vocal cords (maximum vibra-
tional amplitude; Fig. 17.27 and Fig. 18.8, p.393). His-
tologic examination reveals fibrosis with epithelial
thickening and submucosal connective-tissue prolif-
eration.

Symptoms: Typical symptoms are hoarseness, diplo-
phonia, habitual throat clearing, and a foreign-body
sensation.

Treatment: The treatment of choice is voice therapy.
Surgical removal is indicated only in patients with ex-
ceptionally large nodules and for whom voice therapy
has failed. Vocal nodules in puberty generally regress
spontaneously due to longitudinal growth of the vocal
cords.
Malignant Laryngeal Tumors

Laryngeal Carcinoma

Epidemiology: Malignant tumors of the larynx are the most common head and neck malignancies, accounting for approximately 40% of these cancers and for 1–2% of all malignant tumors. The reported incidence in the United States is 4–6 per 100,000 population per year. Internationally, incidence rates range from 2.5–17.1 per 100,000 population per year in men and from 0.1–1.3 per 100,000 population per year in women.

Etiopathogenesis: Besides the epithelial changes described in 17.10, chronic laryngitis must be considered a predisposing factor for laryngeal carcinoma based on the presence of the same risk factors.

Histology, sites of occurrence and classification: The great majority of laryngeal malignancies are keratinized or nonkeratinized squamous cell carcinomas (90–95%). The rest consist of undifferentiated carcinomas, well-differentiated verrucous carcinomas, and other rare entities. Sixty percent of laryngeal carcinomas are located in the glottic plane, 40% in the supraglottic region, and only about 1% in the subglottis. A carcinoma of the glottis, Morgagni pouch, and ventricular fold that has an indeterminate site of origin is called a transglottic carcinoma.

The growth of supraglottic malignancies into the pre-epiglottic fat pad, extension to the glottic plane, the spread of glottic malignancies to the anterior commissure with possible thyroid cartilage invasion, and paraglottic extension down along the thyroid cartilage not only worsen the prognosis but also limit the possibilities for larynx-conserving surgery. Vocal cord fixation by invasion of the muscles inserting on the arytenoid cartilage or by infiltration of the criocarytenoid joint signifies the very advanced growth of a glottic or subglottic cancer and critically affects management. Like all head and neck tumors, laryngeal carcinoma is...
17.10 Carcinogenesis (continuation)

The different line thickness of the black arrows represent the likelihood of regression of the dysplastic epithelial changes or the development of invasive carcinoma. The red arrows represent inevitable, irreversible processes.

- a Normal epithelium
- b Hyperplasia
- c Hyperkeratosis
- d Dysplasia
- e Carcinoma in situ
- f Invasive carcinoma
17.11 Premalignant Lesions

Premalignant (precancerous) lesions are epithelial changes that may give rise to carcinoma. The gross clinical appearance of potentially premalignant lesions in the larynx is highly variable. The degree of dysplasia is a histologically defined criterion that cannot be assessed clinically.

- **Leukoplakia**: whitish patch of mucosa that cannot be rubbed away; may be circumscribed or more diffuse. Gross appearance does not permit reliable benign-malignant differentiation.
- **Erythroplakia**: reddish, nonkeratinized epithelial lesion with a strong likelihood of malignant transformation. In many cases, carcinoma in situ is already present.
- **Pachydermia**: area of epithelial thickening, more or less completely covered by keratin scales.

In all of these lesions, the degree of dysplasia should be evaluated histologically based on the World Health Organization staging system for epithelial dysplasia (see also Fig. in 17.10):

- **Hyperplasia**
- **(Hyper)keratosis**
- **Dysplasia (3 grades: mild, moderate, severe)**
- **Carcinoma in situ**

In principle, squamous cell carcinoma of the larynx can develop from all histologic grades of epithelial dysplasia (see Fig. in 17.10). In a smaller percentage of laryngeal cancers, especially at the supraglottic level, normal epithelium may also undergo a direct malignant transformation. These epithelial changes may be asymptomatic but often cause complaints in the form of hoarseness, dry cough, or a foreign-body sensation. The lesions are nine times more common in men than in women.

The changes visible by indirect laryngoscopy most commonly affect the vocal cords but may involve any portion of the larynx. Microscopic evaluation is indicated for tissue sampling and histologic evaluation and may involve complete removal of the affected mucosa. In cases with vocal cord involvement, the histologic degree of dysplasia will determine whether it is possible to remove the epithelium of the vocal folds while leaving the vocal ligament intact. All cases require regular follow-ups and particularly the elimination of causal agents (cessation of smoking).

Staged according to international UICC guidelines (Table 17.7). Glottic malignancies have a considerably better prognosis than supraglottic or subglottic cancers, not only because of earlier diagnosis but also because of their limited lymphatic drainage (see pp. 342–343). Up to 60% of supraglottic and subglottic malignancies already have ipsilateral lymph-node metastases at the time of diagnosis, and up to 30% have bilateral or contralateral regional nodal metastases, which imply a very poor prognosis. By contrast, less than 10% of tumors confined to the glottic plane are accompanied by regional metastases when diagnosed.

Distant metastases, which most often spread hematogenously to the lung and then require differentiation from a second primary (bronchial carcinoma), are unusual with laryngeal malignancies, and asymptomatic patients require no screening tests for distant metastases other than a chest radiograph. As with all other tumors of the upper aerodigestive tract, the simultaneous or metachronous occurrence of second tumors in the upper foodway and airway is of major significance in patients with laryngeal cancer. The UICC staging system shown in Table 17.8 is based on local tumor extent (T category), regional lymph-node involvement (N category, see Table 16.5, p. 334), and distant metastasis (M category).

**Symptoms**: The symptoms of laryngeal malignancies—foreign-body sensation, habitual throat clearing, dysphagia, respiratory distress, hemoptysis—depend on the location and extent of the tumor. Glottic malignancies, even when small, are likely to cause voice change (hoarseness) as their initial symptom, enabling them to be diagnosed at an earlier stage than supraglottic malignancies, which often remain silent for some time. The cardinal symptoms of subglottic cancers are dyspnea and inspiratory stridor.

Hoarseness that persists longer than 2–3 weeks should be investigated by laryngoscopy.

Earache in the absence of otoscopic abnormalities, known as referred otalgia, may signify tumor-related irritation of the vagus nerve in the larynx and hypopharynx.

**Diagnosis**: Suspicious cases are initially evaluated by indirect laryngoscopy, giving particular attention to vocal cord mobility. The examiner should also palpate the laryngeal skeleton (for extralaryngeal tumor ex-
### Table 17.7  UIICC classification of laryngeal carcinoma

<table>
<thead>
<tr>
<th>T</th>
<th>Primary Tumor</th>
</tr>
</thead>
<tbody>
<tr>
<td>TX</td>
<td>Primary tumor cannot be assessed</td>
</tr>
<tr>
<td>T0</td>
<td>No evidence of primary tumor</td>
</tr>
<tr>
<td>Tis</td>
<td>Carcinoma in situ</td>
</tr>
</tbody>
</table>

#### Supraglottis

| T1  | Tumor limited to one subsite of supraglottis with normal vocal cord mobility   |
| T2  | Tumor invades more than one adjacent subsite of supraglottis or glottis or region outside the supraglottis (e.g., mucosa of the base of the tongue, vallecula, medial wall of the piriform sinus), without fixation of the larynx |
| T3  | Tumor limited to larynx with vocal cord fixation and/or invasion of postcricoid area, preepiglottic space, paraglottic space and/or with minor thyroid cartilage erosion (e.g., inner cortex) |
| T4a | Tumor invades through the thyroid cartilage and/or invades soft tissues beyond the larynx, e.g., trachea, soft tissues of the neck including deep/extrinsic muscle of tongue (genioglossus, hyoglossus, palatoglossus, and styloglossus), strap muscles, thyroid, esophagus |
| T4b | Tumor invades prevertebral space, mediastinal structures, or encases carotid artery |

#### Glottis

| T1  | Tumor limited to vocal cord(s) may involve anterior or posterior commissure with normal mobility |
| T1a | Tumor limited to one vocal cord |
| T1b | Tumor involves both vocal cords |
| T2  | Tumor extends to supraglottis or subglottis or with impaired vocal cord mobility |
| T3  | Tumor limited to the larynx with vocal cord fixation and/or invades paraglottic space, and/or with minor thyroid cartilage erosion (e.g., inner cortex) |
| T4a | Tumor invades through the thyroid cartilage and/or invades soft tissues beyond the larynx, e.g., trachea, soft tissues of the neck including deep/extrinsic muscle of tongue (genioglossus, hyoglossus, palatoglossus, and styloglossus), strap muscles, thyroid, esophagus |
| T4b | Tumor invades prevertebral space, mediastinal structures, or encases carotid artery |

#### Subglottis

| T1  | Tumor limited to the subglottis |
| T2  | Tumor extends to vocal cord(s) with normal or impaired mobility |
| T3  | Tumor limited to the larynx with vocal cord fixation |
| T4a | Tumor invades through the thyroid cartilage and/or invades soft tissues beyond the larynx, e.g., trachea, soft tissues of the neck including deep/extrinsic muscle of tongue (genioglossus, hyoglossus, palatoglossus, and styloglossus), strap muscles, thyroid, esophagus |
| T4b | Tumor invades prevertebral space, mediastinal structures, or encases carotid artery |

### Table 17.7  Continuation

<table>
<thead>
<tr>
<th>N</th>
<th>Regional lymph nodes</th>
</tr>
</thead>
<tbody>
<tr>
<td>NX</td>
<td>Regional lymph nodes cannot be assessed</td>
</tr>
<tr>
<td>N0</td>
<td>No regional lymph node metastasis</td>
</tr>
<tr>
<td>N1</td>
<td>Metastasis in a single ipsilateral lymph node, less than 3 cm</td>
</tr>
<tr>
<td>N2</td>
<td>Metastasis in a single ipsilateral lymph node between 3 and 6 cm; or in multiple ipsilateral lymph nodes, none more than 6 cm in greatest dimension; or in bilateral or contralateral lymph nodes, none more than 6 cm in greatest dimension</td>
</tr>
<tr>
<td>N2a</td>
<td>Metastasis in a single ipsilateral lymph node between 3 and 6 cm</td>
</tr>
<tr>
<td>N2b</td>
<td>Metastasis in multiple ipsilateral lymph nodes, none more than 6 cm in greatest dimension</td>
</tr>
<tr>
<td>N2c</td>
<td>Metastasis in bilateral or contralateral lymph nodes, none more than 6 cm in greatest dimension</td>
</tr>
<tr>
<td>N3</td>
<td>Metastasis in a lymph node greater than 6 cm</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>M</th>
<th>Distant metastasis</th>
</tr>
</thead>
<tbody>
<tr>
<td>MX</td>
<td>Distant metastasis cannot be assessed</td>
</tr>
<tr>
<td>M0</td>
<td>No distant metastasis</td>
</tr>
<tr>
<td>M1</td>
<td>Distant metastasis</td>
</tr>
</tbody>
</table>

### Table 17.8  TNM categories of laryngeal carcinoma

<table>
<thead>
<tr>
<th>Category</th>
<th>T</th>
<th>N</th>
<th>M</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Tis</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>I</td>
<td>T1</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>II</td>
<td>T2</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>III</td>
<td>T3</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>IV A</td>
<td>T4</td>
<td>N0–1</td>
<td>M0</td>
</tr>
<tr>
<td>IV B</td>
<td>T4</td>
<td>N0–3</td>
<td>M1</td>
</tr>
</tbody>
</table>

### Tension and Soft Tissues

Tension and the soft tissues of the neck (for lymph-node metastases). Ultrasonography of the cervical soft tissues is an essential part of the basic work-up in order to detect possible regional metastasis. Computed tomography and magnetic resonance imaging can be helpful in verifying tumor extent. Microlaryngoscopy (Fig. 17.28) is used to define the precise tumor extent and to obtain histologic tissue samples that can differentiate cancer from other laryngeal pathology such as chronic laryngitis or benign endolaryngeal neoplasms. This procedure should include panendoscopy (inspection of the nasopharynx, oropharynx, hypopharynx, tracheobronchial passages, and esophagus) to exclude second tumors.
Fig. Laryngeal carcinomas

a Clottic carcinoma involving the anterior third of the left vocal cord (T1a).
b Clottic-supraglottic carcinoma involving the left side of the larynx (T3; arrows).
c Supraglottic carcinoma (arrows) arising from the epiglottis. Note the edematous swelling of the arytenoid region on the affected side (*).

