Role of Radiologic Imaging in Otosclerosis

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Abstract
Purpose of Review To review the role of imaging in otosclerosis with an emphasis on pre- and post-operative imaging evaluation.
Recent Findings Pre-operative CT imaging can help define the extent of bone involvement in otosclerosis and may help avoid surgical complications due to variant anatomy or unsuspected alternative causes of conductive hearing loss. In patients with recurrent hearing loss after surgery, CT imaging can clarify prosthesis position and re-assess anatomy.
Summary CT imaging complements otologic exam and audiometry findings in patients with suspected otosclerosis, for pre-operative planning, and post-operative assessment for patients with recurrent symptoms.

Keywords Computed tomography (CT) · Otosclerosis · Semicircular canal dehiscence · Otodystrophy · Stapes prosthesis · Prosthesis migration

Introduction
The role of imaging in otosclerosis has expanded over the last several decades since Valvassori first reported his findings using polytomography [1, 2]. While otosclerosis can be diagnosed with history, otologic exam findings, and audiologic testing, and confirmed at surgery [3], access to imaging has become more widespread to complement diagnosis, pre-surgical planning, and evaluation of recurrent symptoms [3, 4].

Non-contrast temporal bone CT is considered the gold standard for otosclerosis imaging [5, 6]. With current technology, slice thickness of 0.5–0.625 mm is routinely achievable, allowing for thin multiplanar reformat capable of detecting abnormalities up to 1 mm in size [7]. Each temporal bone is independently reconstructed with a smaller field of view for imaging review. Using newer multidetector technology, the sensitivity and specificity of CT for evaluation of otosclerosis have been reported as high as 95% and 100%, respectively [3, 8]. Other modalities including magnetic resonance imaging (MRI), single-photon emission computed tomography (SPECT), and cone-beam computed tomography (CBCT) are less well established but may also play a role in otosclerosis imaging [3, 9, 10].

Imaging Findings of Otosclerosis
To best understand the imaging features of otosclerosis, it is helpful to understand the embryologic development of the otic capsule and the histopathology of otosclerosis. The bony labyrinth is a hard and uniformly dense bone with a distinctly different texture than the surrounding parts of the petrous pyramid (except for the modiolus). The otic capsule develops from a preexisting cartilage model and is thus endochondral bone. The inner endosteal layer and the middle (thicker) endochondral layer of the otic capsule form in mid-fetal life and remain relatively inert and unchanged throughout life [11, 12]. Otosclerosis primarily involves the endochondral layer. Otosclerosis is a primary focal spongifying process which alters the ivory-hard avascular endochondral bone of the otic capsule, and irregular, highly vascular (“Schwartz sign”), foci of Haversian bone tissue develops [11, 13]. Histologically, otospongiosis is characterized by enlargement of the perivascular spaces, osteoclastic bone resorption, and new immature bone formation [13]. In the
later stages of the disease, the lamellar bone may be deposited [13]. In 80–90% of otospongiosis, a focus is found just in front of the oval window at the fissula (depression) ante fenestram, the most common site of involvement. The fissula ante fenestram and the fossula post fenestram develop as outpouchings from the vestibule anterior and posterior to the oval window, respectively. They are composed of a strip of altered precartilage which gradually undergoes necrosis and eventually is replaced by connective tissue centrally. As the fissula ante fenestram develops, it enlarges and approaches the anterior aspect of the stapedial area. It is common to have variations in the size, shape, and amount of cartilage around the fissula ante fenestram [14]. The fossula post fenestram develops similarly to the fissula ante fenestram and approaches the posterior aspect of the stapedial area. Cartilaginous changes are less common in the fossula post fenestram compared to the fissula ante fenestram, present in only 5% of temporal bone specimen [15].

