The Forgotten Second Window: A Pictorial Review of Round Window Pathologies

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AJNR Am J Neuroradiol  published online 12 December 2019
http://www.ajnr.org/content/early/2019/12/12/ajnr.A6356
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ABSTRACT

SUMMARY: The round window serves to decompress acoustic energy that enters the cochlea via stapes movement against the oval window. Any inward motion of the oval window via stapes vibration leads to outward motion of the round window. Occlusion of the round window is a cause of conductive hearing loss because it increases the resistance to sound energy and consequently dampens energy propagation. Because the round window niche is not adequately evaluated by otoscopy and may be incompletely exposed during an operation, otologic surgeons may not always correctly identify associated pathology. Thus, radiologists play an essential role in the identification and classification of diseases affecting the round window. The purpose of this review is to highlight the developmental, acquired, neoplastic, and iatrogenic range of pathologies that can be encountered in round window dysfunction.

ABBREVIATIONS: LO = labyrinthitis ossificans; RW = round window

The round window (RW) serves as a boundary between the basal turn of the cochlea anteriorly and the round window niche posteriorly. It, along with the oval window, is 1 of 2 natural openings between the inner and middle ear. The round window is often overshadowed by the “first window” (the oval window) and pathologic “third windows” (eg, superior semicircular canal dehiscence). Nevertheless, numerous developmental, acquired, neoplastic, and iatrogenic processes can affect the round window membrane and niche. Any of these can cause conductive hearing loss because occlusion of the round window prevents propagation of acoustic energy along the cochlear axis. In operative interventions in which a primary goal of surgery is to improve conductive hearing loss or access the round window region for cochlear implantation, accurate preoperative identification of round window abnormalities is essential to first determine whether surgery is a worthwhile endeavor and subsequently to guide the surgical strategy. In this article, the various pathologic entities and surgical considerations of the round window that can be encountered on imaging are reviewed.

Physiology and Functionality
The inner ear “windows” refer to openings in the otic capsule that connect the fluid in the inner ear to either the middle ear or intracranial space. The 2 primary natural openings are the oval and round windows. Other windows include the cochlear and vestibular aqueducts and tiny foramina that transmit vessels and/or nerves to the inner ear and adjacent structures (eg, the petromastoid canal and singular canal).

Functionally, the role of the oval and round windows is related to sound transmission: Vibratory acoustic energy enters through the oval window, is transmitted through the cochlea, and exits into the middle ear cavity via the round window. The fluid in the cochlea through which sound is transmitted is functionally incompressible due to the surrounding osseous structures. Movement of the cochlear fluid is thereby dependent on the mobility of the round and oval window membranes: Inward displacement of the oval window membrane via the stapes by ossicular vibration is matched by outward round window membrane displacement.

Developmental Anomalies
Normal Anatomy. The round window is located along the posterior aspect of the cochlear promontory and measures 1.5–2.1 mm horizontally, 1.9 mm vertically, and 0.65 mm in thickness (Fig 1). The round window membrane is thicker along its edges and
thinner in the middle and is made up of 3 layers: 2 epithelial layers facing the inner and middle ear, respectively, and connective tissue in the core. Contrary to its name, the shape of the round window is typically skewed, ovoid, and nonplanar according to a recent study. The round window niche is primarily defined by the relatively thin overhanging bone that naturally extends from the promontory. This overhanging bone may obscure complete direct visualization of the round window membrane during routine middle ear surgery and cochlear implantation (Fig 2). In addition, most ears have a thin layer of mucosa covering the round window membrane, often called a "pseudo-membrane," that blocks direct visualization of the window if not removed.

Round Window Stenosis and Atresia. Round window absence is a rare abnormality that may be seen in conjunction with various syndromes, including incomplete partition anomalies, mandibulofacial dysostosis, and Coloboma of the eye, Heart defects, Atresia of the choanae, Retardation of growth and/or development, Genital and/or urinary abnormalities, and Ear abnormalities and deafness (CHARGE) syndrome, in addition to cases of aural atresia (Fig 3). In very rare cases, it may also occur without an associated syndrome (Fig 4). Some authors have posited that such nonsyndromic cases may represent an inherited autosomal dominant genetic disorder with variable penetrance. Even in nonsyndromic cases, few reports exist of round window atresia as an isolated finding; most of these patients have associated middle ear or pinna abnormalities.

