Cochlear Dysfunction is not Common in Human Meningioma of the Internal Auditory Canal

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Hypothesis: Cochlear dysfunction is not common in human meningioma of the internal auditory canal.

Background: Meningiomas arising from the cerebellopontine angle and internal auditory canal typically cause hearing loss. Cochlear dysfunction is known to contribute to sensorineural hearing loss induced by vestibular schwannoma, the most common tumor of the internal auditory canal. Detailed cochlear histopathology in meningioma has not been reported.

Methods: Retrospective analysis of cochlear histopathology in five unoperated and five operated meningiomas of the internal auditory canal identified after screening human temporal bone collections from three academic medical centers.

Results: While some dysfunction of all analyzed cochlear cell types was identified, a predominant or exclusive loss of hair cells was not observed in any meningioma. Only 14.3% of temporal bones showed significantly more hair cell damage on the side of the tumor when compared with the contralateral ear; cochlear neuronal damage was more prevalent in meningiomas. The incidence of hydrops, perilymphatic precipitate, endolymphatic precipitate was low.

Conclusions: Substantial cochlear damage in human meningioma of the internal auditory canal is not common. This may explain the anecdotal hearing improvement observed after surgical resection of meningioma. Our findings underline the importance of developing therapeutic strategies to prevent cochlear neuronal degeneration due to tumors of the internal auditory canal.

Key Words: Cochlear dysfunction—Histopathology—Human meningioma—Internal auditory canal.


Meningioma is the most common intracranial tumor in adults (1), and the second most common tumor of the cerebellopontine angle (2). Though usually benign, if located in the cerebellopontine angle and internal auditory canal, meningioma can lead to degenerative changes in the facial, vestibular, and cochlear nerves (1). Vestibular schwannoma (VS) is the most common tumor of the cerebellopontine angle and internal auditory canal, accounting for 85% of tumors in these locations. Both VS and meningioma typically present with hearing loss and tinnitus (3,4). While histopathologic study of human temporal bones reveals that cochlear damage contributes to VS-induced hearing loss (5), it is unclear to what degree cochlear damage also contributes to meningioma-induced hearing loss. To our knowledge, there is only one published case of temporal bone histopathology in a patient affected by meningioma of the cerebellopontine angle (6).

The goal of this study was to analyze a series of human temporal bones with internal auditory canal meningioma and quantify damage to specific cochlear cell types. Our results can inform preoperative counseling about the likelihood of hearing improvement after surgery for internal auditory canal meningioma.

METHODS

Temporal Bone Preparation and Study
Archival collections of human temporal bones from the US Temporal Bone Registry at Massachusetts Eye and Ear (Harvard Medical School), House Research Institute (University of California, Los Angeles), and the University of Minnesota were screened for cases of meningioma located within the internal auditory canal. This inclusion criterion was chosen because VSs often arise in close proximity to the cochlea, while meningiomas can be found throughout the cranial cavity and in the spinal canal. Such a difference in distance of tumor to the fundus of the internal auditory canal may lead to a bias in the analysis of cochlear dysfunction. Written informed consent from all patients had been obtained before death. Out of 26 initially identified temporal bones, 16 were excluded because the tumor
was not in the internal auditory canal (five bones), the cochlear tissue was insufficiently preserved (two bones), the appropriate sections were missing (one bone), or patients had comorbidities that made it impossible to separate meningioma-related damage from sequelae of other diseases, including neurofibromatosis type II with bilateral VSs (four bones), spontaneous VS (two bones), Menière’s disease (one bone), and bilateral Mondini dysplasia (one bone). Of the remaining 10 cases, half had undergone surgical resection. Because surgery could have contributed to cochlear damage, we analyzed the unoperated meningiomas separately (five bones, age range 34–81 yr, median 65 yr, 60% women) and in the context of all identified meningiomas (10 bones, age range 34–89 yr, median 67 yr, 60% women). Patient characteristics are summarized in Table.

Assessment of Cochlear Pathology

Two independent researchers (L.D.L. and F.H.L.) evaluated cochlear sections ipsilateral and contralateral to the internal auditory canal meningioma for inner hair cell (IHC) and outer hair cell (OHC) loss, atrophy of the stria vascularis, and loss of spiral ganglion neurons (SGNs).

The presence of hydrops was determined by a convexity of the stapes footplate or a minimal air–bone gap (less than 10 dB) of greater than 1 cm.

Table 1, predominant SGN damage was most pronounced, with 40% in the unoperated and 50% in the operated group. Cochlear damage was absent or minimal in 20% of temporal bones with meningiomas. Moderate or severe damage to two or more cochlear cell types was observed in 20% of unoperated meningiomas, while presumed iatrogenic damage doubled this number (40%).

RESULTS

Table 1

<table>
<thead>
<tr>
<th>Meningioma (excl. surg.)</th>
<th>Predominant HC Damage</th>
<th>Predominant SGN Damage</th>
<th>Predominant Stria Damage</th>
<th>Predominant HC Damage</th>
<th>2+ Cell Types Damaged</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meningioma (excl. surg.)</td>
<td>1 (20.0%)</td>
<td>0 (0.0%)</td>
<td>2 (40.0%)</td>
<td>1 (20.0%)</td>
<td>4 (40.0%)</td>
</tr>
<tr>
<td>Meningioma (incl. surg.)</td>
<td>2 (20.0%)</td>
<td>0 (0.0%)</td>
<td>3 (30.0%)</td>
<td>1 (20.0%)</td>
<td>4 (40.0%)</td>
</tr>
</tbody>
</table>

Excl./incl. surg. indicates meningiomas excluding/including those that underwent surgical resection; HC, hair cell; SGN, spiral ganglion neuron; stria, stria vascularis.