On CT, the normal otic capsule appears uniformly dense, unlike the surrounding petrous Haversian bone which contains marrow. The imaging findings in otosclerosis largely depend on the phase of the disease and the location of involvement (Fig. 1). Classically otosclerosis is described in two phases, an active otospongiotic phase and an inactive mature otosclerotic phase [16–20]. In the active phase, the earliest CT finding is hypodensity at the fissula ante fenestram [16, 17]. As otosclerosis progresses, disease may extend beyond the fissula ante fenestram to the peri-cochlear region, described as retrofenestral disease. Active retrofenestral disease is predominantly characterized by hypodensity about the cochlea, and a distinctive “double ring” sign can sometimes be seen [18, 19, 20]. In the inactive otosclerotic phase, the findings can be more difficult to appreciate. The degree of sclerosis can be variable, and the only finding may be subtle contour abnormality or scalloping [16, 17, 21•]. CT has been shown to be less sensitive for the evaluation of inactive otosclerotic disease, although active and inactive disease is often found to coexist. It is important to carefully assess for bilateral involvement, as the appearance and symptoms may be asymmetric.

Several CT classification and grading systems have been proposed for evaluation of otosclerosis [6, 22, 23], largely based on active versus inactive disease and location of involvement. These classification systems have shown variable success in prediction of disease severity, prognosis, and likelihood for successful surgery, but currently, no imaging-based classification system has been universally accepted [22, 23]. Similarly, authors have proposed different checklist approaches to reviewing and reporting imaging of otosclerosis [7]. Preliminary investigations suggest that quantitative CT attenuation measurements may serve as an additional complimentary method to detect otosclerosis [2, 24].

MRI findings in the evaluation of otosclerosis are less established. Most patients with suspected otosclerosis present with conductive hearing loss are initially imaged with CT; however, for patients who present with mixed hearing loss, sensorineural hearing loss, or tinnitus, otosclerosis may not be suspected, and these patients may be first imaged by MRI. The normal otic capsule appears hypointense on all MRI sequences, unlike petrous Haversian bone. Additionally, the normal otic capsule is comprised of avascular endochondral bone which does not enhance. MRI findings of otosclerosis are subtle and easily overlooked. Classic antefenestral involvement is less commonly identified on MRI, whereas retrofenestral involvement is more conspicuous and is characterized by intermediate T1 signal and mild-moderate enhancement [9, 25, 26]. Occasionally, curvilinear increased T2 signal in the peri-cochlear region can also be seen [26]. This is not a finding exclusive to otosclerosis, as retrofenestral enhancement is also present in the acute/subacute phase of labyrinthitis ossificans [27].

Currently, there is limited data for the role of SPECT and CBCT in the evaluation of otosclerosis [10, 28]. A study assessing diphosphonate SPECT for the evaluation of otosclerosis demonstrated that increased temporal bone uptake on SPECT was sensitive for otosclerosis and correlated with temporal bone CT findings, although the spatial resolution of SPECT was limited compared to CT [10]. CBCT is more widely used for dental evaluation but can identify bone changes of otosclerosis [29].
Clinical and Imaging Mimics of Otosclerosis

Many alternative causes of conductive hearing loss can be excluded based on history, otoscopy, and audiometric evaluation. However, causes of conductive hearing loss may occasionally be misdiagnosed as otosclerosis, leading to surgical complication or failed surgery. Occasionally chronic middle ear infection, tympanosclerosis, and cholesteatoma (particularly congenital cholesteatoma) in the antrum, fossa incudis, and petrous juxta labyrinth are difficult to detect by otoscopy alone. Small masses such as menigioma, schwannoma, and metastasis can mimic clinical findings of otosclerosis [30]. Imaging may be of particular benefit in cases with mixed or sensorineural hearing loss. It is also important to note if patients have a systemic disease which may have complications mimicking otosclerosis. Rare cases include symptoms related to extramedullary hematopoiesis, inflammatory processes such as chronic inflammatory demyelinating polyneuropathy (CIDP) involving the facial nerve and chorda tympani, among other case reports. The combination of CT and MRI has proved the most valuable for detection of unsuspected cholesteatoma and intratympanic masses in patients with clinically suspected stapedial cochlear otosclerosis [31, 32].

