Imaging Review of the Temporal Bone: Part I. Anatomy and Inflammatory and Neoplastic Processes

From a clinical-radiologic standpoint, there are a limited number of structures and disease entities in the temporal bone with which one must be familiar in order to proficiently interpret a computed tomographic or magnetic resonance imaging study of the temporal bone. It is helpful to examine the region in an organized and systematic fashion, going through the same checklist of key structures each time. This is the first of a two-part review that provides a practical approach to understanding temporal bone anatomy, localizing a pathologic process with a focus on inflammatory and neoplastic processes, identifying pertinent positives and negatives, and formulating a differential diagnosis.

© RSNA, 2013

Learning Objectives:
After reading the article and taking the test, the reader will be able to:

- Identify important anatomic landmarks in the temporal bone
- Recognize clinical presentations associated with important inflammatory and neoplastic conditions in the temporal bone region
- Describe imaging features of some common inflammatory conditions in the temporal bone
- Distinguish between various tumors of the temporal bone based on CT and MR findings

Accreditation and Designation Statement
The RSNA is accredited by the Accreditation Council for Continuing Medical Education (ACCME) to provide continuing medical education for physicians. The RSNA designates this journal-based activity for a maximum of 1.0 AMA PRA Category 1 Credit™. Physicians should claim only the credit commensurate with the extent of their participation in the activity.

Disclosure Statement
The ACCME requires that the RSNA, as an accredited provider of CME, obtain signed disclosure statements from the authors, editors, and reviewers for this activity. For this journal-based CME activity, author disclosures are listed at the end of this article.

Online CME
See www.rsna.org/education/ry_cme.html

1 From the Department of Radiology, Massachusetts Eye and Ear Infirmary, 243 Charles St, Boston, MA 02114. Received April 17, 2012; revision requested June 4; revision received December 14; accepted January 7, 2013; final version accepted January 29. Address correspondence to A.F.J. (e-mail: amy_juliano@meei.harvard.edu).

© RSNA, 2013
There are a limited number of structures and disease entities in the temporal bone with which one must be familiar in order to proficiently interpret a computed tomographic (CT) or magnetic resonance (MR) imaging study of the temporal bone. It is helpful to examine the region in an organized and systematic fashion, going through the same checklist of key structures each time.

The temporal bones comprise the lateral skull base, forming portions of the middle and posterior fossae. Each temporal bone is composed of five osseous parts: the squamous, mastoid, petrous, tympanic, and styloid portions.

Several intrinsic channels, intrinsic fissures, and extrinsic sutures are often apparent on CT images and can mimic fractures (pseudo-fractures) (1). The major anatomic landmarks of the temporal bones are depicted on axial and coronal CT images (Figs 1–5) and are described in the following subsections.

**Temporal Bone Anatomy**

**External Ear**

The external ear includes the auricle and external auditory canal (EAC), which extends medially to the tympanic membrane. The tympanic membrane attaches to the tympanic annulus and measures approximately 10 mm in diameter. The normal tympanic membrane can often be faintly discerned on CT images. The lateral third of the EAC is fibrocartilaginous, while the medial two-thirds of the EAC is surrounded by the tympanic portion of the temporal bone (Fig 5).

The anterior wall of the EAC also forms the anterior aspect of the glenoid fossa. The posterior wall of the EAC forms the anterior margin of the mastoid segment of the temporal bone and is resected during canal wall-down mastiodectomy.

**Middle Ear and Mastoid**

The middle ear is an air-filled cavity within the petrous portion of the temporal bone that contains the ossicular chain and is bounded by the tympanic membrane laterally, the inner ear structures (surrounded by the otic capsule and the cochlear promontory) medially, the tegmen tympani superiorly, and the jugular wall (floor) inferiorly (Fig 5). The scutum is a sharp bony projection to which the tympanic membrane is attached superiorly (Fig 5). The tegmen refers to a thin plate of bone that separates the dura of the middle cranial fossa from the middle ear and the mastoid cavity. The tegmen tympani is the roof of the middle ear, and the tegmen mastoideum is the roof of the mastoid (Figs 4, 5). The posterior wall of the middle ear cavity is irregular and includes the facial recess (also referred to as the facial nerve recess), pyramidal eminence, sinus tympani, and round window niche, from lateral to medial (Fig 3). The pyramidal eminence overlies the stapedius muscle, which inserts onto the head of the stapes. Another important area of the middle ear is the Prussak space (superior recess) (Fig 5). This space is margined by the pars flaccida and scutum laterally, the lateral malleal ligament superiorly, and the neck of the malleus medially.

The middle ear can be further subdivided into the epitympanum (attic) superior to the level of the tympanic membrane, mesotympanum at the level of tympanic membrane, and hypotympanum inferior to the level of tympanic membrane (Fig 5). The hypotympanum contains the opening to the eustachian tube, with the internal carotid artery seen along its medial margin (Fig 4).

The mesotympanum contains the majority of the ossicular chain. The ossicular chain is composed of three bones: malleus (head, neck, anterior process, lateral process, and manubrium), incus (body, short process, long process, and lenticular process), and stapes (head/capitellum, anterior crus, posterior crus, and footplate) (Figs 2–5). The manubrium of the malleus is attached to the tympanic membrane, and the head of the malleus articulates with the body of the incus in the epitympanum forming the incudomalleal joint, which has a characteristic “ice cream cone” configuration on axial sections (Fig 2) (2). The lenticular process of the incus extends at approximately a right angle from the long process of the incus to articulate with the head of the stapes, forming the incudostapedial joint. The footplate of the stapes attaches to the oval window of the vestibule (Fig 5).

There are four suspensory ossicular ligaments: superior malleal, lateral
The malleus, posterior malleus, and posterior incudal. These ligaments are sometimes visible on CT images as thin linear structures. The lateral malleal ligament is most commonly identifiable among the suspensory ligaments (Fig 4) (3). The tensor tympani muscle arises from the superior surface of the cartilaginous part of the eustachian tube, courses posteriorly in the medial portion of the middle ear, turns sharply at the terminus of the cochleariform process, and attaches to the neck of the malleus (Fig 3).