**Treatment:** The main therapeutic options are surgery and radiotherapy; the role of chemotherapy is still uncertain. Surgery remains the primary treatment of choice, however.

A larynx-sparing and voice-sparing partial resection can be done in a large percentage of patients, depending on the tumor location and extent. This is a particularly good option with circumscribed tumors.

With more extensive tumors (T3, T4), which may broadly infiltrate adjacent structures, complete removal of the larynx (laryngectomy) is occasionally unavoidable for oncologic reasons.

Generally a neck dissection is unnecessary with small glottic carcinomas (Tis, T1), but it is indicated for higher tumor categories and especially for supraglottic and subglottic cancers. The need for postoperative radiotherapy will depend on the location and extent of the tumor and on lymphogenous metastasis. **17.12** illustrates several partial laryngectomy techniques in current use.

Early glottic cancers are sometimes treated with radiotherapy alone, the major advantage being superior voice quality after treatment. Disadvantages are the necessary length of treatment (approximately 6

### 17.12 Surgical treatment options for laryngeal carcinoma

#### Glottic carcinoma

**Voice-sparing procedures**

- Vocal cord stripping (Fig. a):
  - **Principle:** removal of the vocal cord epithelium.
  - **Approach:** endolaryngeal by microlaryngoscopy.
  - **Indication:** epithelial dysplasia, carcinoma in situ.

- Partial or complete cordectomy (Fig. b):
  - **Principle:** removal of the tumor-involved portion of the vocal cord.
  - **Approach:** endolaryngeal by microlaryngoscopy with CO₂ laser resection or external access via thyrotomy (incision of thyroid cartilage).
  - **Indication:** T1a laryngeal carcinoma.

- More extensive partial laryngectomies (Fig. c):
  - **Principle:** supraglottic, glottic or subglottic tissue resection for bilateral and advanced glottic malignancies, may include resection of cartilaginous structures from the thyroid and/or cricoid cartilage; can be extended to a hemilaryngectomy
  - **Approach:** external via thyrotomy (incision or resection of thyroid cartilage) or endolaryngeal by microlaryngoscopy with CO₂ laser resection

- **Indication:** T1b or T2 glottic carcinoma; can be extended under certain conditions to remove more extensive tumors

#### Non-voice-sparing procedure

- Total laryngectomy (Fig. d):
  - **Principle:** complete removal of the larynx with separation of the airway and foodway and construction of a permanent tracheostomy.
  - **Approach:** external.
  - **Indication:** advanced laryngeal tumors that cannot be adequately removed by other procedures in accordance with oncologic principles.

#### Supraglottic laryngeal carcinoma

**Voice-sparing procedure**

- Horizontal partial laryngectomy (Fig. e):
  - **Principle:** removal of the supraglottic larynx with preservation of the glottic plane and arytenoid cartilages.
  - **Approach:** external via horizontal partial laryngectomy or endolaryngeal by microlaryngoscopy with CO₂ laser resection.
  - **Indication:** T1 and T2 supraglottic malignancies; may be suitable for more extensive tumors.

**Non-voice-sparing procedure**

- Laryngectomy (Fig. d):
  - **Principle and approach:** see above
  - **Indication:** advanced laryngeal tumors that cannot be adequately removed by other procedures in accordance with oncologic principles

Source of Fig.: Naumann et al., see p. 411.

*continuation next page*
weeks), a higher complication rate in surgery for recurrent or residual disease, and the risk of radiation-induced malignancy. In patients with advanced tumors, radiotherapy alone has so far been used only as a palliative treatment. Modern fractionation schemes for radiotherapy, combined with the use of cytostatics, may open up new possibilities in the future. Today the highest cure rates for laryngeal malignancies are achieved with surgery, which may be followed by radiotherapy in patients with advanced tumors and especially metastases.

**Prognosis:** The prognosis depends on the location and stage of the disease. While T1 glottic carcinomas have a 5-year survival rate of up to 100%, the rates associated with advanced glottic cancer (T4) decline to approximately 50%. Supraglottic carcinoma has a considerably poorer prognosis even in T1 cases, which have a 5-year survival rate of 70–80%. The 5-year survival with advanced tumors is 30–40%.
Fig. Speech valve prosthesis

Principle of a speech valve prosthesis for voice rehabilitation after laryngectomy, illustrated here for a Provox valve.

Functional Sequelae of Surgery for Laryngeal Malignancies

Permanent voice change is a very common sequel to partial laryngectomy, depending on the resection technique. But a much more serious problem for patients is the swallowing difficulty that can result from impairment of the laryngeal sphincter function following a very extensive partial laryngectomy. This can lead to chronic aspiration with recurrent bouts of pneumonia and may ultimately require separating the respiratory and digestive tracts by performing a total laryngectomy.

Swallowing function is generally not impaired after a total laryngectomy, but laryngeal voice production is lost.

Laryngectomy Effects and Voice Rehabilitation

Since a total laryngectomy completely separates the respiratory tract from the digestive tract, laryngectomized patients have respiratory anosmia due to the absence of nasal airflow but still have intact gustatory and olfaction. They can no longer sneeze, blow their nose, or perform a Valsalva maneuver. The nose and pharynx can no longer condition the inspired air, and so the air inhaled through the tracheostomy is neither warmed nor humidified, especially at certain times of the year. This can lead to irritation of the tracheal mucosa (trachitis) with crusting. Shields can be worn to protect the tracheostomy while bathing or showering.

Various options are available for voice rehabilitation after laryngectomy, all of which require intensive speech training by a therapist. It is always helpful to teach the patient esophageal speech, which is produced by swallowing air into the esophagus and forcing it back up against folds of mucosa in the upper esophagus and hypopharynx.

Another method of voice restoration is to create a tracheoesophageal fistula through which air can be forced into the upper part of the esophagus, with mucosal folds in the pharyngoesophageal segment functioning as a speech generator. The tracheoesophageal fistula may be surgically constructed (“neoglottis,” with risk of aspiration during swallowing), or a valve prosthesis may be placed between the trachea and esophagus (Fig. 17.29). Another option is to use electronic devices for speech production. In this method an external sound generator transmits vibrations to the pharyngeal wall and oral floor, setting the air in the resonant chambers (pharynx, mouth, nose) into vibration and producing speech. The role of these devices is limited, however, by the relatively poor quality and mechanical nature of the sound.

It is extremely helpful to refer laryngectomized patients to self-help groups (e.g., the International Association of Laryngectomees) once they have been made aware of the options for voice rehabilitation after laryngectomy.

Oncologic Follow-Up

Regular oncologic follow-up visits are scheduled chiefly for the purpose of detecting a local or regional tumor recurrence while it is still asymptomatic. The high incidence of metachronous second tumors in the upper aerodigestive tract is also a major concern, requiring that all risk-exposed mucosal areas be examined during each follow-up. The standard follow-up examination includes a specific history, a complete otolaryngologic examination, and palpation and ultrasound scanning of the cervical soft tissues. If the findings raise the suspicion of a tumor, or if all mucosal areas cannot be reliably evaluated, it may be necessary to proceed with panendoscopy under general anesthesia. Chest radiographs should also be obtained at regular intervals.

In scheduling the follow-up intervals, it should be kept in mind that the great majority of locoregional recurrences are diagnosed within 2 years after the primary treatment.

Tracheal Tumors

Chondroma, Osteochondroma, Osteoma

These cartilaginous and bony neoplasms may be manifested in the trachea and the main bronchi. They appear endoscopically as a thickening of tracheal or bronchial cartilages and are surrounded by a capsule. Endoscopic biopsy is often unsuccessful due to the hardness of the tumors. They grow very slowly but have the potential to cause extensive bronchopulmo-
nary destruction. Sarcomatous transformation has been described. Treatment consists of complete surgical removal.

**Tracheopathia osteochondroplastica**

This condition is based on a malformation of the tracheal and bronchial cartilages. Abnormal deposits of cartilaginous tissue in the endotracheal mucosa during embryonic development present postnatally as numerous small bony or cartilaginous tumors that project into the tracheal lumen and may cause progressive airway obstruction. There is no causal treatment for this disease, which is manifested by wheezing, coughing, hemoptysis, and increasing respiratory distress. Progressive airway obstruction is relieved by debulking or removing the lesions that cause the greatest degree of luminal narrowing.

**Malignant Tumors of the Trachea**

Primary malignancies of the trachea are very rare. The most common type is adenoid cystic carcinoma (Fig. 17.30). It is much more common, however, for malignant tumors to invade the trachea from adjacent structures (larynx, hypopharynx, esophagus, thyroid gland). Surgical treatment is the best option whenever it can be done with curative intent. Otherwise radiotherapy and/or chemotherapy should be considered, depending on the tumor histology. An essential palliative measure is airway maintenance. This can be done by repeated (laser) tumor debulking, stent insertion, or tracheotomy.

![Adenoid cystic carcinoma of the trachea](image)

**Fig.** Adenoid cystic carcinoma of the trachea

- **a** MRI appearance
- **b** Resection specimen

- **a** Arrows indicate the carcinoma.
- **b** Surgical specimen following resection and end-to-end anastomosis.

Source: Kramann, see p. 411.
17.8 Airway Management

The principal methods of airway management—establishing an airway, stenting, preventing blood aspiration, ventilation—are intubation (see textbooks of anesthesiology and emergency medicine), cricothyrotomy, and tracheotomy. Special care measures are needed in tracheotomized patients.

Tracheotomy and Cricothyrotomy

Tracheotomy refers to an incision of the trachea below the larynx. In the technique that is now most favored, the isthmus of the thyroid gland is transected and an opening is made between the second and third tracheal rings.

Elective Tracheotomy

An elective tracheotomy (indications Table 17.9) is performed under controlled surgical conditions. The opening in the cartilage should not exceed a critical size; otherwise subsequent closure of the tracheotomy could lead to tracheal stenosis due to the loss of cartilaginous substance. It is particularly important in children to avoid resecting any cartilaginous structures, as this could lead to refractory postoperative complications.

The anterior wall of the trachea is sutured to the skin of the neck to create an epithelialized tract (Fig. 17.31). Unlike a nonepithelialized tracheostomy, this type of tracheotomy eliminates the danger of false passage during cannula changes and avoids the risk of a descending pretracheal inflammation.

Emergency Tracheotomy

Otrotracheal or nasotracheal intubation is the usual primary method of airway intervention in patients with acute respiratory distress caused by an upper airway obstruction between the dental arch and larynx. Today this method is rapidly available, largely standardized, and has a low complication rate (see textbooks of anesthesiology). Occasionally, however, intubation is unsuccessful even when done under operative conditions and endoscopic control (fiberoptic intubation), and it may be necessary to open the trachea in order to establish a secure airway. When the head is fully extended and neck anatomy is undistorted, the cervical trachea is palpable and visible in the midline beneath the skin. In a highly acute emergency situation, incision of the trachea may be hampered by subcutaneous tissue layers and especially by bleeding from the thyroid isthmus.

---

Table 17.9 Indications for elective tracheotomy

- Laryngeal stenosis caused by:
  - Tumors
  - Swelling (e.g., postirradiation)
  - Bilateral vocal cord paralysis
  - Subglottic stenosis
- Tracheal stenosis above the proposed stoma site
- Prolonged mechanical ventilation (see 17.6, p. 365)
- Pulmonary diseases (to facilitate bronchial toilet and reduce dead space)
- Postoperative airway management following upper respiratory tract surgery

In children, very strict criteria should be applied in assessing the need for tracheotomy.

