ABSTRACT

Objectives: To present a case of cervical vagal schwannoma and describe our experience with the clinical presentation, surgical management and outcome of an elderly patient with this pathology.

Methods: 

- **Design:** Case Report
- **Setting:** Tertiary Public Hospital
- **Patient:** One

Results: A 65-year-old lady presented with a recently enlarging, pulsatile right sided neck mass that had been asymptomatic for 15 years. Contrast CT revealed a circumscribed non-enhancing heterogenous 4 x 4 x 7 cm mass splaying the right internal jugular vein and common carotid artery. A neurogenic tumour was considered, and the mass was excised from the vagus nerve with preservation of adjacent structures. Final histopathologic reading was schwannoma. However, the patient succumbed to complications following a second surgery for expanding hematoma.

Conclusion: Schwannomas are benign, slow growing tumours that arise from Schwann cells of the nerve sheath. Cervical schwannomas originating from the vagus nerve are rare but should be considered in patients presenting with solitary neck masses. Surgical extirpation is still the treatment of choice for nerve sheath tumours and recurrence is uncommon. Efforts should be made to preserve unaffected structures and patients should be counseled preoperatively on the possible high risk of morbidity especially in the elderly group where close follow up and aggressive rehabilitation should be instituted following surgery.

Keywords: Cervical schwanna, vagus nerve, neurogenic tumour; parapharyngeal space

The head and neck region is a source of swellings of various types of pathology. Nonetheless, neurogenic tumours arising from these regions are uncommon in the adult population and are rare in the pediatric group. On the other hand, these tumours are commonly found arising from the parapharyngeal spaces. The reported sites of origin of neurogenic tumours are the cranial nerves 9-12, the sympathetic chains, the cervical plexus and the brachial plexus. Cervical schwannomas are rare, slow-growing tumours of nerve sheath origin that may originate from any of these nerves. They are also commonly referred as neuromas and neurilemmomas. They are usually asymptomatic benign lesions and complete surgical resection is the treatment of choice. Malignant transformation is unusual. No gender predilection is noted and they are usually reported to occur in patients between 34 and 67 years of age. We report the case of a patient diagnosed with cervical schwannoma of vagal nerve origin, which is rare.
CASE REPORT
A 65-year-old lady with painless swelling on the right side of the neck for 15 years consulted at our centre for recent enlargement. She had no dysphagia, dyspnoea or other associated symptoms nor was there a past history of cervical trauma. Her past medical and surgical histories were unremarkable. On examination, the swelling was oval in shape and situated beneath the upper and middle third of the right sternocleidomastoid muscle. It measured 5 x 3 cm with normal overlying skin. On palpation, the mass was firm in consistency, pulsatile and could be displaced from side to side but not up and down. No bruit was noted on auscultation. Regional lymph nodes were not enlarged. The intraoral, indirect laryngeal and post nasal mirror examinations were normal. No cranial nerve deficits or Horner's syndrome were noted. Flexible nasopharyngolaryngoscopy showed no significant findings. In view of the pulsatility of the neck mass, contrasted computed tomography scan (CT) of the neck was performed and revealed a well circumscribed non-enhancing heterogeneous mass measuring 4x4x7cm over the right side of the neck. It extended from below the mandible to the level above the cricoid cartilage. The mass splayed the right internal jugular vein and right common carotid artery. The patient was diagnosed with neurogenic tumour and underwent surgical extirpation via a transverse cervical skin incision. All great vessels were isolated and controlled before the mass was excised. Intraoperatively, the tumour was tracked to the vagal nerve in origin and separated en-bloc from the vagal trunk. The glossoopharyngeal, hypoglossal, lingual, accessory nerves were all preserved. The patient was transferred to the intensive care unit after the operation. Laryngeal functions were noted to be normal. During the early postoperative period, the patient developed pneumonia and the neck wound was complicated with an expanding hematoma despite a functioning surgical drain in-situ. The patient underwent a second surgery for wound exploration, hemostasis and hematoma removal after which she was readmitted to the intensive care unit but was not recuperating well. She was kept ventilated and unfortunately succumbed to sepsis secondary to pneumonia one week after surgery. The resected specimen reported findings that were consistent with typical features of schwannoma.