The epitympanum communicates with the mastoid via the aditus ad antrum (Fig 1). Normally, the mastoid is an air-filled cavity divided into numerous compartments by mastoid septations. The mastoid air-cell size and configuration are highly variable (4). The mastoid air cells are traversed by the Koerner septum, which is a thin bony structure formed by the petrosquamous suture that extends posteriorly from the epitympanum, separating the mastoid air cells into medial and lateral compartments. The medial mastoid air cells are separated from the adjacent sigmoid sinus by the sigmoid plate.

Inner Ear

The inner ear is situated within the petrous portion and comprises the osseous labyrinth, which includes the cochlea, vestibule, and semicircular canals (Figs 2–5). The cochlea contains the end organ for hearing while the vestibule and semicircular canals are responsible for balance and equilibrium. The densest portion of the temporal bone that surrounds the osseous labyrinth is termed the otic capsule (Fig 2). The osseous labyrinth encapsulates the membranous labyrinth, which contains endolymph and is surrounded by perilymph. The structures of the membranous labyrinth (cochlear duct, utricle, saccule, semicircular ducts, and endolymphatic duct and sac) cannot be discerned at CT.

The cochlea is a spiral-shaped structure with 2½ to 2⅔ turns, including the basal, middle, and apical turns, which are separated by interscalar septae (Figs 2–4). The osseous spiral lamina is a microanatomic structure that can be faintly
discerned on thin-section temporal CT images; it extends from the modiolus, paralleling the interscalar septae (5). The lateral aspect of the basal turn of the cochlea bulges into the middle ear cavity, forming the cochlear promontory (Fig 5). The nerve of Jacobson (branch of cranial nerve IX) courses over the cochlear promontory. The cochlear nerve passes from the internal auditory canal through the bony canal for the cochlear nerve (also referred to as the cochlear fossa or cochlear aperture) into the modiolus (Fig 2), which is a crown-shaped structure centered within the cochlea that transmits branches of the cochlear nerve to the organ of Corti. The organ of Corti is the end organ for hearing but is not visible on CT images.

The bony vestibule is an ovoid space located superior and posterior to the cochlea, which connects to the semicircular canals (Fig 2). There are three semicircular canals—superior, posterior, and lateral (also called horizontal)—which are oriented orthogonal to one another (Figs 1, 5). Each semicircular canal has a dilatation termed ampulla at one end. The posterior and superior semicircular canals share a common crus formed by the fusion of the posterior crus of the superior semicircular canal and the anterior crus of the posterior semicircular canal, which then opens into the superomedial part of the vestibule. The lateral semicircular canal, on the other hand, has two separate openings into the vestibule.

The endolymphatic duct extends from the posterior aspect of the vestibule toward the posterior cranial fossa, and ends in a blind pouch, the endolymphatic sac, at the posterior margin of the petrous ridge. The bony vestibular aqueduct (Fig 3) surrounds the endolymphatic duct and normally measures up to 1 mm at the midpoint and 2 mm at the operculum, according to the Cincinnati criteria.

The cochlear aqueduct is a narrow bony channel that surrounds the perilymphatic duct and extends from the basal turn of the cochlea, anterior to the round window, to the subarachnoid space adjacent to the pars nervosa of the jugular foramen. The cochlear aqueduct normally measures up to 0.1–0.2 mm in its midpoint and is widest at the medial orifice (6).

**Internal Auditory Canal**

The internal auditory canal (IAC) is a channel in the petrous bone that is somewhat variable in size, shape, and orientation. The medial opening of the IAC is termed the porus acusticus. The lateral end of the IAC is termed the fundus and abuts the labyrinth. At the fundus, a transverse crest (crista falciformis) divides the IAC into superior and inferior compartments (Fig 5). A vertical crest (“Bill’s bar”) divides the superior compartment into anterior and posterior components. The facial nerve is located in the anterosuperior compartment, the cochlear nerve in the anteroinferior compartment, and the superior and inferior vestibular nerves in the superoposterior and inferoposterior compartments, respectively. A useful mnemonic devised for the location of the nerves in the anterior compartment is “Seven (cranial nerve VII) Up Coke (cochlear nerve) down.”

**Facial Nerve (Cranial Nerve VII)**

Although the facial nerve itself cannot be directly visualized on CT images, its bony canal defines its course. The facial nerve exits the lateral pons, traverses the cerebellopontine angle (CPA) (cisternal segment), passes through the porus acusticus, courses through the IAC in the anterosuperior quadrant above the crista falciformis (canalicular segment), and through the petrous bone anterior to the cochlea (labyrinthine segment) to reach the geniculate ganglion (Fig 1), from which the greater superficial petrosal nerve arises. From here (anterior or first genu), the facial nerve turns posteriorly toward the medial aspect of the middle ear cavity ( tympanic segment). On coronal CT images, the proximal portion of the tympanic segment and the distal portion of the labyrinthine segment of the facial nerve can be visualized superior to the cochlea, resulting in the appearance of “snake eyes” or “snail eyes” (Fig 4). The tympanic segment of the facial nerve runs inferior to the lateral semicircular canal and superolateral to the oval window (Fig 5). On MR images, enhancement is normally seen at and distal to the anterior genu of the facial nerve related to peri- and epineural venous plexuses. The intracanalicular and labyrinthine segments do not normally enhance with use of contrast material. The bone covering the tympanic segment of the facial nerve can be dehiscent, presenting a potential hazard during ossicular replacement surgery. In the posterior wall of the middle ear cavity adjacent to the facial recess, the facial nerve turns inferiorly forming the posterior or second genu. The mastoid segment of the facial nerve courses through the medial portion of the mastoid bone until it exits the skull through the stylomastoid foramen and enters the parotid gland. The chorda tympani branches off of the mastoid segment of the facial nerve and returns to the middle ear cavity via the canaliculus of the chorda tympani.