Fig. Principle of elective tracheotomy

a Incision of the anterior tracheal wall.
b Schematic cross section of an epithelialized tracheostomy.
Cricothyrotomy

Synonym: coniotomy

Cricothyrotomy, in which the airway is opened through the cricothyroid ligament between the thyroid cartilage and cricoid cartilage, is still an important procedure in emergency situations (Fig. 17.32). This area of the larynx is subcutaneous, is easily palpated, and can be opened with a transverse incision. The instrument used to make the luminal incision (e.g., pocketknife) should be held perpendicular to the skin and should not be withdrawn until the tube has been introduced between the thyroid and cricoid cartilages; otherwise the tissue planes would shift and block access to the lumen. It is very helpful to include a ready-to-use cricothyrotomy set in every emergency kit (Fig. 17.32). A cricothyrotomy should be converted to a standard tracheotomy as soon as possible due to the danger of cricoid cartilage injury with intralaryngeal stenosis.

Percutaneous Tracheotomy

In this technique, a needle is inserted between the tracheal rings, a guidewire is introduced, the tract is dilated, and finally the tracheotomy tube is introduced over the guidewire into a palpable area of the trachea. Percutaneous tracheotomy is most commonly used in emergency medicine and to establish a temporary tracheotomy in intensive-care settings. Commercial sets are available for percutaneous tracheotomy, which requires a familiarity with cervical anatomy.

The Tracheotomized Patient

Tracheotomized patients with an intact larynx can speak by plugging the tracheotomy tube with a finger or cap. Voice quality will depend on the extent of pathology in the laryngeal region. A speaking tube with a valve suspended in the inspiratory/expiratory air stream will also allow for phonation. Fig. 17.33 shows an assortment of commonly used tracheostomy tubes. Tracheotomized patients and their families should be taught the specifics of tracheostomy care before leaving the hospital. This includes accessories for tracheotomized patients such as a suction device, an inhalation and/or room-humidifier unit, dressings, and tube care materials. The selection of the tube material (plastic, silver, cuffed or noncuffed) depends on the underlying disease, the necessary duration of the tracheostomy, and local findings. Respiratory distress in tracheotomized patients usually signifies obstruction of the trachea due to crusting. Initial emergency measures are decannulation, tracheoscopy, and removal of the crusts as needed.

In a temporary tracheotomy, the tube is withdrawn at the end of the tracheotomy period and the stoma is closed with an airtight adhesive seal to make certain that the patient can breath adequately on his own. This tentative closure, combined with removal of the tube, will cause the stoma to shrink and facilitate surgical closure. Especially with an epithelialized tracheostoma, plastic surgical technique should be used in making the definitive closure.
17.9 Neurogenic Disorders of the Larynx

Paralysis of the vocal cord muscles supplied by the recurrent laryngeal nerves often has an iatrogenic cause (e.g., thyroid gland surgery) as well as unknown idiopathic causes. The cardinal symptom of unilateral vocal cord paralysis is hoarseness, whereas bilateral paralysis is manifested by dyspnea, which often necessitates emergency intervention. Isolated paralysis of the superior laryngeal nerve and vagus nerve is relatively rare.

Recurrent Laryngeal Nerve Paralysis

It is necessary to distinguish between unilateral and bilateral recurrent laryngeal nerve paralysis due to their different cardinal symptoms and treatment strategies.

Unilateral Recurrent Laryngeal Nerve Paralysis

Epidemiology and etiopathogenesis: The typical causes of recurrent laryngeal nerve paralysis are listed in Table 17.10. The most frequent cause is a lesion sustained during thyroid surgery, with reported lesion rates of 0.14–5.0% in initial operations and up to 20% in revision procedures. These figures pertain to unilateral recurrent laryngeal nerve lesions detected immediately after surgery. With lesions that do not disrupt the continuity of the nerve, it is reasonable to expect that functional recovery will occur within weeks or months in a certain percentage of cases. Intraoperative exposure of the recurrent laryngeal nerves during thyroid surgery lowers the lesion rate.

"Idiopathic" vocal cord paralysis is diagnosed by exclusion upon the completion of comprehensive diagnostic tests (see below). This condition is more common in men than women, has a peak incidence at 20–30 years of age, and affects the left side more often than the right side. It is assumed to have a viral etiology, as it is often preceded by an upper respiratory infection.

Symptoms: The chief complaint of unilateral recurrent laryngeal nerve paralysis is hoarseness. Respiratory distress is generally not observed.

Diagnosis: At telescopic laryngoscopy, unilateral vocal cord immobility is noted during respiration. Usually the vocal cord is fixed in a paramedian position (Fig. 17.34 and Fig. 17.35), but the vocal cord position alone is not a reliable indicator of lesion location. If a cause cannot be identified, a diagnosis of "idiopathic" paralysis should not be made until further tests have been performed. These include examination of the thyroid gland and neck (ultrasonography), the mediastinum (computed tomography or magnetic resonance imaging), serologic tests, and a possible history of exposure to harmful agents.

Electromyography (EMG), which may employ needle electrodes inserted transorally or transcervically into the vocal cords under local or general anesthesia without muscle relaxation, supplies information useful in assessing the cause, extent, and prognosis of the lesion. EMG is also helpful in differentiating a recurrent laryngeal nerve lesion from the rare conditions of arytenoid dislocation, which is usually caused by intubation.

<table>
<thead>
<tr>
<th>Table 17.10 Causes of recurrent laryngeal nerve paralysis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Unilateral</strong></td>
</tr>
<tr>
<td>• Iatrogenic nerve injury, especially in thyroid operations (most frequent cause, usually affecting the left side) and surgery of the esophagus, trachea, mediastinum, lung, or heart. Variations in the course of the nerve (e.g., a &quot;nonrecurrent&quot; recurrent nerve) are a predisposing factor</td>
</tr>
<tr>
<td>• Infiltration or compression of the nerve by tumors of the thyroid gland, larynx, trachea, or mediastinum</td>
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<tr>
<td>• Laryngeal trauma</td>
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<tr>
<td>• Nerve compression by mediastinal masses (known as Otter syndrome, e.g., cardiac hypertrophy, aortic aneurysm, malignant lymphoma, sarcoidosis)</td>
</tr>
<tr>
<td>• Infectious or toxic neuritis (e.g., influenza viruses, herpes viruses, alcohol, lead, arsenic, vinca alkaloids)</td>
</tr>
<tr>
<td>• (Poly)neuritis in the setting of a rheumatoid or (auto)immune disease</td>
</tr>
<tr>
<td>• Infectious diseases: Lyme disease, syphilis, mononucleosis</td>
</tr>
<tr>
<td>• Diabetic (poly)neuropathy</td>
</tr>
<tr>
<td>• Idiopathic (approximately 20% of cases; viral and other etiologies have been proposed)</td>
</tr>
<tr>
<td><strong>Bilateral</strong></td>
</tr>
<tr>
<td>• Preexisting unilateral recurrent laryngeal nerve paralysis, with one of the above causes supervening</td>
</tr>
<tr>
<td>• Both recurrent laryngeal nerves affected by one of the above causes</td>
</tr>
<tr>
<td>• Iatrogenic bilateral nerve injury (usually in thyroid operations, especially malignant tumor resections and resections of large or recurrent goiters; also surgery of the trachea and esophagus)</td>
</tr>
<tr>
<td>• Simultaneous bilateral recurrent nerve paralysis caused by infectious or toxic agents is relatively rare but may occur</td>
</tr>
</tbody>
</table>
trauma, and arytenoid fixation by scar tissue (detectable muscle activity).

An examination of the larynx to assess vocal cord mobility is essential before and after thyroid surgery (strumectomy, thyroidectomy).

**Treatment:** The treatment of unilateral recurrent laryngeal nerve paralysis depends chiefly on its cause. If the paralysis occurs immediately after a surgical procedure in the territory of the nerve, it should first be determined whether the nerve is transected or whether it still has continuity and may have been functionally compromised due to pressure (hemotoma, nerve caught in a stitch). Because the recurrent laryngeal nerve supplies nerve fibers to both agonistic and antagonistic muscles (abductors and adductors), there is little hope of success in reapproximating a completely severed nerve, even if the ends can be found, due to the erratic nature of regeneration by axonal sprouting. But if continuity of the nerve is preserved, an immediate revision procedure can be done to decompress the nerve and promote regeneration. Whenever possible, causal therapy should be attempted first for nontraumatic nerve lesions, depending on the established or presumed cause. An unsevered nerve can regenerate, and it is very likely that recovery will occur within 6–12 months. It is rare for spontaneous recovery to occur after 1 year.

Long-term unilateral recurrent laryngeal nerve paralysis leads to muscular atrophy with excavaation of the vocal cord and persistent vocal dysfunction. Voice therapy may help to reduce hoarseness in patients with persistent unilateral vocal cord paralysis. Other options are augmentation of the paralyzed vocal cord by the injection of Teflon or autologous collagen and a medialization procedure to improve vocal cord position (see 18.2, p. 394).

**Bilateral Recurrent Laryngeal Nerve Paralysis**

**Etiology:** See Table 17.10.

**Symptoms:** The chief complaint is dyspnea—ranging in severity from resting dyspnea to dyspnea on exertion—because the paralyzed vocal cords assume an almost closed position due to the relative predominance of the adductor muscles over the abductors (see Table 17.1, p. 341). The severity of the dyspnea depends on the residual glottic gap. Additional mucosal swelling due to intubation or infection can exacerbate the respiratory distress.

**Diagnosis:** Telescopic laryngoscopy during respiration usually shows the vocal cords fixed in a paramedian position and displaying only passive motion in response to transglottic airflow. The diagnostic methods described under unilateral recurrent laryngeal nerve paralysis should be applied as needed.

**Treatment:** When bilateral vocal cord paralysis is noted immediately after a surgical procedure (e.g., strumectomy), it should first be determined whether it is advisable to perform an immediate revision procedure with neurolysis of one or both nerves. If the nerves have not definitely been severed, this type of surgery is generally indicated.

An immediate tracheotomy is unavoidable in the great majority of cases. The next step is to wait and see if one or both nerves regenerate. A prognosis can be offered based on the EMG findings.

Surgical procedures to widen the glottis should be considered no earlier than 6 months after the onset of paralysis.
Laterofixation of a vocal cord

In cases of persistent bilateral recurrent laryngeal nerve paralysis, the glottic aperture can be sufficiently widened for respiration by lateralizing one of the vocal cords. The drawings illustrate a technique in which two narrow mucosal triangles are resected in the false vocal cord and over the arytenoid cartilage, which is also removed. Source: Naumann, see p. 411.

| Table 17.11 Symptoms of unilateral paralysis of the superior laryngeal nerve |
|---------------------------------|-----------------------------|
| Low-frequency vocal range       |                             |
| Deeper, monopitched speaking voice |                        |
| Inability to sing higher notes  |                             |
| Breathy voice that fatigues easily |                     |
| Decrease in maximum phonation time |                     |

Source: Modified from Berendes, see p. 411.

| Table 17.12 Causes of vagus nerve paralysis |
|-------------------------------------------|-----------------------------|
| Intramedullary (see also textbooks of neurology): |             |
| Congenital: Arnold–Chiari syndrome, Dandy–Walker syndrome, syringobulbia, myelomeningocele, Klippel–Feil syndrome, aplasia of vagal nuclei (Gerhardt syndrome) | |
| Inflammatory: poliomyelitis, herpes zoster, Guillain–Barré syndrome, diphtheria, tabes dorsalis | |
| Vascular: thrombosis, hemorrhage, angioma, Wallenberg syndrome | |
| Tumors of the brainstem and floor of the fourth ventricle | |
| Disseminated encephalomyelitis | |
| Extramedullary: | |
| Tumors in the area of the jugular foramen (e.g., paragangliomas) | |
| Basal skull fractures | |
| Vagus neurinomas (cardinal symptom: coughing fits triggered by palpation of the mass) | |
| Trauma: stab wounds and gunshot injuries, iatrogenic lesion caused by local anesthesia, neck dissection, revascularizing procedures on the carotid artery | |

Laterofixation always aims for the best tradeoff between voice quality and respiratory function for the individual patient. The surgical procedure should also be tailored to the specific patient (age, desired voice quality).

