Dorsal nasal deformity: an unusual initial presentation of nasal schwannoma
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Nasal schwannomas are very rare neoplasms. We present the case of a 54-year-old woman who had dorsal nasal deformity and partial right-sided nasal obstruction over 5 years. Computerized tomography scan of the nose and paranasal sinuses showed a rounded soft tissue mass (21x18x16 mm) in the anterior part of the right nasal cavity with expansion of the overlying bony wall. Complete endoscopic excision of this mass was performed revealing benign schwannoma with no recurrence over 3 years of follow-up. According to the best of our knowledge, this case is the first reported case of a nasal schwannoma presenting with dorsal nasal deformity in the English literature. This case highlights the importance of thorough endoscopic nasal examination in every patient presenting with a dorsal nasal deformity as well as the consideration of nasal schwannoma in the differential diagnosis of a dorsal nasal deformity.

Keywords:
- nasal deformity
- neurilemmoma
- schwannoma

Introduction
Schwannomas or neurilemmomas are neurogenic tumors arising from Schwann cells of the neural sheath of the peripheral, cranial, or autonomic nerves [1]. Olfactory and optic nerves cannot give rise to schwannomas because they lack Schwann cells in their sheaths. Between 25 and 35% of all schwannomas occur in the head and neck region. However, fewer than 4% of head and neck schwannomas involve the nasal cavity and paranasal sinuses [2]. Clinically, these tumors usually present like other benign nasal masses with nonspecific symptoms. In a retrospective review of patients with sinonasal schwannomas, headache was the most common chief complaint, followed by unilateral nasal obstruction and epistaxis [3]. In this article, we describe our reported case of a nasal schwannoma presenting initially with dorsal nasal deformity.

Case report
A 54-year-old woman presented with a 5-year history of progressive dorsal nasal deformity. Three years later, she developed partial right-sided nasal obstruction. No history of nasal trauma was reported. Physical examination revealed that bony dorsal nasal deformity (diffuse prominence of the right lateral bony wall of the nasal pyramid) can be seen and felt (Fig. 1). Nasal endoscopy revealed a pink firm polypoid mass in the right nasal cavity, which originated from the lateral nasal wall very close to the nasal valve angle and anterior to the middle turbinate. Examination data were otherwise irrelevant.

Computerized tomography (CT) scan of the nose and paranasal sinuses showed a rounded soft tissue mass (21x18x16 mm) in the anterior part of the right nasal cavity expanding the overlying lateral bony wall of the nasal pyramid. There was no evidence of bone destruction (Fig. 2).

Preoperative punch biopsy was taken from the mass under local anesthesia revealing schwannoma. As schwannomas can be multiple and bilateral when associated with neurofibromatosis type 2 (NF2) and they are characteristic of NF2, thorough neurologic examination with formal audiologic testing and a gadolinium-enhanced MRI of the brain with thin cuts through internal auditory canals have been performed excluding the presence of vestibular schwannomas, meningiomas, and gliomas. A spinal MRI has not been requested because there were no signs or symptoms suggestive of myelopathy. Ophthalmologic examination was performed to rule out the presence of posterior subcapsular lenticular opacities.

After exclusion of the diagnostic criteria for NF2, the patient was scheduled for endoscopic excision of the mass under general anesthesia. First, uncinctomy with middle meatal antrostomy was performed to expose the mass completely. Thereafter, complete tumor excision including an adequate margin of the surrounding mucoperiosteum was carried out. Finally, frontal sinustomy was performed to allow drilling of the frontal process of maxilla to remove any tumor remnants at the site of attachment.
Histopathological examination of the excised specimen came up with a final diagnosis of benign schwannoma. The examination revealed a soft tissue tumor formed of alternating zones of compact spindle cells arranged in interlacing fascicles (Antoni A) and loose hypocellular zones (Antoni B). The nuclei in the more cellular areas were arranged in a palisaded pattern (Verocay bodies). No evidence of malignancy was found (Fig. 3). Immunoreactivity staining of the tumor cells for S-100 protein was intensely positive (Fig. 4).

The postoperative treatment consisted of antibiotics and nasal douches with endoscopic debridement to remove any nasal crustations during the follow-up visits, which were scheduled once per week for 2 months, and then every 4 months up to 3 years, with no endoscopic evidence of tumor recurrence over the 3-year follow-up. No complications were reported during or after the operation. A signed informed consent has been obtained from the patient for the publication of photographs and data.

