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<td>doi:10.1055/s-0034-1544112</td>
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A Rare Finding of Schwannoma of the Vidian Canal: A Case Report

Alisa Yamasaki1,2 Ahmad R. Sedaghat1,2 Giant C. Lin1,2 William T. Curry3 Helen A. Shih4 Stacey T. Gray1,2

1Department of Otolaryngology—Head and Neck Surgery, Massachusetts Eye and Ear Infirmary, Boston, Massachusetts, United States
2Department of Otology and Laryngology, Harvard Medical School, Boston, Massachusetts, United States
3Department of Neurosurgery, Massachusetts General Hospital, Harvard Medical School, Boston, Massachusetts, United States
4Department of Radiation Oncology, Massachusetts General Hospital, Harvard Medical School, Boston, Massachusetts, United States

Address for correspondence Stacey T. Gray, MD, Department of Otolaryngology, Massachusetts Eye and Ear Infirmary, 243 Charles St, Boston, MA 02114, United States (e-mail: stacey_gray@meei.harvard.edu).


Abstract

Background Schwannomas of the vidian canal are an extremely rare type of intracranial tumor that can have variable clinical presentations including headache, facial pain, facial muscle paralysis, decreased lacrimation, or nasal dryness. We present an atypical case of an incidentally identified asymptomatic vidian canal schwannoma.

Case Description A 49-year-old woman with a history of multiple sclerosis presented for routine surveillance magnetic resonance imaging that detected an ovoid mass originating in the vidian canal. Given the unusual location of the lesion, an endoscopic endonasal biopsy was performed and confirmed the diagnosis of a vidian canal schwannoma, for which the patient chose to receive fractionated radiation therapy.

Keywords ► vidian canal ► schwannoma ► fractionated radiation therapy

Conclusion When a vidian canal tumor is identified, endoscopic endonasal biopsy can be used to confirm the diagnosis before pursuing either surgical or radiotherapy treatment. In particular, fractionated radiation therapy offers a radiobiologically safe means of delivering radiation when there is concern for late radiation-related side effects following treatment completion.

Background

Schwannomas are rare tumors of the nerve sheath that arise from the primary glial cells of the peripheral nervous system. As such, schwannomas may arise from any myelinated peripheral nerve. They are generally well-circumscribed lesions that rarely undergo malignant degeneration. Although benign and slow growing, schwannomas can cause significant morbidity due to direct involvement of nerves or by local mass effect that can pose particular challenges for treatment.

Approximately 25 to 45% of all schwannomas are found in the head and neck, where they can originate from virtually any cranial nerve.1 The most common schwannoma is the vestibular schwannoma that arises from the vestibular branch of the eighth cranial nerve and accounts for 51% of all nerve sheath tumors found anywhere in the body.2 This is followed by schwannomas of the trigeminal nerve that comprise up to 8% of all schwannomas.3 By contrast, schwannomas in the nasal cavity or paranasal sinuses—and vidian canal schwannomas in particular—are much rarer. The vidian canal, also referred to as the pterygoid canal, extends from the pterygopalatine fossa to the foramen lacerum in the floor of the sphenoid sinus4 and contains the vidian artery as well as the vidian nerve. The vidian
nerve arises from the joining of the greater superficial petrosal nerve and the deep petrosal nerve. The greater superficial petrosal nerve consists of preganglionic parasympathetic fibers from the geniculate ganglion of the facial nerve. The deep petrosal nerve consists of postganglionic sympathetic fibers from the carotid sympathetic plexus.\(^5\)–\(^8\) The parasympathetic fibers of the vidian nerve synapse in the pterygopalatine ganglion, whereas the sympathetic fibers innervate the lacrimal gland and secretory pathways of the nasopalatine glands.\(^5,6\)