Conductive hearing loss due to ossicular discontinuity, fusion, or fracture is usually evident by direct visualization, pneumatic manipulation, or acoustic reflex testing, but occasionally, these findings may not be detectable by clinical exam alone [30]. In these cases, CT can help define ossicular chain integrity. While imaging resolution has improved, a negative CT scan does not exclude otosclerosis or mimics such as incus lentiform process necrosis/osteolysis.

CT is also useful to define extent of other otodystrophies which may present with conductive hearing loss including Paget’s disease, osteogenesis imperfecta, fibrous dysplasia, and osteopetrosis. In Paget’s disease, demineralization tends to be more severe and extensive [33, 34] and is visible throughout the calvarium and skull base, not restricted to the classic location for otosclerosis [35]. Additionally, pagetoid involvement tends to be asymmetric [36]. Imaging findings of osteogenesis imperfecta tend to be more severe, with bilateral involvement, and sometimes with involvement of the ossicular chain, a finding not seen in otosclerosis (although, generally, the diagnosis is known prior to imaging due to systemic features) [37]. Fibrous dysplasia typically appears as osseous expansion with a ground-glass matrix. Rarely, fibrous dysplasia can be isolated to the ossicles, resulting in pure conductive hearing loss. Osteopetrosis involvement of the temporal bone is often diffuse, symmetric, and bilateral with marked bony sclerosis that can result in narrowing of the internal auditory canal, without the prominent volume expansion seen in fibrous dysplasia [36, 38].

Other processes such as otosyphilis and post-radiation changes can cause a demineralized appearance of the otic capsule, mimicking otosclerosis on imaging. In otosyphilis, demineralization tends to be more irregular, hazy, and permissive. Additionally, otosyphilis can involve the ossicular chain, a finding which does not occur with otosclerosis [39, 40, 41, 42]. Although the clinical context may allow for differentiation, in the case of post-radiation change, permissive demineralization should be visible in the central and posterior skull base in addition to any visible changes in the otic capsule [36].

Pre-surgical Imaging

Imaging can help detect known risk factors for stapes surgery and findings associated with failed stapes surgery (Fig. 2). Obliterative mature otosclerosis can be evaluated by CT, characterized by extensive thickening of the stapes footplate and filling of the oval window niche producing a “lobster-claw” fixation of the stapes footplate [43]. Pre-operative knowledge of obliterative otosclerosis can be helpful for appropriate planning and patient counseling [43, 44*, 45, 46]. Additionally, CT can help identify whether otosclerosis involvement extends to the round window and whether there is obliteration, overgrowth, or occlusion of the round window which can result in failed stapes surgery [30, 47]. Third window defects can also result in failed stapes surgery but are readily detected by CT. Most frequently, third window defects are due to dehiscence of the superior semicircular canal [48], but careful inspection of the lateral and posterior semicircular canals, vestibule and vestibular aqueduct, and cochlea are important to avoid missing subtle defects elsewhere [49]. Labyrinthitis ossificans can also be detected by CT, characterized by various degrees of mineralization/ossification within the periotic labyrinth and membranous labyrinth. Labyrinthitis ossificans may contribute to sensorineural and/or mixed hearing loss which would not be corrected by prosthesis placement and can also contribute to failed stapes surgery [44*].