Patients with round window atresia typically have a mixed but predominantly conductive hearing loss, with an associated typical air-bone gap of 30–40 dB. Attempts to surgically recreate the round window do not always produce substantial gains in hearing. The reasons remain uncertain because few such surgical
case reports exist. However, the inconsistent results may be related to the presence of associated anomalies such as congenital stapes fixation, otosclerosis, or re-ossification following an operation.

Unfortunately, because surgeons often have incomplete exposure of the round window niche during routine otologic procedures and because of its rarity, round window atresia can easily be overlooked intraoperatively. It also is frequently missed on imaging, and many patients with round window atresia are not diagnosed until middle ear exploration. Close attention must therefore be paid to the morphology of the round window and round window niche on imaging performed for conductive or mixed hearing loss.

Stenosis of the round window membrane and/or recess is more common than complete round window absence. Like round window atresia, stenosis of the round window can have syndromic associations and contribute to hearing loss. The lower limit of normal size for the round window niche is generally cited as being 1.5 mm. Nevertheless, imaging characteristics of stenotic recesses can be quite variable, ranging from mild to severe (Fig 5).

**Acquired Abnormalities**

Acquired abnormalities that affect the round window include a range of traumatic, inflammatory, and iatrogenic processes. Otitis media associated with mucosal thickening and effusion may obscure the round window or adjacent niche (Fig 6). Less commonly, barotrauma or increased pressure may cause the round window membrane to rupture, leading to perilymph fistula with sensorineural hearing loss and vertigo. Temporal bone fractures through the round window may disrupt the membrane. Several primary osseous processes such as Paget disease, fibrous dysplasia, otosyphilis, and osteogenesis imperfecta can affect the temporal bone, including the annular ring of the round window. Below, several of the most common causes of acquired abnormalities of the round window are discussed, with imaging correlates.

**Labyrinthitis Ossificans.** Labyrinthitis ossificans (LO) refers to ossification within the membranous labyrinth, most frequently occurring within the scala tympani. Typically, LO is secondary to inflammatory changes from infection such as supplicative otitis media, labyrinthitis, or meningitis, though trauma and otosclerosis have also been indicated as inciting processes. In cases of LO associated with bacterial meningitis, the basal turn of the cochlea is often preferentially affected because infection may spread from the subarachnoid space through the cochlear aqueduct to the proximal scala tympani (though spread may also occur through the modiolus). In some cases, LO may involve the round window, involvement thought to occur when the inciting event is otitis media or meningitis that spreads through the round window along the scala tympani (Fig 7). The etiology of
LO cannot be predicted on the basis of mineralization patterns seen on CT.\textsuperscript{27} LO is associated with the development of sensorineural hearing loss and may render cochlear implantation more challenging and outcomes less rewarding.\textsuperscript{28,29} Extensive LO may preclude the placement of a cochlear implant. Expected CT findings of LO involving the round window include thickening/high attenuation along the membrane, likely with coexistent ossific material in the basal turn of the cochlea or elsewhere in the membranous labyrinth. In the early stages of labyrinthitis, fibrosis may be evident only on heavily weighted T2 imaging, where low signal is seen within or partially replacing normally high-signal perilymph.

Otosclerosis. Otosclerosis is an acquired condition in which spongiotic bone replaces mature endochondral bone of the otic capsule.\textsuperscript{30} Typically, otosclerosis involves the oval window and stapes footplate in the region of the fissula ante fenestram.\textsuperscript{30,31} Fenestral otosclerosis may also secondarily involve the round window in approximately 3\%–13\% of cases.\textsuperscript{32,33} Isolated round window otosclerosis has been reported, though the prevalence is rare (approximately 0.3\% of cases).\textsuperscript{32} Nevertheless, secondary involvement or isolated involvement of the round window by otosclerosis is an important radiologic observation because it portends a poorer chance of surgical success.\textsuperscript{30} Specifically, round window involvement diminishes the movement of perilymphatic fluid following stapedectomy and prosthesis placement, the surgery typically used in cases of otosclerosis.\textsuperscript{34,35}