Schwannomas of the vidian canal can present with a variety of clinical symptoms. Damage to the vidian nerve itself can result in decreased or absent lacrimation, corneal desiccation as well as dryness of the nasal mucosa, and cluster headaches.\(^5\) Additionally, vidian schwannomas have been reported to compress the adjacent internal carotid artery or bulge through the lateral wall of the sphenoid sinus to expand into the cavernous sinus.\(^5,9\)

To date, only 4 cases of vidian nerve schwannomas and 11 cases of histologically unconfirmed peri-vidian tumors have been reported in the literature.\(^9\) Here we present a case of biopsy-proven schwannoma of the vidian canal. We discuss the patient’s initial presentation, imaging characteristics, histologic findings, and treatment for this unusual tumor and review the other reported cases of vidian canal schwannomas in the literature.

**Case History**

The patient is a 49-year-old woman with a 24-year history of multiple sclerosis (MS) who presented for follow-up evaluation after an acute flare of MS. In the past, her MS flares had been relatively infrequent, with motor and/or peripheral sensory deficits that were well controlled with short-term courses of steroids. During her most recent flare, she experienced numbness of her left head, arm, and leg, with full resolution of symptoms after steroid treatment. Her past medical history was otherwise unremarkable.

A routine surveillance brain magnetic resonance imaging (MRI) was performed to evaluate for MS progression. MRI showed minimal progression of disease compared with imaging from 8 years prior, with extensive T2, fluid-attenuated inversion recovery, and proton-density prolonged signal in the periventricular, pericallosal, and subcortical white matter consistent with MS. Incidentally, the MRI also revealed a 19 × 13 × 10 mm ovoid mass along the inferior margin of the right cavernous sinus that was not present on previous MRIs. The lesion was well circumscribed and homogeneously enhancing, with T1 hypointense and T2 hyperintense signal. A repeat dedicated MRI was performed that demonstrated the lesion originated within an expanded vidian canal (\(\text{Fig. 1}\)). A sinus computed tomography scan also showed bony erosion of an expanded vidian canal (\(\text{Fig. 2}\)). From a symptom standpoint, the patient noted some mild nasal congestion in the mornings but was otherwise asymptomatic without any facial pain or paresthesias, abnormal lacrimation, or dry eye symptoms. Physical examination was similarly unremarkable with a normal nasal endoscopy and no focal neurologic deficits. An endoscopic
Endonasal biopsy was performed with image guidance. The sphenoid ostium was identified medial to the superior turbinate in a transethmoidal fashion. The ostium was enlarged laterally and inferiorly, and a wide sphenoidotomy was created. The mass was easily identified within a noticeably expanded vidian canal along the lateral floor of the sphenoid sinus (see Fig. 3). The canal was opened along the floor of the sphenoid sinus to uncover the mass and a biopsy was obtained. The final histopathologic assessment confirmed a diagnosis of schwannoma. The patient's recovery was unremarkable, and she had no problems with dry eye or nasal irritation postoperatively. Two weeks after surgery, her endoscopic examination showed a well-healed sinus cavity.

Options of endoscopic surgical resection, single-fraction stereotactic radiosurgery (SRS), and conventional fractionated radiotherapy using stereotactic radiotherapy (SRT) were reviewed with the patient, and she decided to pursue treatment with radiation. Given the benign histology, a background of MS that could confer additional sensitivity to radiation, and the patient's young age, the patient decided to pursue SRT to minimize late normal tissue injury risks. SRT was delivered at 1.8 Gy per fraction to 50.4 Gy. Treatment was well tolerated; the most notable symptom was mild fatigue.

Discussion

Schwannomas of the vidian canal are an extremely rare type of intracranial tumor with variable presentation. Symptoms of vidian schwannomas can result from compression of surrounding structures or extension into surrounding areas that can cause headaches, facial pain and paresthesias, and facial muscle paralysis. Dysfunction of the vidian nerve itself may cause nasal dryness, decreased lacrimation, and corneal desiccation. Here we present an unusual case of a vidian nerve schwannoma incidentally identified in a patient who was asymptomatic from the lesion.