**External Auditory Canal**

EAC cholesteatoma.—EAC cholesteatoma is a rare lesion with an incidence of approximately 0.1%–0.5%. Most cases are spontaneous or idiopathic, although these lesions can also occur secondary to prior trauma, surgery, or radiation (7). Most patients are in an older age group and present with chronic dull pain and otorrhea, most commonly unilaterally (7). Pathologically, it is characterized by local invasion of the lining squamous epithelium of the EAC into the underlying bone, resulting in canal wall erosions and periostitis. On CT images, the lesion is characterized by focal soft tissue within the EAC (typically the inferior wall), with erosion extending into the underlying bone (Fig 6). These imaging findings are nonspecific, however, and may be mimicked by entities such as carcinoma and otitis externa. Therefore, correlation with clinical symptoms is essential.

Keratosis obturans.—Keratosis obturans represents an expansile accumulation of keratin debris within the EAC.
In contrast to EAC cholesteatomas, it occurs in younger patients and tends to be bilateral (9). Clinically, these patients have severe pain and conductive hearing loss, with otorrhea being relatively rare. There is an association of this entity with sinusitis and bronchiectasis (8,9). On CT images, there is diffuse widening of the EAC by an epidermal plug, which may cause smooth scalloping of the surrounding bone, but there is no erosion or periostitis (Fig 7).

**Malignant otitis externa.**—Necrotizing or malignant otitis externa occurs most commonly in elderly diabetic patients and other patients in immunocompromised states. The patients present with severe otalgia and otorrhea, and there is a high mortality rate (10). The infection is usually caused by *Pseudomonas aeruginosa*. Clinically one sees granulation tissue in the inferior EAC at the bone-cartilaginous junction. Infection can then spread via fissures of Santorini into the soft tissues beneath the skull base, leading to skull base osteomyelitis, which can be heralded by cranial neuropathies. On images, soft-tissue thickening in the external canal is noted, often with bone destruction and inflammatory changes in the mastoid. Skull base osteomyelitis results from extension of the destructive process into the clivus, jugular foramen, and prevertebral soft tissues (Fig 8) (10). Abscesses can develop in the epidural space, brain parenchyma, and the prevertebral space as a complication. MR and CT are complementary modalities for evaluation of this entity, with the bone windows at CT showing the destructive process to greater advantage, and MR imaging better demonstrating associated soft tissue complications. The imaging differential diagnosis includes nasopharyngeal carcinoma with secondary obstruction of the eustachian tube orifice causing a secondary otomastoiditis.

**Middle Ear**

**Acute otitis media.**—Acute otitis media is primarily a disease of infants and young children. Patients present with fever, otalgia, and a red bulging tympanic membrane. The infection is usually caused by bacteria such as *Streptococcus* species or *Haemophilus influenzae*. Imaging is usually not necessary in uncomplicated acute otitis media (11). If imaging is performed, one sees opacities in the middle ear and mastoid, with possible fluid levels. There is preservation of the mastoid trabeculae and the overlying cortex. Complicated mastoiditis is suggested clinically by the presence of postauricular erythema, tenderness, and edema. In this clinical situation, imaging is crucial to exclude complications, which may be intratemporal or intracranial. Coalescent mastoiditis represents destruction of the mastoid trabeculae, which may extend to the inner or outer cortex either by means of enzymatic resorption or pressure erosion of inflamed mucosa (Fig 9) (11). If there is destruction of the outer cortex of the mastoid, a subperiosteal abscess may develop within the overlying soft tissues and the patient may present with a postauricular collection (Fig 10). A Bezold abscess develops when there is bone defect at the mastoid tip medial to the insertion of the
posterior belly of the digastric muscle (digastric notch) (Fig 10). Inflammatory changes are then directed inferiorly deep to the sternocleidomastoid muscle by the fascial planes. In young children, the mastoid tip is usually not aerated; as a result, Bezold abscess in young children is uncommon (12).

When present, petrous apex air cells are connected to the mastoid air cells via numerous supra- and infralabyrinthine air channels that facilitate spread of infection from the mastoid air cells and middle ear to the petrous apex, resulting in petrous apicitis. *Streptococcus pneumonia*, *Haemophilus influenza*, and *Staphylococcus aureus* are the most common causative organisms (13). Petrous apicitis occurs in the setting of a pneumatized petrous apex (present in 30% of the population). It is characterized by septal and cortical destruction, osteitis, and adjacent meningeal inflammation. Because of the close proximity of the fifth and sixth nerves to the petrous apex, patients can develop sixth nerve palsy and deep retroorbital pain in the V1 distribution in association with otomastoiditis and petrous apicitis. This constellation of findings is termed Gradenigo syndrome (14). Imaging findings include opacification of an aerated petrous apex with fluid levels and destruction of the inner septations or the cortex (Fig 11). On MR images, meningeal enhancement is better appreciated. Osteomyelitis can occur in a nonpneumatized petrous apex by means of direct medial extension of necrotizing otitis externa or by retrograde spread of thrombophlebitis along the venous plexus of the petrous carotid canal (15).

Intracranial complications of acute mastoiditis include dural venous sinus thrombosis, most commonly the sigmoid and transverse sinuses (Fig 10), epidural abscess, subdural empyema, meningitis, and brain abscess.

**Chronic otitis media.**—Chronic inflammation of the middle ear is known as chronic otitis media, or chronic otomastoiditis if there is mastoid involvement. It is characterized by a variety of signs, symptoms, and physical findings that result from long-term damage to the middle ear by infection and inflammation. Common mechanisms for developing chronic otitis media include underlying eustachian tube dysfunction and tympanic membrane perforation. Some of the important sequelae of chronic otitis media that may be seen at imaging include middle ear effusion, granulation tissue, cholesterol granuloma, and cholesteatoma. On CT images, all these lesions may have very similar appearances. Granulation tissue is a sequela of inflammation of the middle ear and mastoid. It encases middle ear structures but does not destroy or displace them, and there is no mass effect. Granulation tissue enhances intensely since it is highly vascularized. On gadolinium-enhanced MR images, this is readily visible. However, enhancement may be difficult to appreciate on CT images.