The appearance of otosclerosis on imaging depends on the disease phase. In the active (otospongiotic) stage, the affected bone surrounding the round window will appear lucent and demineralized (Fig 8).\textsuperscript{36,37} Later on, these areas are replaced by sclerotic bone during the nonactive (otosclerotic) stage.\textsuperscript{37} As this replacement occurs, the round window membrane may become thickened or irregular. Heaped-up osseous plaques may narrow the round window or adjacent niche.\textsuperscript{34}

Mansour et al\textsuperscript{32} categorized otosclerosis of the round window on the basis of the extent of involvement of the membrane and adjacent structures. Under this classification scheme, RW-I represented hypodensities about the round window edge, RW-II had partial thickening of the membrane, RW-III showed global membrane thickening with a persistent air-filled recess, RW-IV had obliteration of the recess, and RW-V demonstrated overgrowth of otosclerotic foci. Predictably, higher grades of round window involvement were found to be associated with more severe hearing loss, likely related to increased impedance within the scala tympani.\textsuperscript{32}

Jugular Bulb Dehiscence and Diverticula. The positional anatomy of the jugular bulb is variable throughout a person’s life.\textsuperscript{38} Jugular bulb abnormalities consequently develop with time and are typically acquired by the fourth decade of life.\textsuperscript{38} High-riding jugular bulbs are found in approximately 8\% of patients, both on pathologic specimens and CT images.\textsuperscript{38} Dehiscent jugular bulbs and jugular bulb diverticula are more rare, occurring at a rate of 2.6\% and 1.2\%, respectively.\textsuperscript{39,40}

High-riding jugular bulbs, dehiscent bulbs, and jugular diverticula are often asymptomatic.\textsuperscript{41} However, they may also present with pulsatile tinnitus or, less commonly, conductive hearing loss, likely related to encroachment by the bulb on the round window, ossicular chain, or tympanic membrane (Fig 9).\textsuperscript{41,42} The incidence in which the round window membrane is specifically involved is rare; 1 histologic analysis of temporal bones identified 2 such cases in 1579 specimens (0.1\%).\textsuperscript{43}

Imaging findings vary on the basis of the type of jugular bulb abnormality. High-riding bulbs typically occur as an isolated finding, in which the dome of the jugular bulb is within 2 mm of the internal auditory canal (IAC) floor (though definitions vary).\textsuperscript{44,45} A high-riding bulb may extend further superolaterally, erode the sigmoid plate, and protrude into the middle ear cavity; this dehiscence is best seen on CT as thinning and/or absence of bone between the bulb and middle ear structures.\textsuperscript{44}
Finally, diverticula will appear as distinct outpouchings from the bulb. Any jugular bulb anomaly seen on CT should be closely examined for the presence of abutment of the bulb on the round window membrane, niche, or other middle ear structures.

Neoplastic Processes. Several neoplastic processes may affect the round window membrane or round window niche. Some, such as metastases and primary osseous tumors, are centered in the bone; secondary round window involvement by these lesions depends on their size and location. Notably, most primary osteodystrophies and osseous neoplasms spare the otic capsule, given divergent embryology and composition. Other tumors have an anatomic proclivity to involve the round window membrane. For example, intralabyrinthine schwannomas may extend through the round window membrane and into the round window niche (dashed arrows), better demonstrated on the follow-up axial (D), Pöschl (E), and coronal (F) CT images.

Similarly, jugular paragangliomas—comprising glomus jugulare and glomus tympanicum tumors—may also involve the round window. Glomus jugulare tumors arise from the tympanic nerve (Jacobson nerve) and grow within and along the cochlear promontory; if large enough, they may extend into and occlude the round window niche. Glomus jugulare tumors begin within the jugular foramen. Although histologically benign, glomus jugulare tumors are locally destructive and can erode into adjacent middle ear structures such as the round window niche (Fig 11).

