Clinical and histopathological principles for the diagnosis of a recurrent paraganglioma of the jugular foramen initially diagnosed as a middle ear adenoma: illustrative case

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BACKGROUND Paragangliomas (PGLs) are rare neoplasms that may be associated with hereditary PGL syndromes and variable risk of metastasis. Middle ear adenomas are extremely rare tumors with no known hereditary predisposition and extremely low risk of metastasis. Although often easily differentiated, they may share clinical and pathological features that misdirect and confuse the diagnosis.

OBSERVATIONS The authors discussed a 35-year-old woman with left-sided hearing loss and bleeding from the external ear canal who presented to an outside hospital. She underwent resection of a middle ear and mastoid mass, initially diagnosed as a middle ear adenoma with neuroendocrine features, with later mastoidectomy and ligation of the sigmoid sinus with microsurgical excision of persistent tumor in the jugular foramen and temporal bone. Histopathologically, her tumor was vascular, composed of benign-appearing epithelioid cells with "salt and pepper" neuroendocrine chromatin arranged in vague nests. Lesional cells were GATA3-immunopositive, glucagon-negative, and succinate dehydrogenase-immunonegative, consistent with PGL rather than middle ear adenoma, and required further workup for hereditary PGL syndromes.

LESSONS This case demonstrates potential challenges in differentiating a PGL from a middle ear adenoma. The authors offer clinical, histopathological, and imaging principles to aid in diagnosis and workup.

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KEYWORDS paraganglioma; middle ear adenoma; neuropathology; jugular foramen; skull base; middle ear neuroendocrine tumor; NET

Paragangliomas (PGLs) are usually benign neuroendocrine tumors (NETs) derived from paraganglion cells of the autonomic nervous system. Middle ear adenomas are rare, benign tumors with epithelial and neuroendocrine properties that originate from middle ear mucosal epithelium and often present with nonspecific findings on imaging. Here we describe an unusual jugular foramen and middle ear tumor initially presenting with hearing loss and bleeding from the left ear that was first concerning for middle ear adenoma with neuroendocrine features and later determined to be a GATA3-positive and glucagon-negative PGL to illustrate clinical and pathological features of these two tumors.

Illustrative Case

A 35-year-old woman with history of migraine headaches presented to an outside hospital with left-sided hearing loss and bleeding from the left external ear canal. She did not have a personal history of hypertension, anxiety, palpitations, or syncope and denied a family history of pheochromocytoma or PGL. Imaging revealed a left mastoid and middle ear mass, which was partially resected during a left tympanoplasty and mastoidectomy. On pathology, her original tumor was diagnosed as a middle ear adenoma with neuroendocrine features possibly concerning for carcinoma. Five months later, she presented to our hospital with follow-up magnetic resonance imaging (MRI) that revealed persistent
tumor in the left jugular foramen extending down the jugular vein to the C1–2 level. She also complained of fatigue, persistent left hearing loss, tinnitus, difficulty with balance and coordination, frequent headaches, irregular heartbeat, and weight loss within the previous year. She denied dysphagia, hoarseness, or weakness lifting her shoulders.

On examination, her cranial nerves (CNs) were intact, with midline palatal elevation, full-strength bilateral sternocleidomastoid and trapezius muscles without atrophy, and tongue protruding midline and without atrophy. Her examination was otherwise unremarkable, with full strength and intact sensation in all extremities, normal reflexes, and absence of gait disturbance or dysmetria. T1-weighted MRI revealed a contrast-enhancing lesion of the left jugular foramen extending to C2 along the jugular vein and superiorly into the inferior aspect of the middle ear, with involvement of the petrous internal carotid artery (Fig. 1). Surgeons with neurosurgery and otolaryngology expertise performed a combined lateral skull base procedure, with mastoidectomy, left upper neck dissection, and ligation of the sigmoid sinus with placement and subsequent removal of a lumbar drain.