**Cholesterol granuloma.**—In the setting of eustachian tube dysfunction, there may be build-up of negative pressure or vacuum phenomenon in the middle ear cavity, leading to mucosal edema and rupture of blood vessels (13). The breakdown of erythrocytes and tissue elements release cholesterol, which incites a foreign body giant cell reaction leading to formation of a chronic
granuloma, termed cholesterol granuloma (16). This lesion is also referred to as a cholesterol cyst, chocolate cyst, or blue-domed cyst. Common locations for this entity include the middle ear and the petrous apex; it can also rarely occur in a mastoidectomy cavity. In the middle ear, patients will present with a blue tympanic membrane, hemotympanum, or conductive hearing loss. In the petrous apex, these lesions may be asymptomatic or patients may have nonspecific symptoms such as vertigo, dizziness, or cranial neuropathies relating to cranial nerves V, VI, VII, and VIII. In the petrous apex, an expansile lesion with imperceptible bone margins may be seen on CT scans (Fig 12a). On MR images, the characteristic finding is the presence of intrinsic T1 shortening (hyperintensity) due to the presence of blood products (Fig 12b). On T2-weighted MR images, signal intensity is usually heterogeneous hyperintense (Fig 12c) (17). A hypointense rim on T2-weighted images may be present, which is believed to represent hemosiderin or a preserved rim of bone (18). Cholesterol granulomas do not enhance. An important differential diagnosis for this entity is entrapped simple fluid or a petrous apex effusion. However, although an effusion may mimic a cholesterol granuloma by way of MR signal characteristics, it does not cause expansion or destruction of the petrous apex air cells, a distinction that is best evaluated with CT (Fig 13).

Figure 12: Images of cholesterol granuloma. (a) Axial CT image in a 37-year-old woman with diplopia and headache demonstrates an expansile mass in the left petrous apex with imperceptible bone margins (arrows). (b) Axial T1-weighted MR image in a different patient demonstrates characteristic T1 hyperintensity in an expansile petrous apex lesion (>). (c) On an axial T2-weighted image (in same patient as in b), the lesion is heterogeneously hyperintense, with a rim of hypointensity (arrows).

Figure 13: Axial CT image of petrous apex effusion in a 36-year-old man with right-sided sensorineural hearing loss demonstrates opacification of the pneumatized right petrous apex (>). It does not have expansile margins, distinguishing it from a petrous apex mucocele or a cholesterol granuloma. The internal septations and cortex are intact, distinguishing it from petrous apicitis.
the tympanic membrane. Congenital cholesteatoma is postulated to arise from epithelial rests in the middle ear in a child with no previous history of otorrhea, tympanic membrane perforation, or otologic procedures. In the petrous apex, cholesteatomas can be either acquired or congenital. Both congenital and acquired cholesteatomas can be seen as a pearly white lesion at otoscopy. Clinically, patients with acquired cholesteatomas present with chronic foul-smelling otorrhea, tympanic membrane perforation, and a retraction pocket in the pars flaccida. Congenital cholesteatoma may be an incidental finding in an asymptomatic patient, or the patient may have conductive hearing loss.

CT is the imaging mainstay of temporal bone inflammation (21). Acquired pars flaccida cholesteatoma is seen as a rounded expansile lobulated lesion in the Prussak space eroding the scutum, with medial displacement and erosion of the ossicles (21) (Fig 14). The lesion may extend superiorly into the attic, and to the mastoid air cells via the aditus ad antrum. In contrast, pars tensa cholesteatoma tends to occur medial to the ossicles and displace them laterally (21) (Fig 13). Congenital cholesteatoma is commonly located in the anterior superior quadrant of the middle ear cavity just above the opening of the eustachian tube.

Some common complications of cholesteatoma include erosion of the roof of the middle ear ( tegmen tympani ), with or without formation of a meningoencephalocele; erosion of the bony facial canal in its tympanic or mastoid portion; and erosion of the bony covering over the lateral semicircular canal, which may result in formation of a communication between the middle ear and inner ear, also known as a labyrinthine or perilymphatic fistula. Mural cholesteatomas drain their cystic contents through the tympanic membrane into the EAC, with only the matrix lining remaining (22). This results in a characteristic “automastoidectomy” appearance, with an extensive bone destructive process in the middle ear/mastoid resembling a surgical mastoidectomy cavity (Fig 16). At MR imaging, a cholesteatoma is hypointense on T1-weighted images, is mildly hyperintense on T2-weighted images, and does not enhance (Fig 17). Reduced diffusivity may be seen on diffusion-weighted (DW) images in cholesteatomas. Non–echo-planar turbo spin-echo–based DW imaging techniques can achieve thinner sections (2 mm) with less susceptibility artifact at the air-bone interface when compared with echo-planar imaging sequences. MR imaging is also very useful in evaluating for complications such as meningoencephaloceles resulting from tegmen erosion. MR imaging can also help differentiate a recurrent cholesteatoma from cholesterol granuloma, granulation tissue, or scar (Table 1). The sensitivity of DW imaging for the detection of cholesteatomas decreases in the postoperative setting, although the specificity remains high (23).

Facial Nerve

Bell palsy.—Bell palsy refers to facial nerve paralysis with no identifiable cause. Viral inflammation, possibly prior herpes simplex virus infection, has been implicated. It is characterized by very rapid onset of facial nerve paralysis, which progresses over a few hours to up to 3 weeks. In 80% of patients symptoms improve spontaneously; if symptoms persist for more than 3 weeks, more ominous causes such as neoplasm should be investigated, and investigation should include the use of MR imaging. On imaging studies, there is enhancement of the canalicular, labyrinthine, and geniculate portions of the facial nerve (Fig 18) (23,24).

Ramsay Hunt syndrome.—Ramsay Hunt syndrome is also referred to as herpes zoster oticus. It is thought to represent reactivation of varicella zoster virus latent in the geniculate ganglion. Reactivation may occur in the setting of immunocompromised states, systemic diseases, or aging. Clinically, the patient...
presents initially with burning pain in the ear; this is followed 1 day to 4 days later with vesicular eruption, facial paralysis, hearing loss, and/or vertigo. On MR images, there is enhancement of cranial nerves VII and VIII, the labyrinth, and/or the pontine facial nucleus (25).