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(MB2) experienced a late-onset bilateral progressive sensorineural hearing loss and died of acute pulmonary edema and atherosclerotic heart disease at the age of 81 years. An accumulation of psammoma bodies in the right internal auditory canal is visible (consistent with a small psammomatous meningioma), along with bilateral moderate patchy atrophy of the stria vascularis in the cochlear apex (16.4% on the right) and bilateral moderate to severe primary neuronal degeneration in the cochlear base. Inner and outer hair cells remained intact.

**DISCUSSION**

To our knowledge, this is the largest histopathologic study of human temporal bones affected with internal auditory canal meningioma. Although audiometric data from patients who donated these temporal bones are not detailed (see Table, Supplemental Digital Content 1, http://links.lww.com/MAO/A556), audiometric thresholds in general do not predict cell type-specific damage (8), and our results provide new insight into common clinical observations. For decades, clinicians have observed hearing improvement after surgical removal of meningiomas and published detailed accounts of this phenomenon (9–14). Even among patients with preoperative deafness, presumably caused by infiltration of the auditory nerve and subsequent degeneration of cochlear structures, recovery of hearing has been observed in 1.8% of patients (15). Conversely, as even subtle improvement in hearing thresholds is rare after surgical excision of a VS, the primary goal of VS surgery is to preserve hearing at the preoperative level (5,16–20).

When comparing our results to the published VS study (5) to provide a possible mechanistic insight into this disparity, several limitations of such a comparison have to be pointed out: the small sample size, the historic nature of the VS data, and the limited available information regarding tumor size prevent any generalization. Nonetheless, Roosli et al. (5) reported that “tumor size, distance from the cochlea, and nerve of origin did not correlate with structural changes.” In terms of median age, our study population is comparable to the VS study (5), where median age was 72 years (range, 25–100 yr). Sex distribution in both cohorts is also relatively characteristic for each tumor, because meningioma is more common in women than men (21) while VS equally affects women and men (18 men and 14 women in the Roosli et al. study) (22). Mean distance from fundus to tumor was 1.76 cm by Roosli et al. (data obtained after contacting the authors). The VS study reported that

**TABLE 2.** Percent damage to different cell types per temporal bone ipsilateral to meningioma

<table>
<thead>
<tr>
<th></th>
<th>IHCs</th>
<th>OHCs</th>
<th>Stria</th>
<th>SGNs</th>
<th>Hydrops</th>
<th>EL</th>
<th>PL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meningioma (excl. surg.)</td>
<td>6/15 = 40.0%</td>
<td>6/15 = 40.0%</td>
<td>6.5/15 = 43.3%</td>
<td>7.5/15 = 50.0%</td>
<td>0/5 = 0%</td>
<td>1/5 = 20.0%</td>
<td>1/5 = 20.0%</td>
</tr>
<tr>
<td>Meningioma (incl. surg.)</td>
<td>12/30 = 40.0%</td>
<td>12/30 = 40.0%</td>
<td>14/30 = 46.7%</td>
<td>19.5/30 = 65.0%</td>
<td>0.5/10 = 5.0%</td>
<td>1.5/10 = 15.0%</td>
<td>2.5/10 = 25.0%</td>
</tr>
</tbody>
</table>

For each cell type (IHCs, OHCs, Stria, SGNs), the number of maximal points is 3 per temporal bone (i.e., 15 for 5 unoperated meningiomas, 30 for 10 meningiomas incl. surg.). For hydrops, endolymphatic precipitate (EL), and perilymphatic precipitate (PL), the number of maximal points is 1 per temporal bone.

Excl./incl. surg. indicates meningiomas excluding/including those that underwent surgical resection; IHC, inner hair cell; OHC, outer hair cell; SGN, spiral ganglion neuron; Stria, stria vascularis.

**FIG. 1.** Hematoxylin and eosin stain of mid-modiolar sections of left (L) and right (R) temporal bones of a patient (MB2) with a R-sided meningioma of the internal auditory canal and cerebellopontine angle. Stria (adjacent to circles), spiral ganglion neurons (adjacent to asterisks), and hair cells (adjacent to arrows) have similar appearance between the two sides. Magnified view in the central inset depicts psammoma bodies. Scale bars = 1 mm (white) in overview images, 100 μm (black) in close-ups.
18.8% of VSs demonstrated a predominant or exclusive loss of hair cells, while only 3.1% showed more pronounced SGN damage; 14 of these 32 unoperated specimens (43.8%) had moderate or severe damage to two or more cochlear cell types. The average damage per cell type in the ipsilateral ear revealed differences at the level of OHCs (60.4% for VSs). Furthermore, cochlear hydrops was present in 21.9% of VSs with endolympathic and perilymphatic precipitates observed in 31.3% and 43.8% of cases, respectively. Of the patients where ipsi- and contralateral temporal bones were available, the number with significantly more ipsilateral HC damage rose to 28.1% (9/32) in VSs.

Although precise reasons for the apparent differences between VS- and meningioma-induced cochlear damage remain to be determined, one possible contributor is a difference in tumor-secreted ototoxic and neurotoxic factors. We have previously shown that VSs secrete soluble molecules (23) and extracellular vesicles (24) that can cause direct cochlear damage. Such cochlear damage may exacerbate hearing loss due to tumor-induced compression of the auditory nerve and prevent hearing improvement after surgical tumor resection and decompression of the auditory nerve. Soluble factors secreted by meningioma have not been studied in the context of hearing loss.

Taken together, our findings emphasize the importance of developing therapeutic strategies to prevent cochlear nerve degeneration in meningiomas of the internal auditory canal.

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REFERENCES