On pathological examination, the gross tumor was a red-pink soft mass without hemorrhage. A frozen section was taken that was concerning for a PGL versus recurrent middle ear adenoma. Permanent section revealed a vascular tumor with focal nodular pattern and sclerosing pattern with extensive collagen deposition. Tumor cells were small to medium with fine, speckled chromatin and scanty cytoplasm. On immunohistochemistry (IHC), tumor cells stained positive for synaptophysin and were negative for multiple cytokeratin markers (pankeratin, CK5/6, CK7, CAM5.2) and p63. Mucicarmine stain and tyrosine hydroxylase immunostain also stained negative. Of note, tumor cells were positive for GATA3 with nuclear staining by IHC. IHC for SSTR2A revealed a granular cytoplasmic staining as opposed to an expected membranous staining pattern and was considered equivocal. Sustentacular cells stained immunopositive with S-100 and SOX-10. Staining for CD45 revealed rare inflammatory cells, and CD31 demonstrated rich vasularity. Ki-67 proliferative index was low (~3% to 5%). IHC staining for succinate dehydrogenase (SDH) B was positive in endothelial cells, highlighting the vascular nature of the tumor and serving as an internal control, but was negative in tumor cells. Tumor cells were also negative for glucagon. This staining profile was consistent with an SDH-deficient PGL rather than a middle ear adenoma (Fig. 2).

A local neck lymph node was resected and exhibited reactive features with no evidence of metastatic PGL.

The patient followed up with the endocrinology department, which recommended genetic testing for SDHB mutations in family members. She had normal prolactin level, which ruled out concurrent prolactinoma, and an unremarkable urinalysis upon screening for hematuria and renal cell carcinoma. She did not have elevated urine catecholamines or metanephrines when evaluated for associated pheochromocytoma. Genetics studies did not reveal a mutation in SDHA, SDHB, SDHC, SDHD, VHL, MEN1, NF1, RET, or VHL genes. Four months postoperatively, an octreotide scan showed residual uptake in the tumor bed, although temporal bone computed tomography (CT) and MRI of the internal auditory canal only revealed expected postsurgical change without evidence of persistent tumor. Although SSTR2A IHC was not interpretable in this case, positive SSTR2A can suggest that a tumor is amenable to 68Ga-Dotatate imaging1 and 117Lu-Dotatate therapy.2 Multidisciplinary discussions are ongoing to determine her optimal course for surveillance imaging in the future.

FIG. 1. A: Preoperative axial CT demonstrating a mass (red arrow) with associated bony erosion at the left jugular foramen and into mastoid air cells. B: Preoperative axial postcontrast T1-weighted MRI revealing a contrast-enhancing mass (red arrow) in the left jugular foramen with involvement of the petrous internal carotid artery.

FIG. 2. A: Original magnification ×40. Hematoxylin and eosin stain of the lesion showing moderately cellular lesion with sclerosing pattern. B: Original magnification ×400. Hematoxylin and eosin stain showing vascular lesion with vague nodules and some nuclear atypia, without mitoses or necrosis. C: Original magnification ×400. IHC for synaptophysin in lesional cells demonstrates strong cytoplasmic granular staining. D: Original magnification ×400. On IHC for GATA3, nuclear immunostaining is present in lesional cells. E: Original magnification ×400. IHC for SDHB. Note granular cytoplasmic staining in vessels with visible lumen and no staining in lesional cells of PGL. F: Original magnification ×400. IHC for glucagon shows no staining in PGL cells.
Discussion

Observations

In this case, the patient had a recurrent tumor involving the middle ear and jugular foramen that was initially diagnosed as an extremely rare tumor, a middle ear adenoma, and later correctly diagnosed as a rare, although more common, PGL. Although their typical clinical manifestations are often distinct, these tumors may also present with nonspecific findings on imaging, during surgery, and on histopathology. In these scenarios, correct diagnosis is vital because PGLs in this location have a strong association with germ-line mutations and should trigger a genetic workup. Because these tumors straddle a position between neurotology and skull base neurosurgery, it is valuable to delineate their clinical, surgical, and histopathological features to aid in future accurate diagnosis.

