Schwannoma of cervical sympathetic chain: assessment and management

Introduction

Schwannoma or neurilemmoma are benign neoplasms of Schwann cell origin. The Schwann cell surrounds peripheral nerve tissue and is believed to originate from the neural crest. They are typically solitary, well-encapsulated, benign tumours characteristically running along the course of a nerve or attached to peripheral, cranial, or sympathetic nerves. Malignant transformation is rare. In the parapharyngeal space, schwannomas may arise from the last four cranial nerves or the autonomic nerves, the vagus being the most common site. Cervical sympathetic chain schwannomas are uncommon and most often appear as an asymptomatic, slow-growing, solitary neck mass; Horner’s syndrome is rarely apparent on physical examination. Pre-operative diagnosis may prove difficult, for evaluation of neurilemmoma,
contrast-enhancing computed tomography (CT) or magnetic resonance imaging (MRI) and ultrasound (US) are the most appropriate, carotid angiography is not necessary. Post-operative ipsilateral Horner’s syndrome is to be expected following removal of the tumour, due to the necessary division of the cervical sympathetic chain, but this does not appear to have an adverse effect on the patient. Herein, a patient with sympathetic schwannoma arising from the cervical sympathetic chain is described.

Case report

A 42-year-old Italian male presented with an asymptomatic left neck mass found on routine physical examination. The mass, measuring approximately 4.5 X 3 cm, was located below the upper portion of the sternocleidomastoid muscle. It was mobile, non-tender, and non-pulsing, with no associated bruit. The mass had been progressively increasing in size but was not associated with any other symptoms such as dysphonia, dysphagia, pain or weight loss; repeated US examination revealed that the mass had increased by 0.8 mm in 3 months. The patient denied smoking or consumption of alcohol. Diagnostic studies, including CT and MRI, confirmed a well-circumscribed mass in the left parapharyngeal space (Fig. 1), with anterior-lateral displacement of the common carotid artery and the internal jugular vein (Fig. 2). The mass extended from the base of the neck to the area of the thyroid gland. On MRI examination, the mass was slightly heterogeneous, with low signal intensity on T1 and high signal intensity on T2-weighted images and moderate enhancement postcontrast (Fig. 3). In addition to sympathetic schwannoma, the pre-operative differential diagnosis included vagal schwannoma, metastatic or reactive lymphadenopathy, and paraganglioma. Fine-needle aspiration revealed spindle-shaped cells with irregular fusiform nuclei in a vague interwoven pattern. The mass was excised through a transverse left cervical skin incision. The tumour was found to originate from the cervical sympathetic chain and did not involve the vagus, hypoglossal, spinal, accessory, glossopharyngeal, or lingual nerves. It could not be resected without sacrificing a portion of the cervical sympathetic chain. Post-operatively, the patient showed mild left pupillary miosis with ptosis, enophthalmos (Fig. 4) and facial anhydrosis of the ipsilateral face. The presence of all these features was indicative of a preganglionic lesion. The post-operative course was uneventful and the patient was discharged on the fifth post-operative day. Histology showed the tumour to be a benign schwannoma originating within the cervical sympathetic chain.
Ocular examination showed that the best corrected visual acuity was 20/20 in each eye (-3.75 sph.). Intra-ocular pressure was normal. Bilateral fundus: myopic aspect. On Hertel exophthalmometry, the left eye showed an enophthalmos of 2 mm and a normal pupillary reaction with a bright diameter of the right eye 2.5 mm and 2 mm for the left. The marginal reflex distance (MRD1) was 5 mm in the right eye and 3 mm in the left. The elevator function was 16 mm in the right and 15 mm in the left. After one year, ophthalmologic symptoms were unchanged.

Discussion

A review of the literature disclosed fewer than 45 cases \(^\text{3-6,8}\) of schwannoma sympathetic chain. This is a benign nerve sheath tumour that occurs along the length of the nerves. It is a solitary, slowly growing, tumour generally observed in patients between 20 and 50 years of age. Frequency is the same in both sexes and malignant change is rare \(^\text{5}\).

Microscopically, we found a fibrous capsule within the Antoni A and B areas. Antoni A regions are composed of more densely arranged cells with specific areas of palisading nuclei arranged in rows; Antoni B regions tend to be more hypocellular, with a loose and disorderly arrangement (Fig. 5).

Pre-operative diagnosis is difficult and further investigations are needed such as MRI, CT, US and angiography. The patient described here was examined by MRI, CT and US. Imaging studies are necessary in the diagnosis of head and neck schwannoma. The principal aim is to distinguish between a vagal or sympathetic schwannoma and a paraganglioma. On CT examination without contrast, a Schwannoma is generally hypodense compared to the muscle; with contrast, this lesion shows some degree of enhancement (Fig. 2). MRI reveals low signal intensity on T1 and high signal intensity on T2-weighted images (Fig. 3). Paraganglioma, on the other hand, is classically isodense when compared to muscle on pre-contrast CT, with more reliable homogeneous enhancement post-contrast \(^\text{9}\). The post-gadolinium MRI sequences of paraganglioma show extremely bright contrast enhancement in a characteristic “salt-and-pepper” pattern, representing the low signal intensity of vascular flow voids which is not, however, pathognomonic for paraganglioma, but may be found with hypervascular lesions \(^\text{9}\). Usually, the origin of the paraganglioma is more cranial in the superior-medium latero-cervical neck region with respect to the schwannoma; the mass, in our patient, was situated at the base of the neck in the thyroid gland area. It may also be difficult to distinguish which is the nerve of origin of the schwannoma.

The tumour may arise from the neural structures of the parapharyngeal space, including cranial nerves IX, X, XI, and XII, but, in this case, the differential diagnosis was between a schwannoma of the vagus

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**Fig. 4.** Ptosis resulting from Horner’s syndrome; NB ptosis of upper left eyelid and very mild “upside ptosis” of lower eyelid.

**Fig. 5.** Histological appearance of sympathetic chain schwannoma. (Haematoxylin and eosin stain – orig. X100). A) Peripheral areas of Schwannoma with Antoni A region (closed arrow) and Antoni B region (open arrow). B) Detail of Antoni A region with dense arrangements of cells and palisading nuclei (orig. X200). C) Detail of Antoni B region with loose organization and lower cell count (orig. X200).
nerve and a schwannoma of the cervical sympathetic chain. Often the schwannoma of the vagus nerve grows between the common carotid artery and the internal jugular vein, causing a separation between the two vascular structures. In schwannoma of the cervical sympathetic chain, no separation is observed between the internal jugular vein and the common carotid artery (Fig. 2) 11. Fine-needle aspiration, which may be conclusive in many cases of neck masses, provides far less valuable information for the compact neural tumour. The most appropriate surgical excision of the parapharyngeal tumour is the external approach not only in order to gain control of the large vessels but also to avoid injury to other nerves in the area. In our patient, the mass could not be dissected from the sympathetic chain due to involvement with the nerve that would require resection of a segment of the sympathetic chain. Post-operatively, the patient developed complete Horner’s syndrome with facial anhydrosis of the ipsilateral face (Fig. 4).

Surprisingly, despite the clinical ophthalmologic findings, the patient did not present any adverse effects or complaints. The ptosis due to paralysis of Müller’s muscle can be repaired through slight advancement of the levator aponeurosis, or resection of the conjunctiva and Müller’s muscle 12 13. Horner’s syndrome is a common sequela of the schwannoma that originates from the section of the cervical sympathetic chain and should be discussed during pre-operative counselling. Since these tumours are very rare, the physician may not be familiar with the lesion and, indeed, have limited knowledge on the subject. Close follow-up is mandatory.

References