**Superior Laryngeal Nerve Paralysis**

Paralysis of the superior laryngeal nerve may be iatrogenic following thyroid surgery (usually affects only the motor component) or laryngeal surgery (also affects the sensory component). It is less common than recurrent laryngeal nerve paralysis and usually does not produce motor symptoms. Only singers and professional speakers will notice the (unilateral) loss of innervation to the cricothyroid muscle (Table 17.11). The vocal cord on the affected side appears flaccid at laryngoscopy because the posterior commissure is deviated toward the side of the paralysis, causing a slight length discrepancy between the vocal cords. Voice therapy may be appropriate in some cases. A more serious concern is the impairment of laryngeal sensation that occurs in cases with internal branch involvement. Bilateral lesions are particularly likely to cause dysphagia with aspiration, and care should be taken to avoid these lesions during laryngeal surgery. If these symptoms fail to improve with swallowing exercises, laryngectomy may be considered as a final recourse on functional grounds.

With persistent bilateral recurrent laryngeal nerve paralysis, the goal of further treatment is to widen the glottic gap and eliminate the need for tracheotomy. This can be done either by suturing portions of one vocal cord in a lateraled position (laterofixation, (Fig. 17.36) or by partially resecting one vocal cord and/or an arytenoid cartilage. This may be done inter-
Vagus Nerve Paralysis

Etiology and pathogenesis: Vagus nerve lesions are much less common than recurrent laryngeal nerve paralysis. Possible causes are listed in Table 17.12.

Symptoms: The symptoms basically represent a combination of the symptoms associated with recurrent laryngeal nerve paralysis and superior laryngeal nerve paralysis. In the rare cases with bilateral vagus nerve lesions, the dominant feature is dyspnea. The chief complaint in unilateral paralysis is hoarseness.

Diagnosis: Telescopie laryngoscopy again shows an immobile vocal cord, which is usually described as being fixed in an intermediate position. This may be true in the early phase, but as other factors supervene (e.g., atrophy of the denervated muscles) the vocal cord position is not a reliable site-of-lesion indicator. Depending on the presumed location of the lesion, the patient should be referred for a neurologic workup that includes cranial imaging and/or an examination of the neck (ultrasound, computed tomography, magnetic resonance imaging).

Treatment: Treatment is tailored to the established or presumed etiology. The treatment options described for recurrent laryngeal nerve paralysis are also appropriate for vagus nerve paralysis.

17.13 Superior laryngeal neuralgia

Shooting, throbbing pain lasting seconds to hours or a constant ache lasting several hours or days in the side of the neck may be caused by superior laryngeal neuralgia (Awellis syndrome). The point of maximum pain intensity is usually at the site where the internal branch enters the thyrohyoid membrane. The pain may radiate to the ear, tongue, and throat. Differentiation is mainly required from dentogenic causes, sialolithiasis, and glossopharyngeal neuralgia (which may coexist with superior laryngeal neuralgia). The treatment of first choice is local infiltration of the nerve with a long-acting local anesthetic. Carbamazepine has also proved beneficial. Another option is surgical division of the nerve (for functional sequelae, see p. 382).
Voice Disorders

18.1 Clinical Voice Physiology and Diagnostic Procedures
Basic Principles of Speech Production
Changes in the Voice with Ageing
Diagnostic Procedures

18.2 Clinical Aspects of Voice Disorders
Organic Dysphonia
Functional Dysphonia
Mutational Voice Disorders
Endocrine Dysphonia
Vocal Cord Paralysis
18.1 Clinical Voice Physiology and Diagnostic Procedures

Verbal communication is the essential foundation for interpersonal contacts and for human culture in general. It is inextricably linked to the integrity of receptive language function (hearing and understanding) and of expressive language abilities. In this textbook of otolaryngology, we are concerned mainly with the laryngologic aspects of communication disorders.

Basic Principles of Speech Production

Physiology

The most important anatomical structures for speech production are the vocal cords (vocal folds), which form the lateral boundaries of the glottis (see also 17.1, Anatomy of the Larynx, pp. 338–345).

The primary voice signal is produced in the larynx by the vibrations of the vocal cords and is sustained by a dynamic equilibrium between the expiratory air pressure and the muscular tone of the vocal cords. Vocal cord vibration consists of a complex three-dimensional motion:

• The basic motion is produced by the predominantly mediolateral vibration of the vocalis muscle and vocal ligament (Fig. 18.1a).

• The mobility of the superficial mucosa relative to the vocalis muscle during phonation generates a surface wave in the epithelium. This “traveling wave motion” is superimposed over the transverse vibrations of the vocalis muscle (Fig. 18.1b). Normal traveling wave motion is a prerequisite for what is subjectively perceived as “good” voice quality. As the voice becomes louder, the glottis remains closed for a relatively longer time during the vibratory cycle, which maintains a constant period (Fig. 18.2).

Based on a simple concept called the “source-filter model,” the primary voice signal is modulated by the resonant cavities of the pharynx, mouth, and nose (the “vocal tract”) and is emitted from the mouth as a complex voice sound (Fig. 18.3). In an ordinary speaker, no interaction takes place between the vocal cord vibrations and the vocal tract. Trained singers or professional speakers learn how to manipulate these interactions to enhance the quality and timbre of their voices.

Pathophysiology

Changes in the elasticity of the connective tissue, the tone of the vocalis muscles, and the properties of the epithelium can significantly limit and alter the normal traveling wave motion of the vocal folds. This change is clearly audible, even to laypersons, as a voice disorder.

If the vocal cords snap together too forcefully due to pathology, the vibratory cycle becomes jerky and the voice sounds harsh and grating. If the cords adduct too weakly or are unable to meet, the voice sounds weak and breathy and its loudness cannot be increased. In this case, the vocal cord vibrations can become temporarily desynchronized, causing the right and left sides to vibrate at different fundamental frequencies.

Changes in the Voice with Aging

When an infant cries for the first time, the larynx begins to function as an organ of phonation. The voice of infants and small children has a very high fundamental frequency (400 Hz), which decreases to approximately 300 Hz as the larynx continues to grow. Laryngeal growth undergoes a sex-specific response to hormonal changes during puberty. As a result, the fundamental frequency of the voice in girls decreases by a third to a fifth to 220–250 Hz, while in boys it falls by more than an octave to 110–140 Hz (Fig. 18.4). This physiologic change or “breaking” of the voice is termed mutation.

In adults as well, the phonatory function of the larynx is influenced by hormonal changes (e.g., during pregnancy and menopause). By about 70 years of age, the voice acquires a similar pitch in males and females (“senile voice”) and loses much of its variety of expression.

Diagnostic Procedures

All diagnostic evaluation of the voice is based on an endoscopic examination of the larynx. Other basic tests involve the recording, measurement, and evaluation of voice quality and performance. Optional tests such as electromyography (EMG) and respiratory function may also be added.
Glottic width

a The basic vibratory pattern of the vocal cords is a mediolateral motion. 
b A traveling wave motion of the mucosa is essential for a “normal” sounding voice.

Endoscopic Examination

Today, the larynx is always examined with the aid of endoscopic illumination and magnification (rigid telescopic laryngoscope with a 90° or 70° viewing angle; see also 17.2, pp. 346–349). Vocal cord vibrations at a fundamental frequency of 100–400 Hz cannot be observed with an ordinary light source, but they can be viewed by stroboscopy (Fig. 18.5). In this technique, the vibrating vocal cords are illuminated with a stroboscope at a rate that is matched to the fundamental frequency of the voice.

An experienced examiner can evaluate the muscular tone and organic symmetry of the vocal cords based on the stroboscopic examination. The findings can be recorded on video tape for analytical playback and to help patients understand the nature of their condition.

Perceptual Evaluation of the Voice

Despite the availability of quantitative test procedures, the quality of a voice is still evaluated mainly by subjective auditory perception. The following dimensions of voice quality are evaluated in the RBH system:

- Roughness
- Breathiness
- Hoarseness
These properties are described more fully in Table 18.1 and, with a little practice, are easy to distinguish from one another. They are graded on a four-point scale:

- 0 = no disorder
- 1 = mild disorder
- 2 = moderate disorder
- 3 = severe disorder

It should be noted that in the RBH system, roughness and breathiness are rated separately from each other, while hoarseness is considered an “overall measure” of voice quality. Thus, the value of H is always equal to the greater of the R and B values. **Hoarseness** is the end result of a voice disorder. Because of this, the quality and degree of hoarseness depend little on the specific cause—i.e., various disorders can produce the same hoarseness. Conversely, a particular disease in a patient can cause different types of hoarseness that may vary at different times of day, for example.

Rating voice quality and identifying the cause of a voice disorder are inherently ambiguous, therefore.

The standard clinical workup can be supplemented by the **voice range profile**, in which the subject’s pitch and loudness ranges are plotted against each other in a two-dimensional graph called a phonetogram. Connecting the upper and lower points in the graph outlines the area within which normal phonation can occur (Fig. 18.6). The larger the enclosed area, the greater the “capacity” of the voice.

This test requires a cooperative patient who can sing musical notes, and so it is subject to uncontrolled influences.

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### Table 18.1 Dimensions of voice quality in the RBH system: definition and source

<table>
<thead>
<tr>
<th>Sound impression</th>
<th>Acoustic definition</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Roughness</td>
<td>Noise components caused by aperiodicities in the fundamental vibration of the voice sound, with superimposed components</td>
<td>Irregularities of vocal cord vibrations, additional sound sources</td>
</tr>
<tr>
<td>Breathiness</td>
<td>Noise components caused by turbulence of unmodulated expiratory airflow</td>
<td>Incomplete glottic closure</td>
</tr>
<tr>
<td>Hoarseness</td>
<td>Noise components in speech</td>
<td>All deviations from the normal vibratory pattern of the vocal cords, additional sound sources</td>
</tr>
</tbody>
</table>

Source: Wendler, see p. 411.
The voice mutates differently in girls and boys during puberty. Mutation disorders (see p. 393) can result in an abnormally high-pitched or low-pitched voice.

**Fig. 1** Principle of stroboscopy

- **a** Flash rate \( f_b \) = vocal cord frequency \( f_s \)
  - Flash rate \( f_b \)
  - Vocal cord frequency \( f_s \)
  - Steady image

- **b** Flash rate \( f_b \) < vocal cord frequency \( f_s \)
  - Flash rate \( f_b \)
  - Vocal cord frequency \( f_s \)
  - Visible vibration (frequency = \( f_s - f_b \))

**Fig. 2** Phonotogram
18.2 Clinical Aspects of Voice Disorders

A basic distinction is drawn between organic and functional voice disorders (dysphonias). The organic correlates of voice disorders involving the larynx are covered in 17.3–17.7 (pp. 350–377) and 17.9 (pp. 380–383). The present deals with the functional aspects of these disorders.

Organic Dysphonia

Classification: Organic voice disorders are distinguished from functional voice disorders and are classified under the following etiologic headings (Table 18.2):
- Malformations (“dysplastic dysphonia”)
- Trauma
- Inflammation
- Tumors
- Functional disorders

Laryngeal malformations are very rare and may be manifested by the cardinal symptoms of dyspnea, dysphonia, and dysphagia. The main functional problem in newborns with laryngeal anomalies is dyspnea (see p. 350), while dysphonia and dysphagia become more important with ageing. The main functional priority in children is airway maintenance. The main goal in teenagers and adults is to compensate for the disorder or adapt to the functional deficit. Generally, this is accomplished by functional training and in rare cases by surgical intervention.