Inner Ear
Inner ear inflammation or labyrinthitis can occur as a result of infections (viral or bacterial) or may be autoimmune or posttraumatic. Labyrinthitis ossificans is the late stage of labyrinthitis, in which there is pathologic ossification of spaces within the lumen of the bony labyrinth (comprised of the cochlea and the vestibular system). The most common region of cochlear ossification occurs in the scala tympani of the basal turn, and the most extensive cases are seen as a complication of meningitis. The evolution of labyrinthitis is characterized by three stages: acute, fibrous, and ossification (labyrinthitis ossificans) (26). In the acute stage, enhancement of the inner ear is noted on MR images, but the CT scan may appear normal. In the intermediate fibrous stage of labyrinthitis, there is loss of fluid signal intensity on heavily T2-weighted sequence images (e.g., three-dimensional constructive interference in steady state [CISS]) or fast imaging employing steady-state acquisition [FIESTA]), while the CT scan may still appear normal. In the late ossific stage, one sees replacement of the normal cochlea, vestibule, and/or semicircular canals by bone attenuation on CT scans (Fig 19). CT lags behind MR imaging in the detection of labyrinthitis ossificans.

Figure 17: MR images of cholesteatoma in a 15-year-old male patient with a history of draining ear. (a) Axial T2-weighted image demonstrates a heterogeneously hyperintense lesion in the left middle ear/mastoid (*). (b) Axial gadolinium-enhanced T1-weighted image demonstrates the lesion to be nonenhancing (*). (c) Coronal DW image demonstrates reduced diffusivity within the lesion (arrow).

<table>
<thead>
<tr>
<th>Tissue Type</th>
<th>T1-weighted MR Imaging</th>
<th>T2-weighted MR Imaging</th>
<th>Enhancement</th>
<th>Diffusion Restriction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cholesteatoma</td>
<td>Hypointense</td>
<td>Hyperintense</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Granulation tissue</td>
<td>Hypointense</td>
<td>Hyperintense</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Scar</td>
<td>Hypointense/intermediate</td>
<td>Hypointense/intermediate</td>
<td>No/delayed</td>
<td>No</td>
</tr>
<tr>
<td>Cholesterol granuloma</td>
<td>Hyperintense</td>
<td>Hyperintense/intermediate</td>
<td>No</td>
<td>Variable</td>
</tr>
</tbody>
</table>

Tumors of the Temporal Bone
To assess a tumor in the temporal bone region, it is helpful to localize the lesion to one of the following areas, each of which houses a unique set of pathologic conditions: (a) IAC/CPA, (b) middle ear, (c) EAC and mastoid, and (d) petrous apex. We will consider each area separately in the following sections.

IAC/CPA
The most common masses in this area are vestibular schwannomas, meningiomas, epidermoids, and nonvestibular posterior fossa schwannomas (such as trigeminal, facial, and glossophysical). Less frequently seen are arachnoid cysts, lipomas, dermoids, and malignancies such as lymphoma, melanoma, and metastases. In addition, tumors related to the petrous bone (e.g., chondrosarcoma) and brain (e.g., glioma) may extend into the IAC/CPA.

Vestibular schwannoma.—The most common tumor in the IAC/CPA is the vestibular schwannoma, which accounts for 60%–90% of all tumors in this region (27–29). It is also thought to be the most common intracranial nerve sheath tumor (30). Incidence is highest in the 5th through 7th decades of life (31), although they commonly present in the first 2 decades in the setting of neurofibromatosis type II; the
phakomatosis associated with bilateral vestibular schwannomas, multiple schwannomas of other cranial nerve origin, meningiomas, and ependymomas in the brain and spine (32). Patients present with sensorineural hearing loss, tinnitus, disorientation, and/or decreased speech discrimination, secondary to pressure by the tumor on the cochlear and vestibular divisions of cranial nerve VIII (33); facial nerve manifestations are relatively uncommon (33,34). Larger tumors may compress the trigeminal nerve (causing facial numbness), the lower cranial nerves (causing cerebellar signs), or the fourth ventricle (causing hydrocephalus).

As vestibular schwannomas enlarge, they may expand medially into the CPA and laterally toward the fundus and/or into the cochlear aperture. When the former occurs, the porus may be expanded (Fig 20), and the spherical cisternal component usually forms an acute angle with the petrous bone at the bone-tumor interface (35). When a vestibular schwannoma extends into the cochlear aperture, the tumor is termed impacted; this finding correlates with a decreased chance of hearing preservation following surgery (36).

Most vestibular schwannomas are slow growing, with growth rates ranging from 0.2 mm to a few millimeters per year, though some may enlarge by more than 10 mm per year. Whenever feasible, surgical resection allows eradication of the tumor (37). There are three main approaches—retrosigmoid/suboccipital, middle fossa, and translabyrinthine (38–40). The latter leads to complete hearing sacrifice and is therefore generally reserved for patients with poor hearing or large tumors. Alternatively, stereotactic radiosurgery may be considered, especially in high-risk patients, those with bilateral tumors, and those with residual tumors following initial treatment (41).

On CT images, most vestibular schwannomas are isodense with the cerebellum and are diffuse without contrast material enhancement (28). However, if the tumor is large and causes expansion of the porus acusticus, this may be readily seen on CT bone window images, with the porus on the affected side asymmetrically wider. Calcification and hemorrhage are rare unless the tumor has been treated (42). Enhancement is usually avid and homogeneous (43), but this may be difficult to detect on CT images, especially if the tumor is small.