Lessons

PGLs are a genetically diverse set of paraganglion-derived NETs that exist on a spectrum of metastatic risk. The term glomus tumor is sometimes used by clinicians in referring to PGLs found in specific locations such as the jugular foramen (glomus jugulare) or the middle ear (glomus tympanicum), although there may be diagnostic overlap between them as well (jugulotympanic glomus tumor). Of note, middle ear PGLs make up approximately 30% to 40% of head and neck PGLs. In recent years, it has also become clearer that there are multiple hereditary PGL syndromes with associated genetic mutations in SDH in addition to well-known associations with VHL, NF1, and RET mutations. Diagnosis of PGL carries the highest genetic predisposition of any tumor, especially in head and neck locations, and warrants an extensive workup.

Classically, clinical suspicion for PGL or pheochromocytoma is based on symptoms of catecholamine excess, such as episodic headache, sweating, palpitations, or hypertension, although head and neck PGLs are usually nonspecific, as in this case. The most common presenting symptoms for jugulotympanic PGLs are pulsatile tinnitus (~90%) and hearing loss (~80%), although lower CN deficits may also be seen. Suspicion may also be raised if patients present with associated diagnoses from known tumor syndromes such as von Hippel-Lindau or hereditary PGL syndromes or with family history of tumors or symptoms consistent with PGL or pheochromocytoma. Although these tumors are highly vascular and may bleed easily, it is uncommon for patients to present with bleeding from the ear because diagnosis is usually made before the tumor can erode through the soft tissues of the external ear canal. Traditional markers for diagnosis are plasma-free metanephrines and urinary fractionated metanephrines, although they are infrequently produced by head and neck PGLs, in which case plasma methoxytyramine levels may be more useful.

Given their highly vascular nature, it is vital to image possible PGLs before attempting biopsy if they are grossly accessible in the external ear canal because bleeding may be difficult to control once it begins. These tumors tend to erode through neighboring structures and follow the path of least resistance, particularly through the temporal bone in the case of jugular PGL, which is easily visualized on CT, and into vessels, possibly invaginating into the lumen of jugular vein. They are highly contrast enhancing on both CT and MRI, classically revealing a nonspecific “salt and pepper” appearance, although MRI is more sensitive and detects smaller PGLs. On diagnostic angiography, these tumors also reveal a clear vascular blush, which is absent in the case of middle ear adenoma.

Treatment options include stereotactic radiosurgery and open surgery, frequently with preoperative embolization. Radiosurgery primarily controls growth of the tumor and may reduce symptoms as a primary or secondary modality. Resection is another treatment option, particularly if there is existing CN compression with associated neurological deficit. Embolization has been used preoperatively since the 1980s to reduce intraoperative blood loss, although it is not curative by itself and may be associated with higher risk of postoperative CN deficits.

Histopathology classically reveals a Zellballen pattern of nested cells with prominent vasculature and sustentacular cells circumscribing them. Sclerosing, pigmented, and clear patterns have also been recognized. Cells will often show “salt and pepper” nuclei, staining positive for neuroendocrine markers such as synaptophysin or chromogranin A and staining negative for cytokeratins. GATA3, a transcription factor binding to promoter GATA motifs that was initially recognized in breast duct and urethelial cells, is routinely used in IHC staining and is positive in up to 95% of cases of pheochromocytoma and 89% of cases of PGL, although it may be variable. Another key in the histopathological diagnosis of PGL is recognizing mutated SDH genes and their clinical implications. SDH is a multunit enzyme in the Krebs cycle and electron transport chain that is vital for cellular respiration. The most common SDH mutation in familial head and neck PGL is in the SDHD subunit (~66%), with SDHB being second most common (~15%), and then SDHC (~6%). Functional SDH activity may be seen with granular staining of SDHB in the cytoplasm of tumor cells, corresponding to the mitochondrial location of the enzyme complex. Absence of staining in tumor cells, with concomitant positive staining in nontumor tissue as an internal control, is suggestive of an SDH mutation. It does not reveal which specific SDH gene, however, because mutation in any subunit will alter the conformation of the entire complex and affect staining. Genetic subtyping is required for final diagnosis. SDHB-mutated PGL (PGL syndrome type 4) is the most concerning clinically because there is an elevated risk of metastasis (23% versus low or 3% to 4% in other subtypes) and associated pheochromocytoma, renal cell carcinoma, gastrointestinal stromal tumor, and pituitary adenoma. There is also variable penetrance of this mutation, with approximately 21% penetrance by age 50 and 42% by age 70; typically, the gene must be inherited from the father to develop tumors. The loss of tumoral SDHB immunoreactivity in this patient in the absence of a detected SDH gene mutation is currently unexplained but occasionally occurs epigenetically as a result of SDHC promoter methylation.