The functional sequelae of laryngeal trauma depend on the extent of the trauma. They can range from hoarseness to aphony, from respiratory distress to apnea, from odynophagia to dysphagia, and from coughing to hemoptysis. The degree of residual dysfunction depends on the extent of the primary injury. Treatment in the acute stage focuses on airway maintenance and if necessary may include the surgical reconstruction of structures that have been destroyed. In patients who have residual dysfunction, the only option is symptomatic therapy aimed at promoting compensatory mechanisms. Surgical procedures such as adhesiolysis are of very limited value.

Voice disorders in acute laryngitis generally resolve with regression of the organic lesion. From a functional standpoint, it is important to maintain voice rest during the inflammatory stage in order to prevent maladaptive compensation. Patients in certain occupations may require a 7–10-day leave of absence. If hoarseness persists after acute laryngitis has subsided, the patient should be referred for voice therapy to keep the hoarseness from becoming chronic. Twenty sessions are generally sufficient, preferably in an outpatient setting (inpatient therapy is rarely necessary). In chronic laryngitis, only a few selected patients will require voice therapy to optimize compensatory mechanisms.

The hoarseness associated with a benign laryngeal neoplasm will generally improve after treatment of the neoplasm. In exceptional cases the voice disorder may persist in an attenuated form due to maladaptive compensation, necessitating voice therapy. Postoperative voice therapy may also be tried in patients with residual dysfunction, which is relatively common after the surgical removal of extensive Reinke’s edema. This therapy has only a modest success rate, however, in terms of compensating for the dysfunction.

The surgical resection of malignant laryngeal neoplasms is almost invariably followed by dysphonia. From a phoniatric standpoint, voice therapy should be initiated as soon after the surgery as possible since a more extensive resection does not necessarily imply a poorer prognosis in terms of vocal function. Surgical techniques of glottic reconstruction after a unilateral vocal cord removal, for example, have only a very limited prospect of success.

Functional Dysphonia

Definition: Functional dysphonia is a category of voice disorders in which there is no obvious organic change affecting the phonatory structures.

Etiopathogenesis: Functional voice disorders usually have a multifactorial etiology (Fig. 18.7). Constitutional, habitual, stress-related, and psychogenic causal factors have been identified. It is difficult in any given case to determine the relative contribution of specific factors to the overall picture. Usually multiple factors can be identified that may reinforce one another under certain conditions. A vicious cycle may perpetuate a chronic voice disorder, which may later give rise to morphologic changes (vocal nodules, known also as screamer’s or singer’s nodules; see below and Fig. 18.8).

It is customary to subdivide functional dysphonias into hyperfunctional and hypofunctional disorders.

Symptoms: These disorders may be associated with a general impairment of all aspects of voice function.
### Table 18.2 Differential diagnosis of voice disorders

<table>
<thead>
<tr>
<th>Cause</th>
<th>Special features</th>
<th>For description see:</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Malformations</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Internal and external</td>
<td>Can have a very broad range of symptoms, depending on size</td>
<td>17.2, p. 350</td>
</tr>
<tr>
<td>laryngoceles</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hemangiomas</td>
<td></td>
<td>p. 353</td>
</tr>
<tr>
<td>Laryngomalacia</td>
<td></td>
<td>p. 352</td>
</tr>
<tr>
<td>Congenital webs</td>
<td></td>
<td>p. 352</td>
</tr>
<tr>
<td>Laryngeal stenosis</td>
<td></td>
<td>p. 352</td>
</tr>
<tr>
<td>Vocal cord sulcus</td>
<td>Rarely presents with other laryngeal symptoms</td>
<td>18.1, p. 392</td>
</tr>
<tr>
<td><strong>Post-traumatic</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arytenoid dislocation</td>
<td>Postintubation</td>
<td>pp. 364 and 381</td>
</tr>
<tr>
<td>Intubation granuloma</td>
<td></td>
<td>17.6, p. 365</td>
</tr>
<tr>
<td>Laryngeal stenosis</td>
<td></td>
<td>pp. 364-367</td>
</tr>
<tr>
<td><strong>Inflammatory</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acute laryngitis</td>
<td></td>
<td>pp. 354-356 and 357</td>
</tr>
<tr>
<td>Chronic nonspecific</td>
<td></td>
<td>pp. 358-361</td>
</tr>
<tr>
<td>laryngitis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Epiglottitis</td>
<td>Typical: muffled speech</td>
<td>pp. 356 and 357</td>
</tr>
<tr>
<td></td>
<td>Emergency with no prior history</td>
<td></td>
</tr>
<tr>
<td>Laryngeal edema</td>
<td></td>
<td>pp. 357-361</td>
</tr>
<tr>
<td>Pseudocroup</td>
<td></td>
<td>p. 354</td>
</tr>
<tr>
<td><strong>Neoplasms</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vocal nodules</td>
<td>Organic lesion secondary to a functional disorder</td>
<td>pp. 349-377</td>
</tr>
<tr>
<td>Reinke’s edema</td>
<td>Hoarseness, habitual throat clearing, low-pitched voice</td>
<td>pp. 359-361</td>
</tr>
<tr>
<td>Cysts</td>
<td>Postinflammatory</td>
<td>p. 368</td>
</tr>
<tr>
<td>Laryngeal papillomas</td>
<td>Besides vocal nodules, this is the most important differential diagnosis in</td>
<td>pp. 368-377</td>
</tr>
<tr>
<td></td>
<td>pediatric dysphonia</td>
<td></td>
</tr>
<tr>
<td>Contact granuloma</td>
<td>Most common in hypofunctional dysphonia, often associated with laryngeal reflux</td>
<td>p. 361</td>
</tr>
<tr>
<td>Vocal cord polyps</td>
<td></td>
<td>p. 368</td>
</tr>
<tr>
<td><strong>Functional</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Vocal abuse</td>
<td></td>
</tr>
<tr>
<td><strong>Neurogenic</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Recurrent laryngeal nerve</td>
<td></td>
<td>pp. 380-383</td>
</tr>
<tr>
<td>paralysis</td>
<td></td>
<td></td>
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</tbody>
</table>

**Diagnosis:** Every examination for dysphonia should begin with a detailed history. Besides the objective information contained in the history, a perceptual evaluation of the patient’s voice during speech and the observation of overall posture, muscular tone, mimetics, and gestures can provide further clues to possible causal factors. Information on occupational voice use is essential for evaluating a voice disorder. *Laryngoscopy* and *stroboscopy* (see pp. 387–389) supply important information on the nature of the voice impairment and on secondary morphologic changes.

**Psychogenic dysphonia and aphony:** Much experience is needed in order to assign a psychogenic cause to a voice disorder. These cases present with uncharacteristic laryngoscopic and stroboscopic findings. It is helpful to note, however, that most patients with psychogenic voice disorders can cough and laugh normally.

Psychogenic dysphonia or aphony should never be dismissed as a trivial condition.

Psychogenic dysphonia is a *psychosomatic disorder* whose true significance is often appreciated only during a prolonged course of therapy.

**Treatment:** Causal factors and underlying diseases should be corrected or eliminated as completely as possible before treatment is initiated. The actual voice therapy consists of an individualized program of vocal exercises. Today, numerous competing methods known collectively as “functional voice training” take into account not just the laryngeal aspects of voice production but also breathing, posture, and personality issues. The basic goal is the rapid restoration of physiological and economical phonation and general communication skills. For workers in most occupations, the voice is an indispensable tool for communication.

Many patients with pronounced dysphonia can keep their jobs only by undergoing an intensive, rigorous course of voice therapy, which may even require an inpatient program.

**Children,** as a rule, are not good candidates for formal voice therapy because they are not cooperative enough to follow the protocol. A better focus of treatment in these patients is to optimize motor functions while improving concentration and play behavior (e.g., ergotherapy).
Fig. Causes of functional voice disorders

External causes

General stresses:
- Emotional conflicts
- Circadian rhythm (night job)
- Noxious exposure (smoking, alcohol)
- Climate

Speaking situation:
- Ambient noise levels (environmental noise)
- Physical surroundings
- Listeners
- Microclimate

Overuse:
- Teachers:
  - Subject taught (e.g., sports, music, speech)
  - Lack of technical or teaching proficiency
  - Years at work

Leisure time:
- Singing in a choir

Internal causes

General physical constitution:
- Gender
- Endocrine status
- Age

Local constitution:
- Vocal apparatus (respiration, phonation, articulation)
- Hearing
- Kinesthesia

Mental constitution:
- Central and autonomic nervous system
- Interpersonal contacts
- Basic mood
- Affective behavior

Mood disorders and diseases:
- Fatigue
- Excessive stress
- Nervousness
- Upper respiratory tract diseases
- Severe systemic diseases

Dysphonia

18.1 Vocal cord sulcus

With a vocal cord sulcus (Fig.), the mucosa at the free edge of the vocal cord(s) is indrawn and fixed (arrows), making the epithelium less mobile or immobile in relation to the vocalis muscle and vocal ligament. Apparently this disorder is caused by muscle fibers inserting on the vocal cord mucosa or by residual dysfunction following a previous inflammation. The result is a hoarse voice. Treatment consists mainly of symptomatic voice therapy. Surgical release of the muscle fibers from the mucosa or resection of the affected mucosa is not widely practiced due to the risk of postoperative scarring.

Treatment of secondary organic changes: Vocal nodules may be an indication for surgical treatment, depending on whether they are hard or soft. This assessment requires much experience and very careful consideration.

Once vocal nodules have been surgically removed, it is essential to identify and treat the underlying cause of the lesions—e.g., with conservative voice therapy.

Surgical ablation is not indicated for vocal nodules in children. These lesions have a strong tendency to regress spontaneously by puberty, especially in boys.

Treatment of psychogenic voice disorders: Psychotherapy alone is unlikely to relieve hoarseness in these patients, and the plan of treatment should always include somatically oriented voice therapy. An essential task of the physician confronted with psychogenic dysphonia is first to motivate the patient to recognize the psychogenic nature of the complaint and accept the need for help.

Formerly, it was common for speech–language pathologists to use “surprise tactics” with the object of provoking a normal voice.

Surprise tactics are not fair to the patient and are purely of historical interest today.

Hyperfunctional Dysphonia

Epidemiology: Hyperfunctional dysphonia appears to be most prevalent in talkative, extroverted individuals, especially women.

Pathogenesis: A dynamic balance normally exists between subglottic pressure and muscular tone in the larynx. In hyperfunctional dysphonia, this balance is shifted toward higher pressures and tone values, causing the system to function at an uneconomical and inefficient level.

When the patient uses this uneconomical phonation mechanism for an extended period of time, sites of connective-tissue hyperplasia develop at the junction of the anterior and middle thirds of the vocal cords.
The glottic insufficiency in this condition is often not appreciated with conventional laryngoscopy and is seen only during stroboscopy. This study also shows increased amplitudes and an amplified traveling wave motion due to a decrease in vocal cord tension.

**Mixed Dysphonia**

Mixed forms of dysphonia may exhibit both hyperfunctional and hypofunctional features. The mixed pattern can be explained by various mechanisms of decompensation or maladaptive compensation. For example, compensatory participation of the ventricular folds in phonation may occur in a patient with hypofunctional glottic insufficiency. It is very difficult to establish the nature of the primary disorder in any given case.

**Mutational Voice Disorders**

The most common of these disorders involve a **prolonged voice mutation** or **incomplete mutation**. A **premature voice change** may reflect a serious hormonal disorder and should be investigated by a pediatric endocrinologist. A **mutational falsetto** results from increased cricothyroid muscle tension and always has a psychological cause. External downward pressure on the thyroid cartilage (the Gutzmann maneuver) will often evoke a normal-sounding voice. Generally, however, this condition requires a prolonged course of treatment to give the adolescent time to adapt to a normal chest voice. Even if the disorder is presumed to have a psychogenic cause, there is no need for specific psychotherapy in most cases. It is extremely important to educate the parents about the nature of the condition.