On MR images, schwannomas are usually iso- or mildly hypointense to brain parenchyma and hyperintense to cerebrospinal fluid (CSF) on T1-weighted images, mildly hyperintense to brain parenchyma and iso- to hypointense to CSF on T2-weighted images, and enhance avidly. Large tumors may be heterogeneous, with intra- or extrameatal cystic components (28), and may deform and displace the brainstem, causing parenchymal edema, and compress the fourth ventricle (Fig 21). Heavily T2-weighted sequences are helpful for outlining the tumor, as most structures except CSF appear quite dark. CSF thus provides natural contrast around the dark tumor mass (Fig 21). Heavily T2-weighted sequences may also reveal decreased signal intensity of the labyrinthine fluid ipsilateral to the tumor, thought to be related to higher protein content in the fluid (44); this may also be seen as increased signal intensity on fluid-attenuated inversion recovery images.

Meningiomas.—In contrast to vestibular schwannomas, meningiomas are often eccentric to the porus acusticus, centered at the CPA; when they do extend into the IAC, they seldom expand the porus or the IAC (Fig 22). Meningiomas may extend into the middle cranial fossa by means of herniation, growth through the tentorium, or growth through the temporal bone (45,46). They may also extend into the middle ear and cavernous sinus (Fig 22b). As mentioned above, meningiomas tend to be broad-based along the posterior petrous wall, forming an obtuse angle at the bone-tumor interface, appearing either hemispherical or plaquelike (45). Dural
Enhancement extending outward from the margins of the tumor is often seen. On CT images, meningiomas are hyper- or isodense to gray matter on T1-weighted images and are iso- or hypointense to gray matter on T2-weighted images. Enhancement is usually homogeneous, unless there are calcific or cystic foci. On MR images, one might also observe surface flow voids (from marginal pial vessels) and surface CSF clefts (47–49). Treatment is by means of surgical resection, with radiation as an adjunctive or alternative method (50). However, local recurrence is not uncommon (51).

Epidermoid.—Congenital epidermoid cysts (congenital cholesteatomas) are the third most common mass in the CPA (28,29); they consist of stratified squamous epithelial linings surrounding desquamated keratin. They are not true neoplasms and are described in detail under the earlier section, “Inflammatory Lesions.” On CT images, epidermoids are similar to CSF in attenuation; if they are large, smooth remodeling of the adjacent petrous bone may be observed secondary to long-term pressure erosion. At MR imaging, they are iso- or slightly hyperintense to CSF on T1-weighted images and mildly hyperintense to CSF on T2-weighted images (Fig 23) (47). They do not enhance. On fluid-attenuated inversion recovery images, signal intensity remains hyperintense, and on DW images, an epidermoid demonstrates reduced diffusivity because of its solid nature (52). This is in contradistinction to an arachnoid cyst, which does not demonstrate diffusion reduction.

Middle Ear
When a soft-tissue mass is seen in the middle ear, a vascular structure must be excluded. This includes a persistent stapedial artery, a laterally placed or aberrant carotid artery, a carotid artery aneurysm, and an exposed dehiscent jugular bulb. True neoplasms include paraganglioma (most common); facial nerve lesions extending into the middle ear, such as schwannomas and geniculate region hemangiomas, choristomas, and perineural spread of tumor; meningiomas; adenomatous tumor of the mixed pattern type; and malignancies such as carcinomas and metastases (rare).

Most choristomas in the middle ear consist of salivary gland tissue (53,54), and they may be associated with incudostapedial or tympanic facial nerve abnormalities (55).

Adenomatous tumor of the mixed pattern type is also called middle ear adenoma (36). This is a benign tumor that does not demonstrate bone invasion. At CT, the only finding may be an opacity in the middle ear space, and this tumor may therefore be difficult to distinguish from otitis media (Fig 24). At MR imaging, it is hypointense on T1-
Radiology: Volume 269: Number 1—October 2013

Paraganglioma.—Tumors of paraganglia cells are termed paragangliomas, but in the head and neck region, the term glomus tumor has been used ever since discovery of the paraganglion located in the adventitia of the dome of the jugular bulb by Guild in 1941, who named it glomus jugulare (57). Paraganglioma is the second most common tumor involving the temporal bone (28). Glomus tumors in the temporal bone region arise from the jugulotympanic paraganglia along the nerves of Jacobson and Arnold (tympanic branch of the glossopharyngeal nerve and auricular branch of the vagus nerve, respectively). Glomus tympanicum refers to those confined to the tympanic cavity (arising from paranglia cells along the nerve of Jacobson) and glomus jugulare refers to those tumors involving the jugular bulb and the base of skull (arising from paraganglia cells along the nerve of Jacobson or Arnold). Glomus jugulotympanicum has components in both the middle ear and the jugular foramen.

Jugulotympanic paragangliomas occur three to four times more often in women than in men. Patients typically present between the 4th and 6th decades of life. Up to 10% of patients may have multiple paragangliomas, so it behooves the radiologist to continue the search for additional lesions when interpreting a study, whether in the temporal bones, skull base, or neck (58). Paragangliomas are seen in a familial form in multiple endocrine neoplasia (MEN) 2a and 2b. These tumors are slow growing and locally infiltrating, growing along planes of least resistance in existing pathways in the temporal bone, and they rarely metastasize. Most patients present with otologic concerns (conductive hearing loss, pulsatile tinnitus, or a retrotympanic mass) (58,59).

Paragangliomas are highly vascular and therefore enhance avidly. Glomus tympanicum paragangliomas are found against the cochlear promontory and are usually small at presentation since they cause otologic symptoms early on (Fig 25) (60,61). It is important to inspect the margins of the jugular foramen to exclude a glomus jugulotympanicum; localized lytic or permeative bone destruction is characteristic of a glomus tumor involving the jugular foramen, and is well seen on CT scans (Fig 26a) (62).

The soft-tissue mass itself is better appreciated on MR images, showing avid enhancement. In larger tumors, a “salt-and-pepper” appearance may be visible on T1- and T2-weighted MR images (Fig 26b) (63). Multiple internal tumoral vessels manifest as serpentine and arborizing flow voids.

At angiography, enlarged feeding arteries and rapidly draining veins may be seen (64). In tympanicum tumors, blood loss is minimal, and preoperative embolization is not indicated. However, preoperative angiography and embolization of larger jugulotympanicum tumors are commonly performed to help reduce intraoperative blood loss.