Surgical Considerations
Surgical Access and Round Window Visualization. Several conditions may require surgical dissection at the round window niche. Most commonly, the niche is accessed during cochlear implantation. Most cases are performed using a standard cortical mastoidectomy with facial recess. The bony overhang of the round window niche is then drilled to allow direct round window membrane visualization during electrode insertion. Conversely, when surgical access of the round window or niche is performed to treat infectious or neoplastic middle ear processes, the region is typically visualized via a transcanal approach. In addition, there are uncommon situations in which round window occlusion is performed to treat traumatic perilymphatic fistulas or superior canal dehiscence syndrome. In these cases, the middle ear is typically accessed via a transcanal approach, the round window is visualized with an operating microscope or endoscope, and fascia with or without cartilage is packed in the round window niche.

Recently, endoscopes have gained popularity in otologic surgery. Endoscopy allows superior visualization around corners, particularly during removal of a cholesteatoma in the sinus tympani and anterior epitympanum. Thus, endoscopies are sometimes used to augment microscopic techniques in complex cases. However, for visualization and access of the round window, the use of an endoscope does not carry a substantial advantage over microscope visualization in most cases because the round window niche is readily seen when performing either transcanal or transmastoid-facial recess surgery via a direct line of sight.

Postoperative Changes in Third-Window Lesions. Third-window lesions refer to any range of pathology that creates an abnormal connection between the inner ear and either the middle ear or the intracranial cavity. Acoustic energy is lost through these windows, often causing a so-called “pseudoconductive” hearing loss, which manifests as increased bone and decreased air microstructure.
vascular or nervous channels. There are many such connections: vestibular aqueduct enlargement, semicircular canal dehiscence, and any other type of osseous thinning between the inner ear and adjacent vascular or nervous channels. Patients can present with Tullio phenomenon (vertigo symptoms induced by loud noises) or Hennebert sign (similar symptoms induced by increases in pressure within the ear canal) due to deflection of the superior canal cupula by endolymphatic fluid escaping the osseous defect.

For cases of superior semicircular canal dehiscence, surgical plugging of the pathologic osseous defect can be completed, either via a middle cranial fossa craniotomy or transmastoid approach. Alternatively, the round window can be targeted; surgeons may reinforce the round window with overlying tissue (e.g., fascia, cartilage, fat) or occlude the round window niche.

Cochlear Implant. Cochlear implants may be inserted through a cochleostomy adjacent to the round window or directly through the round window membrane. Most surgeons surveyed in 2007 preferred the former approach, though these decisions may be based on historical presumptions; early multichannel leads were thought to traumatize the cochlea if placed through the round window. A more recent survey found that today most surgeons prefer round window membrane electrode insertions, given the natural and less traumatic access provided to the scala tympani—the preferred location of electrode placement. A recent study found no difference in the number of audiometric outcomes or postoperative complications among groups undergoing electrode placement via either approach; however, most cochlear implant surgeons now prefer the round window approach when trying to preserve any residual natural acoustic hearing.

Regardless of the cochlear entry point, complications do occur. The electrode can kink, flip over at its tip, or migrate. Postoperative imaging can also demonstrate various degrees of electrode displacement, including within the semicircular canal, internal carotid artery, internal auditory canal, and vestibule. Across time, electrodes may also migrate from their initial position. Postoperative images should be evaluated in the context of the surgical approach (i.e., via the round window or adjacent cochleostomy) and should include an assessment of electrode position, integrity, and change since prior examinations.
FIG 13. A 64-year-old woman who had poor progress of hearing following the placement of a cochlear implant. Axial CT image at the time of presentation (A) shows that the electrode array is retracted from its expected location, with multiple electrode leads located outside of the cochlea (straight white arrow). Follow-up imaging after surgical revision (B) shows normal positioning of the implant, with the first electrode located approximately 4 mm past the round window (curved black arrow).

CONCLUSIONS
Numerous acquired and developmental processes may affect the round window, presenting with varying clinical symptoms. Such cases can be challenging to radiologists who are unfamiliar with the local anatomy and pathologies. Nevertheless, because the round window and round window niche are often anatomically obscured during an operation, imaging plays a uniquely important role. Thus, radiologists should scrutinize the round window and familiarize themselves with the anomalies and disease processes that may be encountered in this important anatomic region.

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