**Endocrine Dysphonia**

Endocrine dysphonia very often presents clinically as a harsh, cracking, low-pitched voice, frequently combined with loss of the singing voice and rapid fatigability. It is much more common in women than in men. Endocrine voice disorders may be caused by a disease-related hormonal imbalance or, more commonly, by exogenous hormone administration (therapeutic or for athletic doping). The voice may fluctuate during the course of the ovarian cycle. A common disorder in pregnancy is laryngopathia gravidarum. Menopausal women may also develop voice changes. The examiner should give particular attention to any signs of virilization that may reflect an abnormal androgen excess (e.g., due to a hormonally active tumor).

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These vocal nodules (synonyms: screamer's nodules, singer's nodules; Fig. 18.8) represent secondary morphologic changes based on the primary functional disorder.

**Symptoms:** The speaking voice is often raised to a higher pitch. Usually the voice sounds harsh, scratchy, and strained and is reduced in its ability to be raised or modulated. It is also common to find symptoms such as a general increase in muscular tension with neck and throat problems. In many patients, pronounced movements of the shoulder girdle can be observed during breathing (unphysiologic “upper chest breathing”).

**Diagnosis:** Laryngoscopy often shows incomplete glottic closure with an hourglass-shaped glottis and adduction of the false vocal folds during phonation, which may even prevent the true vocal folds from being seen. The epiglottis may be displaced posteriorly and may assume an almost vertical position. The vocal cords may be hyperemic, mimicking their appearance in laryngitis. Stroboscopy often shows a suppression of vibrational amplitudes and traveling wave motion.

**Hypofunctional Dysphonia**

Hypofunctional dysphonia may result from prolonged voice use or from a habitual tendency to speak with too little effort. The dynamic balance of vocal cord vibrations is shifted downward toward low pressures and tone values. The body posture is marked by decreased muscular tension, the breathing is shallow, and the voice sounds weak and breathy.
Treatment aims at reestablishing a normal hormonal balance. Often, however, these voice disorders respond poorly to treatment, and therapeutic exercises are prescribed to achieve optimum compensation.

Vocal Cord Paralysis

Synonym: laryngeal palsy

Symptoms: Unilateral vocal cord paralysis is characterized by a breathy, hoarse, weak voice that is markedly restricted in its pitch and volume range. The voice may deteriorate further over time due to the use of improper compensatory mechanisms. With bilateral vocal cord paralysis, severe respiratory distress may be the dominant feature. Often the voice sounds almost normal.

Diagnosis: The paralyzed vocal cord is fixed in a paramedian position (Fig. 18.9). It may be “excavated” as a result of muscular atrophy. The stroboscopic findings depend on the degree of vocal cord tension. With incomplete paralysis, the traveling wave motion may be detectable even in the absence of respiratory motion.

Treatment: Unilateral vocal cord paralysis is treated initially with conservative voice exercises, which should be supported by synchronous electrical stimulation to prevent muscular atrophy. Electrical stimulation alone is of no value without synchronous voice therapy.

If conservative voice therapy fails to give satisfactory improvement, phonosurgical procedures should be considered: thyroplasty (see 18.2) and arytenoid rotation. As a rule, surgery is not undertaken until at least one year after the onset of vocal cord paralysis to allow time for possible spontaneous recovery. Earlier phonosurgery may be considered for older patients in

18.2 Phonosurgical procedures

Ishikiri type I thyroplasty

Indication: unilateral vocal cord paralysis.

Principle: This procedure is based on the concept that the dynamic equilibrium of vocal cord vibrations can be modified indirectly by altering the cartilaginous framework without causing direct trauma to the vocal cords. The surgery is done under local anesthesia. A piece of thyroid cartilage is excised at the level of the paralyzed vocal cord (a). While the patient phonates, the cartilage is pressed inward until the voice improves (b). The cartilage is then fixed in that position with tissue adhesive or a silicone shim (c).

Nawka arytenoidectomy

Indication: bilateral vocal cord paralysis.

Principle: The glottis is widened by excising the arytenoid cartilage (d) and the posterior part of the vocalis muscle (e), usually on one side. The vocal cord mucosa is sutured to the mucosa of the ventricular fold (f).

Result: Usually an acceptable compromise can be achieved between ventilation and phonation.
18.3 Phono-surgery

**Definition:** Phono-surgery involves the use of surgical procedures to improve the voice. The following **phono-surgical guidelines** should be observed:

A normal traveling wave motion requires integrity of various structures: Reinke's space, the free edge of the vocal cords, and the vocal cord mucosa.

An effort should be made to obtain adequate contact between the vocal cords during the closed phase of the vibratory cycle.

A favorable working point should be found between generating a certain minimum subglottic pressure and speaking without undue force. This is not just a surgical task, however, because effective vocalization can be trained over a broad range with voice therapy.

Every phono-surgical procedure should be preceded by the detailed documentation of findings and adequate informed consent.

**Options:**

**Indirect laryngoscopy:**

*Indication:* Removal of vocal cord polyps, some cysts, and especially discrete marginal edema of the vocal cords.

*Principle:* The patient is conscious and holds the tongue forward with a gauze swab. Topical anesthesia is applied, and the lesions are removed with fine instruments. The vocal cords are observed with a microscope via a laryngoscopic mirror or with an endoscope linked to a video monitor.

*Advantage:* Preserving the muscular tone of the vocal cords makes it easier to see subtle changes and evaluate the immediate functional result by checking the voice intraoperatively.

**Direct laryngoscopy:**

The **indications** for this technique are relatively extensive organic lesions that need to be palpated, removed, or biopsied. It can also be used to remove (pre)malignant lesions that are unrelated to phono-surgical criteria.

*Principle:* General anesthesia is induced either by intubating the patient (see 17.14, p. 349) or using a jet ventilation laryngoscope.

*Advantage:* General anesthesia permits a longer, unhurried operation that allows very precise work.

*Disadvantage:* The quality of the voice cannot be assessed intraoperatively.

**Transcutaneous laryngeal framework surgery:** The principal options are the Ishii type I thyroplasty described in 18.2 and arytenoid rotation.

**Injection techniques:** Collagen and other materials can be injected for the augmentation of vocal cords that are atrophic due to recurrent laryngeal nerve paralysis, and botulinum toxin can be injected for spasmodic dysphonia. These techniques are reserved for highly selected indications.

**Laryngeal reinnervation techniques** are currently in development and are still classified as experimental procedures.

whom conservative voice therapy is ineffectual due to general physical weakness.

**Bilateral vocal cord paralysis** with serious respiratory distress should be managed in the acute stage by intubation or tracheotomy. Tracheotomized patients should be fitted with a speaking tube. This type of tube keeps the tracheostomy open and allows free respiration. On expiration, a small flap valve occludes the tube and the exhaled air can be used for phonation in the usual way. In chronic cases with satisfactory respiration that does not require a tracheotomy, an attempt can be made to widen the glottis with a minor surgical procedure on the vocal cords (arytenoidectomy, see 18.2). However, this almost always causes some degree of voice deterioration due to incomplete glottic closure.
Speech and Language Disorders

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- Normal Language Development 398
- Abnormal Language Development 398

19.2 Important Types of Speech and Language Disorders
- Dyslalia 400
- Dysgrammatism 400
- Rhinolalia 401
- Speech Fluency Disorders 401
- Dysarthria 403
19.1 Principles of Normal and Abnormal Language Development

According to conservative estimates, from 10% to 15% of all children under 6 years of age have a developmental language disorder. Ultimately, this type of disorder can profoundly impair the child’s psychosocial integration. To prevent this, it is essential to know the normal patterns of language development, the possible causes of developmental disorders, necessary test procedures, and the indications and options for treatment.

Normal Language Development

The milestones of normal language development are outlined in Fig. 19.1.

Abnormal Language Development

Causes

The causes of abnormal language development are numerous and diverse. When multiple components are present, it is no longer possible in many cases to determine the contribution of individual factors to the overall disorder. The classification of Nickisch and Gross (Table 19.1) summarizes the main causes and is helpful in understanding the etiopathogenesis of developmental language disorders.

Symptoms

Abnormal language development can have various clinical presentations. The possible deficits may involve:

- Sound recognition, articulation, and distinguishing sounds (dyslalia, stuttering);
- Expressive and receptive vocabulary;
- Speech comprehension;
- The observance of syntactical and grammatical rules (dysgrammatism);
- The comprehension and formulation of spoken or written sentences, reading;
- The contextually appropriate use of language; and
- Other disturbances of nonverbal communication, perception, motor skills, and overall cognitive-intellectual and social development.

Diagnosis

If a developmental language problem is suspected, it is generally necessary to conduct a comprehensive diagnostic evaluation covering all developmental aspects of the affected child. This includes the testing of speech and language in addition to nonlinguistic areas (general and manual motor skills, cognitive-intellectual development) and social interactions. Play behavior, communicative behavior, and social conduct are assessed based on personal observation or reports from others. Nonlinguistic areas of development are generally investigated by a pediatrician.

Every child with a speech or language problem should be tested for a hearing disorder.

The diagnostic target areas for speech and language disorders are as follows:

- Spontaneous speech
- General communicative behavior
- Speech comprehension
- Active vocabulary
- Grammar
- Ability to distinguish and use sounds that differentiate word meanings
- Articulation
- Language learning ability

Treatment

Impaired language development in children can be treated only by a structured exercise program that is tailored specifically to individual deficits and is administered by a speech–language pathologist. The program is based on a combination of play-therapy principles and pedagogic elements, depending on the age of the child.

Treatment should be instituted as soon as possible after the causes have been identified and should incorporate the child’s caregivers.

For children up to 2 years of age, initial intervention is generally indirect and consists of parental counseling. Children over 2 years of age can receive direct therapeutic intervention, but parenteral counseling is still important.

Psychosocial conditions appropriate for language acquisition should be present or created therapeutically. The scheme for improving language abilities follows the normal sequence of language development: language comprehension—vocabulary—grammar—articulation.
### Table 19.1 Causes of developmental language disorders

<table>
<thead>
<tr>
<th>Classification</th>
<th>Causes</th>
</tr>
</thead>
</table>
| **Isolated developmental language disorder** | - Familial language impairment  
- Psychosocial causes: overprotection, deprivation, bilingualism, lack of language stimulation  
- Disturbance of peripheral speech organs  
- Abnormal central coordination of peripheral speech organs (e.g., oral dyspraxia) |
| **Developmental language disorder (DLD) combined with other disorders** | - Peripheral hearing disorders  
- Auditory perception disorder  
- Peripheral visual disorder  
- Visual perceptual disorder  
- Peripheral or central motor disease with normal speech organs  
- Mental disorder (e.g., autism)  
- Congenital (prenatally acquired or hereditary) mental retardation  
  - With associated physical anomalies (syndrome)  
  - Without physical anomalies  
- Perinatally or postnatally acquired mental retardation (traumatic, inflammatory, hypoxic, metabolic, neurodegenerative) |

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Source: modified from Nickisch and Gross, see p. 411.

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The **goal of treatment** is to correct the developmental language deficit before the child enters school to permit mainstream schooling or prevent a delay in school enrollment.

**Prognosis**

The prognosis depends on the individual causes, the severity of the disorder, and the prompt initiation of services and treatment. Primary deficits can be compensated as the child grows older. Secondary disorders such as social withdrawal or aggressive modes of behavior may affect the overall picture. Children who have no severe, progressive, or refractory underlying diseases will generally have a good prognosis.

---

Fig. 19.4 Milestones of language development

1. Cooing period: undifferentiated vocalization  
2. Babbling period: development of language-specific sound differentiation, imitation of own and others’ intonation and stress patterns  
3. First word comprehension  
4. Increasing recognition of word meanings (symbolic function of language)  
5. One-word sentences, vocabulary up to 50 words  
6. Two-word sentences, vocabulary of about 900 words  
7. Multisyllabic sentences, first use of prepositions and pronouns, vocabulary of about 900 words  
8. Complex sentences, vocabulary growth, grammatical and articulation errors less frequent, use and comprehension of more complex questions
19.2 Important Types of Speech and Language Disorders

Given the fundamental importance of speech in all forms of human interaction, many branches of science and several medical specialties are concerned with the phenomena of normal and abnormal speech and language development, viewing speech and language disorders from varying perspectives and with various goals. Several main types of language disorders can be characterized to facilitate a systematic approach to these disorders and create a basis for routine clinical work.