To date, four documented cases of biopsy-proven vidian nerve schwannomas have appeared in the literature. Reported cases include both male and female patients, ranging in age from 13 to 78 years, with both unilateral and bilateral lesions described.10–13 Presenting symptoms included headache and facial palsy, oculomotor palsy and cerebrospinal fluid leak, unilateral hearing loss secondary to obstructive serous otitis media, and facial paresthesias. Complete surgical resection was pursued for all four cases through both open and endonasal approaches, although one patient with bilateral vidian nerve schwannomas had only partial resection of one lesion due to difficulties with surgical access.13 All patients had significantly improved or stable symptoms, and the two patients with the most severe symptoms—one with oculomotor palsy and ptosis and the other with headache and facial nerve palsy—demonstrated the most dramatic postoperative improvement with near complete resolution of symptoms.10,13

Consistent with these cases, the traditional and curative management of peripheral nerve tumors is surgical; recent advancements in transnasal endoscopic skull base approaches frequently enable complete resection. Radiation therapy by either SRS or multiple fraction radiotherapy is also an alternative option for treatment. Conservative management through observation with serial imaging is also reasonable when surgical resection is either not feasible or involves a disproportionate risk of postsurgical morbidity and/or mortality. The choice of treatment modality depends on patient choice, functional status, presenting symptoms, tumor location, tumor volume, and—in rare instances—malignant potential. The ultimate goal of both surgery and radiotherapy is to maximize tumor control while also optimizing preservation of surrounding structures.

In this case, the patient decided to pursue treatment with radiation therapy. Conservative management was not considered, given the significant increase in tumor volume during the eight-year interval of MRI surveillance and the presence of bone erosion. Given the rare presentation of a mass located in the vidian canal, the decision was made to confirm the diagnosis with a biopsy prior to treatment. After histologic confirmation, the schwannoma was deemed favorable to be controlled by radiation. The choice between single-fraction SRS or multiple-fraction radiation was made based on size and proximity of the tumor to potential radiation-sensitive structures. Minimizing the risk of adverse effects from radiation was the determinant for pursuing fractionated radiation.

In recent years, radiation therapy has become a common alternative to surgical resection of intracranial schwannomas.16–21 Options include either single-fraction SRS or conventional daily low-dose fractionation over approximately five weeks. SRS is a well-tolerated procedure, typically completed during a single session of irradiation and far more convenient for the patient. SRS has become a widely accepted treatment option for vestibular schwannomas and has a high success rate of durable tumor control with often partial reduction in tumor volume, with progression-free survival of 93% at 5 years and 92% at 10 years.22 Fractionated radiation has similar results and is chosen over SRS when there is a concern that the size of target
exceeds safe tolerance to SRS, when there are critical radiation-sensitive structures in adjacent areas that may not be able to tolerate the single high-dose collateral exposure, or when there are underlying concerns of the patient to suggest a higher risk of an adverse event. Compared with surgery, radiation therapy has been reported in several studies to result in decreased mortality and fewer treatment-related complications for vestibular schwannoma including hearing loss and facial weakness. 22–25 Recent advances in imaging and radiation delivery offer more conformal treatment options such as SRT—which combines both the superior accuracy of target localization of a stereotactic system and reduction in the size of the radiation treatment fields—and fractionated treatment, a radiobiologically safer means of delivering radiation when there are concerns for late effects of radiation decades after treatment completion.

**Conclusion**

Schwannomas of the vidian canal are extremely rare tumors and may present with symptoms of headaches, facial pain, facial muscle paralysis, decreased lacrimation, and nasal dryness or as an incidental finding. When a vidian canal tumor is identified, endoscopic biopsy can be used to confirm a histopathologically benign diagnosis before considering treatment options. Treatment options include surgical resection and radiotherapy.

**References**