Indium 111 octreotide is useful for detecting multicentric, metastatic, or recurrent tumors. The sensitivity of iodine 123-metaiodobenzylguanidine for head and neck paragangliomas is lower (65–67).

EAC and Mastoid

Tumors in the EAC and mastoid region are often malignant, with squamous cell carcinoma being by far the most common (27). Patients with EAC squamous cell carcinoma frequently have a long history of chronic ear infections. There is aggressive bone destruction, and there may be invasion of surrounding soft tissues including intracranial, infratemporal, middle ear, parotid, carotid, and temporomandibular joint involvement (Fig 27) (68–70). Other malignancies such as basal cell carcinoma, melanoma, lymphoma, myeloma, metastases, chondrosarcoma, and osteosarcoma occur much less frequently.

Aggressive bone destruction may be seen with some benign processes and should be considered in the differential diagnosis. These include granulomatous diseases such as Langerhans cell histiocytosis (Fig 28), tuberculosis, and Wegener granulomatosis. Aggressive infections such as malignant otitis externa and radiation necrosis are additional considerations.

Petrosus Apex

True neoplasms in the petrous apex include chondrosarcoma, chordoma, osteosarcoma, and meningioma. Myeloma, lymphoma, and metastases may also occur. The petrous area may be secondarily involved by regional tumors such as trigeminal schwannoma, jugular paraganglioma, and nasopharyngeal carcinoma. The latter is usually seen along...
the petrooccipital fissure, superior to the fossa of Rosenmüller. Unlike lesions in the IAC and middle ear, lesions in the petrous apex usually reach considerable size before causing symptoms.

Figure 26: Images of glomus jugulotympanicum. (a) Axial CT image demonstrates the moth-eaten, permeative bone destruction around the tumor. On this image through the right temporal bone, one can appreciate destruction of the margins of the jugular foramen (∗), the caroticojugular spine (arrowhead), and the carotid canal (small arrow). Note the tumor extending into the middle ear (large arrow). (b) Axial contrast-enhanced T1-weighted MR image shows the large enhancing tumor with internal flow voids (arrows), creating the so-called salt-and-pepper appearance. This patient initially presented at age 56 years with right ear fullness, hearing loss, and audible whooshing sound that correlated with heart beat. Over the next 4 years, the tumor gradually enlarged, with new right true vocal cord paralysis indicating new involvement of the vagus nerve. He underwent proton beam radiation therapy.

Figure 27: Axial CT image of squamous cell carcinoma of the EAC with extension to the mastoid. Patient was a 64-year-old woman with recurrent and persistent left ear infection not improved with antibiotics; biopsy revealed the diagnosis. CT image demonstrates extensive bone destruction involving the EAC margin and mastoid (black arrowheads), extending to the petrous apex (large arrow). The dense otic capsule is relatively spared (white arrowhead). Abnormal soft tissue fills the EAC, middle ear, and mastoid, extending into the posterior cranial fossa (small arrows). The auricle is also thickened, with ulcerated margins.

Figure 28: Axial contrast-enhanced T1-weighted MR image of Langherhans cell histiocytosis shows solid enhancing tissue involving the EAC and mastoid (black arrows). There is also involvement of the middle ear (white arrow) and sigmoid sinus (arrowhead). Patient was a 2-year-old boy with otitis media starting at 8 months of age, now with pus and bloody discharge and diffuse swelling in the ear region. Biopsy revealed Langherhans cell histiocytosis.

Figure 29: Axial contrast-enhanced T1-weighted MR image of petrous apex chondrosarcoma. Image shows the tumor mass in the petrous apex causing displacement of the petrous carotid artery (arrow). It was hypointense on T2-weighted images (not shown) and enhances avidly.

notochordal remnants and are typically seen in the midline, may be found off midline as well and may mimic a chordoma radiologically (71,72). Additionally, the chondroid subtype of chordoma may be difficult to distinguish from the myxoid variant of chondrosarcoma pathologically (73).

At CT, these tumors enhance mildly to moderately, may contain calcifications, and cause surrounding bone destruction (71,74,75). At MR imaging, the lesions are hypointense on T1-weighted images and hyperintense on T2-weighted images, although their appearance may be quite heterogeneous (71,75,76). Enhancement is marked but may also be heterogeneous (Fig 29).

Endolymphatic sac tumor.—Endolymphatic sac tumor is a locally invasive papillary cystadenomatous tumor (56,77,78). Most cases are sporadic, although a minority may be seen in the setting of von Hippel-Lindau disease (7%,79). The tumor causes local bone destruction in the region of the vestibular aqueduct in the retrolabyrinthine petrous bone.

At CT, the bone invaded by tumor has a moth-eaten, lytic appearance, with intratumoral bone spicules often seen (Fig 30a). At MR imaging, areas of intrinsic T1 shortening are commonly seen, reflecting blood products (Fig 30b). Low-signal-intensity areas may
represent hemosiderin. Enhancement is usually heterogeneous (80).

**Facial Nerve**

The most common benign tumors of the facial nerve are schwannomas and hemangiomas. The most important consideration in the malignant category is perineural spread of tumor along the facial nerve.

**Facial schwannoma.**—Facial nerve schwannomas can involve any segment of the nerve and may span multiple segments (81). The geniculate ganglion is frequently involved (82). Of note, only a minority of patients initially present with facial palsy; many have no facial nerve symptoms at all (82). Facial nerve schwannomas in the IAC/CPA typically manifest as sensorineural hearing loss, presumably because the thinly myelinated sensory fibers of cranial nerve VIII are more sensitive to compressive effects by the tumor than are the thickly myelinated motor fibers of the facial nerve (82). Facial schwannomas in the geniculate region can grow into the middle cranial fossa. Those along the tympanic segment may bulge into the middle ear and compress the ossicular chain, causing conductive hearing loss. Those in the mastoid segment are more likely to present with facial palsy, owing to the surrounding narrow bony canal exerting pressure on the growing tumor (82). Those in the parotid segment present as painless neck masses (83).