Several anatomic variations may affect surgical approach. Variations of the ossicular chain can have surgical implications, in particular variation in the size and morphology of the lentiform process of the incus. Although evaluation of the lentiform process of the incus is near the limit of CT resolution, suspected variations can be conveyed, preparing the surgeon for potential prosthesis adjustment [30, 46]. Coronal CT profiles the tympanic facial nerve canal, for detection of canal dehiscence or nerve prolapse covering the oval window niche and cochlear promontory [50]. In situations where the overhanging facial nerve crosses the oval window, the stapes footplate can directly attach to the facial nerve, placing the patient at risk for facial nerve...
injury if a surgical repair for otosclerosis is planned [30, 50]. Pre-operative CT can define if there is a high riding jugular bulb, dehiscent jugular bulb, or jugular diverticulum, and if there is vascular encroachment upon the round window niche [47]. Non-contrast CT is sufficient to exclude aberrant internal carotid artery, to avoid surgical complication [51, 52]. Associated with the aberrant internal carotid artery, a persistent stapedial artery may appear as a soft tissue mass adjacent to the tympanic segment of the facial nerve [53]. This may cause conductive hearing loss by limiting stapes movement [54] and rarely may occur concurrently with otosclerosis [55]. A careful review of the inner ear structures is critical to identify any dysplasia that may indicate a risk for perilymph gusher at the time of stapedectomy [56]. Finally, a rare but important anatomic variant is a persistent peri-tympanomeningo fissure (Hyrtl’s fissure), which can cause conductive hearing loss and is a potential route for perilymphatic congenital fistula and CSF otorrhea [57, 58]. On CT, the tympanomeningo fissure can be seen inferior to the cochlear aqueduct.

Patients with far advanced otosclerosis [59] who develop sensorineural hearing loss (thought to be due to hyalinization of the spiral ligament [60]) or the result of ototoxic and metabolic factors [61], may benefit from cochlear implantation [62]. Several pre-surgical imaging findings can be beneficial prior to cochlear implantation. As with stapes surgery planning, CT can allow for evaluation of labyrinthitis ossificans which can complicate cochlear implantation. Through CT, the degree of mineralization/ossification can be directly assessed [63]. CT can also evaluate for diffuse retrofenestral disease which has been associated with an increased likelihood of facial nerve stimulation with cochlear nerve implantation, presumably due to bone thinning (spongiotic bone) proximal to the genu/labyrinthine segment leading to decreased impedance between the facial nerve and the electrode within the cochlea [22]. Few reports have described “cavitary otosclerosis” as a potential surgical pitfall for cochlear implantation [64, 65] as the electrode may penetrate into peri-cochlear cavities. Pre-operative knowledge of “cavitary otosclerosis” may help avoid this potential complication.

**Post-surgical Imaging**

The most common cause of persistent or recurrent conductive hearing loss after surgery is prosthesis migration or dislocation (Fig. 3). A well described mechanism is the “lateralized piston syndrome” where the piston migrates laterally out of the oval window, at times so far laterally that it contacts the tympanic membrane. This can lead to a distinctive delayed conductive hearing loss which temporarily improves with changes in middle ear pressure [66]. Occasionally, the piston can instead migrate medially into the vestibule. In addition to migration at the oval window, the prosthesis can dislocate at the incus, appearing on CT as a gap between the incus and the prosthesis.

Another cause of failed surgery is too short or too long of a piston; too short may not reach the oval window and too long may insert too deep into the vestibule and may contact either the saccule or utricle and lead to vertigo [67]. It is possible
Otosclerosis is a common and important cause of hearing loss. Although uncomplicated cases of stapedial otosclerosis can be diagnosed clinically, imaging can play an important role to clarify the diagnosis, reduce operative morbidity, and assess recurrent symptoms.

Fig. 3 Post-surgical evaluation. A. Normal prosthesis placement. Axial CT demonstrates a normally positioned prosthesis tip at the oval window (arrow). B. Medial migration of the piston. Axial CT demonstrates a medially displaced piston, terminating within the vestibule (arrow). C. Insufficient prosthesis length. Axial CT demonstrates the piston terminating lateral to the oval window (arrow). Note streak artifact associated with the metal prosthesis which can complicate evaluation. D. Dislocated prosthesis. Axial CT demonstrates posterior displacement of a non-metallic prosthesis (arrow).

Conclusion

Otosclerosis is a common and important cause of hearing loss. Although uncomplicated cases of stapedial otosclerosis can be diagnosed clinically, imaging can play an important role to clarify the diagnosis, reduce operative morbidity, and assess recurrent symptoms.

Compliance with Ethical Standards

Human and Animal Rights and Informed Consent This article does not contain any studies with human or animal subjects performed by any of the authors.

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