First described in 1976, middle ear adenomas are rare, benign tumors of the middle ear that classically present with unilateral hearing loss and have neuroendocrine and epithelial or glandular properties on pathology. This has led to multiple naming conventions and controversy regarding their cellular origin. Although they have recently been called middle ear adenoma with neuroendocrine features, they are more appropriately called middle ear NETs, in keeping with recent immunohistochemical findings and a World Health Organization initiative aimed to harmonize the nomenclature of NETs across disciplines and locations. Their preoperative diagnosis is difficult because they do not have specific imaging findings and are ideally diagnosed on histopathology. They have presented with facial nerve (CN VII) palsies, although that is rare. On T1-weighted MRI, they are isointense to hyperintense relative to white matter and gadolinium contrast enhancing. Although anatomically they
may be in proximity to Jacobson’s nerve (tympanic branch of CN IX), they are detached and distinct on imaging, unlike the intimate involvement with Jacobson’s nerve that may be seen with glomus tympanicum tumors.8

On histopathology, middle ear adenomas are infiltrative and moderately cellular with variable architectures, including glandular/tubular, trabecular, nested, cribriform, single cell, or cystic patterns and possibly form sheets.26,27 They are characterized by pancytokeratin, CAM5.2, EMA, chromogranin A, and synaptophysin immu-

TABLE 1. Characteristics of jugulotympanic paragangliomas and middle ear adenomas that may be useful in their diagnosis and workup

|                        | PGL                                      | MEA                                      |
|------------------------|------------------------------------------|------------------------------------------|
| Frequency              | Rare                                      | Extremely rare                           |
| Common presenting symptoms | Pulsatile tinnitus, unilateral hearing loss | Unilateral hearing loss; may see nonpulsatile tinnitus; 25% present w/ no symptoms |
| Associated w/ lower CN palsies (especially CN X) | Yes                                      | Uncommon; rare cases of CN VII palsy |
| Wraps around glomus bodies & CNs | Yes                                      | No                                      |
| Bony erosion surrounding tumor | Common, especially in jugular PGL; less common in tympanic | Rare; ossicles may be embedded in tumor |
| Contrast enhancing on CT or MRI | Yes                                      | Yes, although variable                   |
| Vascular blush on angiogram | Yes                                      | No                                      |
| Glucagon IHC            | Negative                                  | May be strongly positive                 |
| GATA3 IHC               | Yes, ~90%                                 | Variable                                 |
| S-100 IHC               | Yes, positive staining in sustentacular cells | Uncommon                                |
| Associated w/ tumor syndromes | Yes, need to work up hereditary PGL syndromes   | No                                      |
| SDH mutations           | Sometimes                                 | No                                      |
| Resection as treatment  | Yes, although SRS may be used             | Yes, surgery is the mainstay of treatment |
| Is SRS effective against it? | Yes                                      | Typically avoided                        |
| Is preop embolization useful? | Yes                                      | Unlikely, although it has not been studied |

MEA = middle ear adenoma; SRS = stereotactic radiosurgery.

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