At CT, the tumor may be seen to cause expansile smooth remodeling of the surrounding bony canal. At MR imaging, an expansile, enhancing mass can be appreciated (Fig 31). It may have a tubular appearance, may span multiple segments, or may have a sausage-link appearance, with areas that are relatively narrower between “links” that are expansile. In the IAC, they may be indistinguishable from vestibular schwannomas (84,85). In the geniculate ganglion region, differential diagnostic considerations include hemangioma and meningioma, although these lesions tend to demonstrate intratumoral bone spicules.

**Hemangioma.**—Hemangiomas may be seen along the intratemporal course of the facial nerve, most often in the region of the geniculate ganglion, followed by the IAC, and are seen least often at the posterior genu. These tumors often grow among bone trabeculae and may form bone; in these cases the term *ossifying hemangioma* is sometimes used (86). Recent literature suggests that these lesions are, in fact, venous malformations (87). These lesions characteristically have an expansile honeycomb appearance and may demonstrate intratumoral bone spicules on CT images (Fig 32a) (88). These may be difficult to distinguish from meningiomas with intraosseous involvement. On MR images, there may be heterogeneous signal intensities and avid enhancement (Fig 32b).

**Perineural spread of tumor.**—With segmental facial nerve thickening and enhancement, an important malignant process to consider is perineural spread of tumor (Fig 33a). The source of malignancy is usually the parotid gland, for example, adenoid cystic carcinoma (Fig 33b) or mucoepidermoid carcinoma, or a nearby skin malignancy that secondarily invades or metastasizes to the parotid gland (89,90). Tumor burden can vary along the course of the nerve, resulting in varying degrees of thickening.
Figure 32: Images of hemangioma in the geniculate region in a 48-year-old woman with new right facial fasciculation in the region of the right eye, progressing to the lower face, and then global weakness of the right face with synkinesis. (a) Axial CT image demonstrates an expansile mass with intratumoral bone spicules (arrow) in the region of the geniculate ganglion. (b) On contrast-enhanced axial T1-weighted MR image, avid enhancement can be seen (arrow).

Figure 33: Axial T1-weighted MR images show perineural spread of tumor along the facial nerve in a 58-year-old woman with transient lower lip numbness and weakness. (a) Contrast-enhanced image shows expansile enhancement involving the geniculate ganglion and tympanic segment of the facial nerve (arrow). The mastoid segment is involved as well (not shown). (b) Nonenhanced image more caudally reveals an adenoid cystic carcinoma in the right parotid gland that extends to the stylomastoid foramen (arrow) as the source; it was subsequently resected.

Table 2

<table>
<thead>
<tr>
<th>Location</th>
<th>Inflammatory</th>
<th>Neoplastic</th>
</tr>
</thead>
<tbody>
<tr>
<td>EAC/mastoid</td>
<td>Cholesteatoma; keratitis obturans; malignant otitis externa</td>
<td>Squamous cell carcinoma; other malignancies</td>
</tr>
<tr>
<td>Middle ear</td>
<td>Otitis media; cholesterol granuloma; congenital cholesteatoma</td>
<td>Glomus tympanicum/jugulotympanicum; adenoma; choristoma</td>
</tr>
<tr>
<td>Facial nerve</td>
<td>Bell palsy; Ramsay-Hunt</td>
<td>Schwannoma; hemangioma (geniculate); meningioma (geniculate); perineural spread of tumor</td>
</tr>
<tr>
<td>Petrous apex</td>
<td>Petrous apicitis; cholesterol granuloma; cholesteatoma</td>
<td>Meningioma; chordrosarcoma; osteosarcoma; chordoma; endolymphatic sac tumor; metastasis; direct extension from nasopharyngeal carcinoma</td>
</tr>
</tbody>
</table>

and enhancement; “skip lesions” may be seen where there are areas of uninvolved nerve between abnormal segments (91). It is important to note that the original tumor at the primary site could be absent if it has been treated; this does not preclude the diagnostic consideration of active perineural spread detectable at imaging.

Conclusion

In this review, we highlighted the major anatomic landmarks in the temporal bone one should be familiar with to be able to describe normal and pathologic findings and provide an interpretation. We described the most frequently encountered pathologic conditions in various locations of the temporal bone, with a focus on inflammatory and neoplastic processes (Table 2). CT and MR are the major tools for imaging evaluation. CT offers excellent delineation of soft-tissue abnormalities against a background of air (middle ear cavity, EAC, mastoid air cells) and allows assessment of bone changes and/or involvement. MR imaging offers more detailed characterization of soft tissue and fluid, allows better visualization of enhancement in small structures, and aids in distinction between soft tissues of different nature (eg, granulation tissue versus tumor).

Disclosures of Conflicts of Interest: A.F.J. No relevant conflicts of interest to disclose. D.T.G. No relevant conflicts of interest to disclose. G.M. No relevant conflicts of interest to disclose.

References

47. Gentry LR, Jacoby CG, Tarski PA, Houston LW, Strother CM, Sackett JF. Cerebellopontine angle–petromastoid mass lesions: compar-
59. 57. 56. 55. 54. 52. 50. 49.
Radiology:

61. 60. 59. 58. 57. 56. 55.

63. 62. 61. 60. 59. 58. 57. 56.

65. 64. 63. 62. 61. 60. 59. 58.

67. 66. 65. 64. 63. 62. 61. 60.

69. 68. 67. 66. 65. 64. 63. 62.

71. 70. 69. 68. 67. 66. 65. 64.

73. 72. 71. 70. 69. 68. 67. 66.

75. 74. 73. 72. 71. 70. 69. 68.

77. 76. 75. 74. 73. 72. 71. 70.

79. 78. 77. 76. 75. 74. 73. 72.

81. 80. 79. 78. 77. 76. 75. 74.

83. 82. 81. 80. 79. 78. 77. 76.

85. 84. 83. 82. 81. 80. 79. 78.

87. 86. 85. 84. 83. 82. 81. 80.

89. 88. 87. 86. 85. 84. 83. 82.

91. 90. 89. 88. 87. 86. 85. 84.