Discussion
Nasal schwannomas are very uncommon neoplasms. They have been postulated to arise from the ophthalmic or maxillary divisions of the trigeminal nerve, the sympathetic fibers from the carotid plexus or the parasympathetic fibers from the sphenopalatine ganglion [4]. However, it is often impossible to identify the nerve of origin at the time of surgery, as was the case in our patient [3].

Figure 1

Dorsal nasal deformity of the right lateral wall of the nasal pyramid.

Figure 2

Preoperative computed tomographic scan (coronal view) shows a rounded mass in contact with the nasal septum and expanding the overlying nasal bone.

Figure 3

Histopathological section (hematoxylin-eosin staining, original magnification ×200).

Figure 4

S-100 protein immunohistochemical staining (original magnification ×400).
Sinonasal schwannomas are manifested with vague and nonspecific symptoms and signs depending on the location and size of the tumor. They are benign slow-growing lesions and invade adjacent structures by bony erosion and direct extension [4]. They rarely undergo malignant transformation. The most common clinical presentations are unilateral nasal obstruction, mucopurulent nasal discharge, hyposmia, facial pain, and recurrent epistaxis. Other less common presentations are exophthalmos, epiphora, or cranial nerve palsies, which may occur with sphenoid schwannomas [5]. As documented in previous reports [6,7] and in our reported case, nasal schwannomas are usually described as polyoid masses of hard to elastic consistency on endoscopic examination. They may be tan-white or red [6].

CT scan is usually sufficient for delineating the extent of benign schwannomas and outlining the bony margins to rule out the invasion [1]. However, MRI is useful for the delineation of intracranial or intraorbital tumor extension and compared with CT, MRI can better evaluate the cause of opacified sinuses (e.g. tumor or inflammation) [8]. In a retrospective review of 12 cases, the characteristic CT and MRI finding of sinonasal schwanna was a well-defined round, tubular, or partially lobulated soft tissue mass, most frequently an expansile mass with pressure remodeling of the adjacent bony walls. The tumors were isodense on CT scans and predominantly isointense on T1-weighted MR images and isointense to hyperintense on T2-weighted MR images, compared with the brain stem. Mild contrast enhancement on CT scans and marked enhancement on MR images were noted in most of the tumors with characteristic cystic or hemorrhagic degenerative changes especially in large ancient schwannomas [9].

Schwannomas were first described as a pathologic entity by Verocay [10] in 1908. In 1920, Antoni [11] described the histological aspect of this tumor distinguishing two alternating histological patterns: Antoni type A (fasciculated pattern) and type B (reticular pattern). Antoni type A areas are composed of compact spindle cells arranged in short or interlacing fascicles. Areas of nuclear palisading with nuclear alignment in rows are known as Verocay bodies. Antoni type B areas are composed of few spindle cells loosely arranged in a fibrillar myxoid stroma [3]. Intense immunoreactivity staining of the tumor cells for S-100 protein is characteristic of schwannomas (particularly in Antoni type A tissues) due to the high density of Schwann cells [12].

Complete surgical excision is the definitive treatment for schwannomas because of the benign nature of these tumors and their resistance to radiotherapy. Recurrences are rare after total removal. The surgical approach is determined according to the location and extent of the lesion [3]. Endoscopic endonasal approaches have been successfully implemented in many reports [2–5] to achieve complete excision of the lesion with the advantages of no external scar, shorter hospital stay, and less blood loss, when compared with traditional approaches [3]. Traditional approaches such as lateral rhinotomy, lateral rhinotony with external ethmoidectomy, Caldwell–Luc, and midface degloving approaches have been used depending on the location and extent of the tumor. Some extensive intracranial tumors require craniotomy [5,7].

Nasal deformity is a very rarely reported presentation of nasal schwannomas. According to a literature review, few cases of nasal tip schwannoma [13,14] presenting as nasal tip deformity and only one case of neurilemmoma of the nasal dorsum [15] in a child presenting as dorsal midline nasal mass have been reported. As far as we know, our patient represents the first reported case of an intranasal schwannoma that initially presents with a bony dorsal nasal deformity. Therefore, the clinician must always have a high index of suspicion with thorough assessment and examination of every patient presenting with dorsal nasal deformity to ensure that this tumor is not missed.

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References