Dyslalia

**Definition:** Dyslalia refers to any of a number of articulation disorders characterized by errors of pronunciation. Individual vocal sounds may be omitted, distorted, or replaced by other sounds. Dyslalia includes the inability to use sounds or sound combinations in a way that clearly differentiates word meanings.

**Classification:** Dyslalia can be classified on the basis of various criteria:

**Quantity:**
- Isolated dyslalia: defective pronunciation of a sound
- Partial dyslalia: defective pronunciation of certain sounds
- Multiple dyslalia: defective pronunciation of many sounds
- Universal dyslalia: defective pronunciation of virtually all sounds

**Constancy:**
- Constant dyslalia: the error is always present
- Inconstant dyslalia: the error is sometimes present

**Variability:**
- Consistent dyslalia: the nature of the error is always the same
- Inconsistent dyslalia: the errors or substitute sounds may vary

**Quality:** Specific articulation errors are named for the Greek letter equivalents of the affected consonants, with the ending “ism” or “tism” added: sigmatism, rhotacism, etc. Dyslalia most commonly affects the sibilants, particularly the “s” sound—a condition known as sigmatism or lisping. The prefix “para” indicates the use of a substitute sound, while the prefix “a” denotes the absence of a sound (“asigmatism”). When clinical findings are interpreted, moreover, it should be noted whether the articulation error occurs in sound groupings—i.e., in syllables, words, or sentences. Thus, while one sound may be correctly pronounced in isolation, it may be mispronounced within the context of syllables or words. In some cases this can result in universal dyslalia.

**Extent:** A key distinction in the diagnostic classification of dyslalia is whether the patient has an isolated (“pure”) articulation disorder or a general impairment of language development. Isolated dyslalia in a child under 3–4 years of age with a normal vocabulary and normal grammar is classified as “developmental dyslalia” or “physiologic dyslalia.” Fig. 19.1 (see p. 399) reviews the normal timetable for the acquisition of vocal sounds.

**Etiology:** “Organic dyslalia” is the term applied to articulation disorders caused by congenital or acquired defects at the peripheral receptive level (hearing impairment), central level, or peripheral expressive level (teeth). Central defects generally cause more complex deficits than just dyslalia, however. This is also true of mental retardation, general physical retardation, and serious systemic diseases. Insufficient language stimulation (deprivation) also tends to cause a more comprehensive disturbance of language development.

Dysgrammatism

**Definition:** Dysgrammatism is an inability to speak grammatically due to errors of word morphology (tense, number, gender) and syntax (sentence structure).

**Occurrence:** Within the context of normal language acquisition, dysgrammatic elements occur physiologically during the second and third years of life. After 3 years of age, however, a marked deviation from the age-specific norm usually indicates a significant developmental language disorder. As a rule, dysgrammatism is associated with other abnormalities of language acquisition. Dysgrammatism rarely occurs as a dominant complaint; when it does, it is often the cardinal symptom of an auditory disorder.

**Diagnosis:** The evaluation of general language development includes testing for dysgrammatism. The extent of the disorder is assessed on the basis of various tasks (spontaneous speech, repeating sentences, telling a story based on a series of pictures, retelling a story shown in pictures).
Treatment: The treatment of dysgrammatism is integrated into a complex, overall treatment plan that is patterned after physiologic language development. The prognosis depends on the extent of the overall disorder. Dysgrammatism may persist for some time as a refractory residual symptom. The prognosis is usually favorable, however.

Rhinolalia

Synonym: rhinophonia, rhinism

Definition and pathogenesis: Rhinolalia is altered speech caused by abnormal airflow through the nose during phonation. It may have an organic or functional cause. While it is technically correct to distinguish between altered voice sounds (rhinophonia) and altered sound production (rhinolalia), both terms are often used interchangeably. Several types of rhinolalia are distinguished based on the voice sound: Hypernasal speech (rhinophonia aperta): Nasal resonance is increased during speech as a result of velopharyngeal dysfunction, creating an ineffectual seal between the oral cavity and nasopharynx. The underlying cause may be a cleft palate or paralysis of the soft palate in a setting of myasthenia. Functional hypernasality may reflect nonuse of the soft palate following a tonsillectomy. Hyponasal speech (rhinophonia clausa): Nasal resonance is decreased during speech. Usually this has an organic cause based on obstructive lesions such as adenoid vegetations in the nose or nasopharynx. Mixed nasality (rhinophonia mixta): Both hyponasal and hypernasal features are present.

Diagnosis: The initial goal in diagnosis is to identify the specific type of rhinolalia that is present. In the mirror test, a cold mirror or a device called a Czemak plate is held beneath the nose while vowels are pronounced. Foggging of the mirror indicates the nasal air escape that characterizes hypernasal speech. A Czemak plate has rings that show the extent of foggging, permitting a semiquantitative assessment of nasal air escape. In the “ah-ee test,” the vowel sounds “ah” and “ee” are spoken in succession. With hypernasal speech, the tester will hear a nasal-sounding change in the vowel sound. This does not occur in hyponasal speech. The cheek inflation test (with the tongue protruded) can detect an organic cause of hypernasal speech. If the soft palate is shortened due to an incomplete cleft, the patient will be unable to inflate the cheeks with air while the tongue is protruded. The “backdrop sign” is an important finding in patients with paralysis of the soft palate (see p. 76). Hypernasal speech can be accentuated by head turning toward the unaffected side.

An underlying neurologic disease may also have to be excluded in some cases.

Treatment: The initial treatment for functional disorders is an intensive voice therapy program, which may be long and arduous in some patients. The treatment for organic hypernasal speech due to a cleft palate is integrated into a comprehensive plan for treating the underlying disorder (see p. 78). Organic hypernasal speech due to an obstructive lesion in the nose or nasopharynx is easily corrected with surgery (see p. 58).

Speech Fluency Disorders

Stuttering

Definition: Stuttering is an impairment of speech fluency that is independent of the will of the speaker. The strict classification of stuttering into tonic and clonic forms is no longer considered essential due to a lack of therapeutic implications. Disfluency occurs physiologically during language acquisition between 2½ and 4 years of age.

Epidemiology: The incidence of stuttering is approximately 1% in the overall population and approximately 4% in children, regardless of language and culture. Males are predominantly affected.

Etiology: The etiology of stuttering is uncertain. Organ theories stress the causal significance of genetic components as well as hyperexcitability of the central nervous system. There are also learning theory models of causation, and there is evidence to support a psychogenic cause in specific cases. For clinical purposes, it is useful to postulate a multifactorial etiology with hereditary influences along with organic, psychological, and environmental factors.

Symptoms: Tonic stuttering is characterized by a blocking of speech at the beginning of words or sentences, usually associated with a generalized rise in body tension. Clonic stuttering is characterized by repetitions of sounds, syllables, or words. Secondary characteristics includes changes in breathing, mimetics, gestures, and autonomic responses that accompany stuttering.

Diagnosis: The diagnostic evaluation of stuttering should cover the following specific points:
- Basic symptoms and accompanying features by number and severity, fear of speaking and avoidance behavior, body image, variability of symptoms, situations that elicit or exacerbate stuttering, or situations that elicit fluent speech.
• Causal factors and progression over time, social milieu (family, friends, acquaintances, work environment)
• Awareness of the stuttering by self and others, attitude toward the stuttering (self-consciousness)
• Speaking demands, quality of information, emotionality, and motivation

In children, diagnosis should begin by assessing the relative importance of the stuttering symptom within the framework of a general developmental language disorder, which is usually present.

Treatment: Based on the assumption of a multifactorial etiology, the treatment of stuttering combines symptom-oriented elements with psychotherapeutic-psychosocial treatment strategies to create an integrative, multidimensional program. The focus of the program is on improving self-perception, desensitizing the stutterer to his or her own disorder, modifying the symptom by the use of specific speaking techniques, generalizing the new speaking techniques into spontaneous communication, and establishing support by referring the patient to a self-help group, for example. Tranquilizers and other medications may be helpful as adjuncts.

In children, an effort is made to direct the patient’s attention to fluent episodes and reduce any fears that are associated with speaking. In exceptional cases, elements of adult therapy may be applied in this age group for the purpose of direct symptom modification. Parental counseling is also emphasized.

Prognosis: Regardless of the age group, approximately one-third of patients will recover completely, one-third will improve, and one-third will experience no change.

Cluttering

Cluttering is a fluency disorder characterized by an abnormally rapid and irregular rate of speech delivery. The speech may be difficult to understand due to the omission of sounds, syllables, or even entire words. The symptoms are improved (even with strangers) by concentrating on the verbal output and deliberately slowing the rate of speaking. The clutterers themselves rarely perceive their speech as being abnormal. The etiopathogenesis of cluttering is as poorly understood as that of stuttering.

After potential causes have been investigated, the main task in diagnosis is to differentiate cases that require treatment from cases that have no pathologic significance.

As a rule, no specific treatment is necessary in adults because usually they are not self-conscious about their disorder. Cluttering in children is usually part of a general developmental language disorder and is treated within that context.

Aphasia

Definition: Aphasia refers to the partial or complete loss of speech (formulation, memory, comprehension) after the completion of language acquisition.

Epidemiology: The incidence of aphasias syndromes is estimated at 60 new cases per 100,000 population per year, with a rising trend.

Etiology and pathogenesis: Predisposing underlying diseases are arterial hypertension, disorders of fat metabolism, diabetes mellitus, and generalized atherosclerosis. More than 80% of cases have a cerebrovascular cause. More than 90% of these cases are caused by a thrombotic or embolic vascular occlusion, and less than 10% by an intracranial hemorrhage.

Symptoms: Aphasia may present clinically as a disturbance of both written and spoken language, meaning that all the coding functions for the same information may be impaired. All linguistic components—phonology, lexicon, syntax, semantics—may be affected. Based on the cerebrovascular etiology, various patterns of injury may be encountered in aphasic patients, and other cognitive deficits such as altered consciousness or a decline in intelligence may accompany the language disorder.

The peripheral speech organs and especially the peripheral auditory apparatus are not directly affected by aphasia.

Classification: There is still no universally accepted scheme for the classification of aphasia. The classification of Poeck et al., based on the dominant features of different aphasias, is perhaps the most widely used system in German-speaking countries. It is the classification upon which the Aachen Aphasia Test (AAT) is based, of which an English-language version (EAAT) is also in use.

Global aphasia is the most common cerebrovascular type of aphasia and is considered the most severe form. It results from damage to the brain in the territory of the middle cerebral artery. Regardless of the affected hemisphere, global aphasia is characterized by a severe impairment of all receptive and expressive language functions.

The term Wernicke's aphasia is currently preferred over older terms such as "sensory," "receptive," "acoustic" and "posterior" aphasia, because the latter terms do not reflect the potential complexity of the disorder and are not helpful in directing treatment. Wernicke's aphasia results from lesions in the territory...
of the posterior temporal artery. The cardinal symptoms are poor speech comprehension and a fluent but often unintelligible output (“fluent nonsense”). Thus, Wemicke’s aphasia bears a close resemblance to global aphasia from the standpoint of language comprehension.

**Amnestic aphasia** is caused by damage to the temporoparietal area of the brain. The principal causative lesions are brain tumors, temporal lobe abscesses, and degenerative processes. Its cardinal symptom is word-finding difficulty, and the patient tries to compensate for this by paraphrasing. As a rule, amnestic aphasia has the best prognosis of all the aphasias, barring progression of the underlying disease.

The term **Broca’s aphasia** is preferred over the older terms “motor” and “expressive” aphasia, which do not accurately describe the nature of the disorder. The main features of Broca’s aphasia are a marked increase in speaking effort combined with poor articulation and altered prosody (the “melody” of speech). The disturbance is never confined to language expression and always includes comprehension difficulties. Dysgrammatism or agrammatism is also present. The classification also includes special forms of aphasia, which are beyond our present scope.

**Diagnosis:** Regardless of the type of aphasia present, it is currently believed that every aphasic patient is fundamentally impaired at all language levels, and that different forms differ mainly in the predominance of certain features. The **Aachen Aphasia Test (AAT)** is among the instruments that can be used to differentiate and quantify these features. **Audiologic–phoniatric testing** is performed in aphasic patients for the classification and quantification of language disorders in both receptive and expressive functions.

**Differential diagnosis:** Aphasia requires differentiation from **dysarthria**, which results from the paralysis or incoordination of respiratory, voice and speech muscles due to an organic brain lesion. Differentiation is also required from **developmental language disorders**, **language disorders in psychiatric conditions**, and from **apraxia**, which is an inability to perform voluntary and intentional movements with one part of the body in the absence of muscular weakness or paralysis.

**Treatment:** The goal of treatment in aphasia is to restore the communication skills needed to cope with everyday situations, achieve social reintegration, and ideally return the patient to work. The overriding goal is to restore overall communication more than rehabilitate its separate components. The treatment of aphasia can be divided into three phases:

- During the **activation phase** in the first 6 months after the onset of aphasia, treatment focuses on the general activation of seeing, listening, reading, and writing. Automatic terms such as number sequences, days of the week, months, etc., can be used to stimulate more voluntary words and phrases. Intact abilities are used to “unblock” impaired functions, proceeding for example from speaking with the therapist to repeating words after the therapist and finally naming objects.
- In the **exercise phase**, which begins after the aphasic syndrome has stabilized, the therapy consists of exercises tailored to the specific disorder. Largely intact modalities are used as an access point, linguistic principles are conveyed, and drills are performed to condition and improve certain language skills. Compensatory strategies are developed to cope with irreversible deficits.
- In the **consolidation phase**, further disorder-specific work will no longer improve the patient’s language skills. The emphasis in this phase is on reestablishing the patient into everyday life and reestablishing contact with the environment.

**Dysarthria**

**Synonym:** dysarthrophonia

**Definition:** Dysarthria is a speech disorder based on an impairment of the neural mechanisms that control speech movements.

**Etiopathogenesis and symptoms:** Given its neurologic cause, dysarthria may affect one or more of the systems involved in the production of speech: respiration, phonation, and articulation. Prosody, or the melody of speech, is also affected in most cases. The speech of dysarthric patients is characterized by altered muscular tone, weakness, slowing, incoordination, or hyperkinetic symptoms. Several forms of dysarthria can be distinguished based on clinical criteria:

- **Spastic dysarthria:**
- **Etiopathogenesis:** Spastic dysarthria is the most common form occurring after severe craniocerebral trauma. It appears to involve a spastic paralysis caused by damage to the first motor neuron, leading to acceleration-dependent muscular hypertonicity with a decrease in strength and fine motor skills. Reflex functions (swallowing, coughing, gagging) are preserved.
- **Symptoms:** The voice sounds strained, consonants are pronounced imprecisely, and the speech has a slow, monotonic quality. Unilateral lesions may be associated with subtle clinical findings.
- **Rigid hyperkinetic dysarthria:**
- **Etiopathogenesis:** The most frequent cause is Parkinson’s disease, in which rigid hyperkinetic dysarthria is an early symptom.
- **Symptoms:** The voice sounds soft and breathy and has a limited range.
Ataxic dysarthria:
Etiopathogenesis: This form is presumably based on deficits in cerebellar motor functions.
Symptoms: Patients typically have variable articulation errors and uncontrolled variations in vocal pitch and loudness. The voice may sound harsh and strained, and voice breaks are frequent.

Hyperkinetic and dystonic forms:
Etiopathogenesis: The causes are diverse and include brainstem lesions as well as rare neurologic disorders such as Huntington's disease or athetosis.
Symptoms: Tremor and myoclonias can cause variable impairment of the voice and speech, depending on the affected muscle group.

Hypotonic dysarthria:
Etiopathogenesis: Hypotonic dysarthria is a characteristic feature of bulbar palsy like that occurring in association with brainstem lesions (e.g., amyotrophic lateral sclerosis). A lesion of the second motor neuron leads to a flaccid paralysis of the affected muscles.
Symptoms: Characteristic features are muscular hypotonicity, loss of strength, and the limitation or abolition of voluntary and reflex movements. The voice sounds soft, breathy, and hypernasal. Consonant articulation is slowed and imprecise.

Diagnosis: The diagnosis of dysarthria is always based on a neurologic examination to investigate possible causes, determine the location of the lesion, and assess the extent of the underlying neurologic disease.

Phoniatric tests evaluate phonation and articulation, oral communication, reading, and writing. A quantitative assessment can be made by rating the degree of various criteria on a point scale, such as:
• Ability for voiced phonation
• Voice quality (harsh, breathy, strained?)
• Habitual pitch and volume
• Voice stability (pitch and loudness fluctuations, trembling?)
• Jaw movements (visual assessment)
• Vowel and consonant articulation
• Resonance (nasality?)
• Slow or rapid rate of speech during repetition or reading
• Fluency of speech (pauses, iterations, voice arrests?)
• Intonation, accent (monotoned speech?)

Treatment: Initial treatment is directed toward improving the underlying disease. Treatment planning should take into account the individual circumstances of the patient (Table 19.2). The cornerstones of treatment are reviewed in Table 19.3.
Various techniques are used to promote the motor functions essential for speech: special exercises, therapeutic aids (e.g., bite block or pointing board), prosthetic measures (e.g., palatal lift prosthesis), adaptive measures (changing the mode of communication, technical aids such as a Palm Pilot), or surgical procedures (velopharyngoplasty, posterior pharyngeal wall augmentation to improve nasopharyngeal closure). The prognosis in all forms of dysarthria depends on the underlying neurologic disease. Generally the disorder cannot be corrected but often it can be improved.
### Table 19.2 Factors that influence treatment planning

<table>
<thead>
<tr>
<th>Classification</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neurologic status and prior history</td>
<td>Location, nature, and extent of brain damage, time since lesion onset, progression or stability of the disease</td>
</tr>
<tr>
<td>Age</td>
<td>Younger patients usually have more rehabilitation potential owing to a better health status and psychosocial conditions</td>
</tr>
<tr>
<td>Personality</td>
<td>Optimistic, goal-oriented patients benefit more from treatment than uncompromising, less confident, or perfectionistic patients</td>
</tr>
<tr>
<td>Psychosocial status</td>
<td>A stimulating social environment and the availability of helpers will reinforce the transfer of relearned skills</td>
</tr>
<tr>
<td>Associated neuropsychiatric deficits</td>
<td>Deficits of attention, memory, learning, planning, language, affect, disease insight, perception, drive, and motivation can adversely affect therapeutic response</td>
</tr>
<tr>
<td>Degree of handicap</td>
<td>This assessment is based on individual requirements and on social and occupational demands</td>
</tr>
</tbody>
</table>

Source: Ziegler et al., see p. 411.

### Table 19.3 Guidelines for the treatment of dysarthria

<table>
<thead>
<tr>
<th>Guidelines</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Begin treatment early</td>
<td>To prevent maladaptive responses. Muscular functions that are more intact tend to become overactive and thus inhibit the recovery of other motor skills. Optimize residual skills in progressive diseases</td>
</tr>
<tr>
<td>Focus on the key disorders</td>
<td>First work on the function that is uppermost in the hierarchy of the causal chain</td>
</tr>
<tr>
<td>Modify the underlying disorder</td>
<td>Start with measures to restore physiologic function: regulation of muscle tone, postural correction, etc.</td>
</tr>
<tr>
<td>Promote compensatory behavior</td>
<td>Compensation is necessary for the maximum utilization of residual motor capacities</td>
</tr>
<tr>
<td>Promote speech awareness</td>
<td>The patient can no longer rely on speaking “automatically” or without special effort</td>
</tr>
<tr>
<td>Differentiate self-perception</td>
<td>The ability for self-perception needs to be developed</td>
</tr>
<tr>
<td>Create motivation for treatment</td>
<td>Motivation should not be confused with hope for recovery</td>
</tr>
</tbody>
</table>

Source: Ziegler et al., see p. 411.
Appendix: Emergencies and Primary Measures
Overview and Referrals for Detailed Information

Emergency situations are common in the head and neck region due to the complex anatomical relationships and the broad spectrum of possible diseases. They may present as life-threatening conditions associated with one or more of the following symptoms: hemorrhage, respiratory distress (dyspnea, stridor), acute dysphagia, and signs of local or systemic inflammation. Acute hearing loss, acute tinnitus, acute vertigo, and facial nerve paralysis are not life-threatening but still represent an acute situation. Skull fractures are also classified as emergencies. The diseases that are relevant in emergency medicine are covered in separate chapters and are reviewed in Table 1, which lists not only life-threatening events but also diseases and injuries in which immediate intervention is necessary to prevent complications or permanent disability.

Otologic Emergencies

Life-threatening hemorrhages are extremely rare, but any bleeding from the ear canal should be investigated in order to determine its cause and confirm the integrity of the tympanic membrane, ossicular chain, and inner ear. Injuries to the tympanic membrane and ossicular chain require immediate evaluation by a specialist, one reason being to exclude a lesion of the inner ear (damage to the auditory and/or vestibular apparatus).

Inflammations of the ear are mainly considered emergencies when the process transcends the boundaries of the ear (mastoiditis, brain abscess, meningitis, sepsis), there is evidence of facial nerve paralysis, and/or there are definite signs of inner ear involvement (labyrinthitis, hearing loss, vertigo). This not only applies to acute inflammations but also to cholesteatoma, which is likely to produce complications. Auricular inflammations and trauma carry a risk of cartilage damage.

Acute vestibulocochlear dysfunction that presents with hearing loss, tinnitus, and/or vertigo generally requires immediate diagnostic and therapeutic intervention. The same applies to paralysis of the facial muscles, which requires immediate investigation and treatment regardless of whether the facial nerve damage has an idiopathic, inflammatory (otitis media, cholesteatoma), neoplastic (parotid malignancy, vestibular schwannoma), or traumatic cause (temporal bone fracture).

Sinonasal Emergencies

Nosebleed (epistaxis) is the most common emergency, although life-threatening bleeding is relatively rare. Acute sinusitis becomes an emergency when the inflammation spreads to involve the orbit or eye and/or the meninges or frontal lobes of the brain, and immediate action is required.

Inflammations of the external nose can lead to cartilage liquefaction and requires appropriate treatment due to the risk of angular vein thrombosis and cerebral venous thrombosis.

Septal hematomas and abscesses can occur after nasal trauma and generally require immediate intervention. Emergencies in the broad sense include fractures of the nasal bone, zygoma, or orbital floor and intranasal foreign bodies.

Oropharyngeal Emergencies

Besides the inflammatory complications of tonsillitis (peritonsillar and parapharyngeal abscess), postoperative bleeding after tonsillectomy is a typical emergency situation. The erosion of blood vessels by malignant tumors can also provoke massive bleeding. Impalement injuries may create a nidus for abscess formation.

Bilateral choanal atresia in newborns generally requires immediate intubation because newborns are obligate nose breathers, especially while feeding.

Laryngotracheal Emergencies

Obstruction of the airways by swelling, a tumor, a foreign body, and/or blood and secretions necessitates emergency airway intervention (intubation, cricothyrotomy, tracheotomy).

Neck Emergencies

The most common emergencies in the neck are injuries (stabbing, gunshot, strangulation, blunt trauma), which are marked by bleeding and airway obstruction, and inflammations (parapharyngeal abscess, cervical cellulitis), which may progress to mediastinitis.
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<td>Acute sensorineural hearing loss: sudden hearing loss (pp. 267 – 269), acoustic trauma, shock-wave trauma, explosion trauma (pp. 260 – 262), barotrauma (p. 251)</td>
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<td>Tympanic membrane and middle ear injuries (pp. 250 – 251), temporal bone fractures (pp. 302 – 305)</td>
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<tr>
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Fig. 17.12a  Comp. photo. Fa. Richard Wolf, Knittlingen
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