DISCUSSION
Schwannomas are rare, benign tumours of nerve sheath origin that may originate from any of the cranial, peripheral or autonomic nerves.\(^1\) Over the last few decades, much literature has been written on the studies on intracranial schwannoma with regards vestibular neuroma. Nonetheless, a majority of schwannomas are non-vestibular and extracranial. Kang et al.\(^6\) reported merely 6 cases of vagal schwannoma identified over the period of 10 years retrospective study.\(^6\) Extracranial non-vestibular head and neck schwannomas usually present as asymptomatic, slow-growing lateral neck masses that can be palpated along the medial border of the sternocleidomastoid muscle. This renders preoperative diagnosis difficult as many vagal schwannomas do not present with any neurological deficits.\(^1\) Therefore, possible differentials include metastatic cervical lymphadenopathy, malignant lymphoma, carotid body tumour, branchial cyst, as well as aneurysm. When symptoms are present, hoarseness is the most common symptom for vagal schwannomas. Occasional palpation of the mass may induce paroxysmal coughing.\(^4\) Horner’s syndrome may prevail when the tumour has pressured on the cervical sympathetic chains, or when these chains are themselves the origin of the tumour.

Imaging studies play an important role in the diagnosis of head and neck schwannomas. Regardless of the nerve of origin, schwannomas in general are hypodense in relation to muscle tissue on CT without contrast. With contrast, these lesions may show some degree of enhancement, often peripheral.\(^13\) More contrast-enhancing lesions should merit differentials of possible vascular lesions. In such instances, angiography or magnetic resonance angiography may be employed to outline the feeding vessels and preoperative embolization could be planned. According to Lin et al.\(^1\) the site and the way the major neck vessels are displaced could give further clues on the type of schwanna. Vagal schwannomas typically separate the internal jugular vein and carotid arteries but do not usually widen the carotid bifurcation. On the contrary, sympathetic chain schwannomas mildly splay the carotid bifurcation but do not separate the great vessels. The splaying of the carotid bifurcation is usually more prominent in carotid body tumours. This is called the “lyre sign,” which, along with significant contrast enhancement, is rarely associated with schwannoma.\(^14\)

Differentiating amongst the origin of schwanna can be challenging. In one study, the sympathetic chain was shown to be the most usual site of origin for parapharyngeal schwannoma.\(^4\) Meanwhile, Maniglia et al. stated that the vagus nerve is the site of origin in approximately 50% of cases.\(^19\) The identification of the exact nerve trunk from which the schwanna arises can be difficult to ascertain. Anatomic evaluation by direct vision intraoperatively is an acceptable way but may not be feasible in all cases. Otherwise, the derivation may be presumed based on the presenting features or postoperative morbidity experienced by the patient.\(^5\) Nonetheless, adjacent nerves could also be damaged during surgical extirpation, resulting in error of the assumption of origin. Therefore, some diagnoses of the site of origin of schwannomas are definitive; others, presumptive.\(^15\)

Histologically, on hematoxylin and eosi staining, schwannomas show typical clusters of alternating areas of compact hypercellular with spindle shape cells (Antoni A) with areas of loose hypocellular patterns...
Furthermore, schwannomas also show positivity in S100 protein immunohistochemical studies. Although neurofibromas are of schwann cell origin, they are differentiated from schwannomas by their lack of true capsules and presence of only loose interlacing bands of spindle cells. Moreover, the nerve fibers seen in neurofibromas are scattered throughout the tumour body whereas those of schwannomas are compressed to one side.

Treatment of vagal schwannomas should encompass complete surgical extirpation with preservation of the neural pathway, as in this case. If the plane of resection is inadequate and preservation is technically difficult, the involved segment of the nerve trunk may be resected with an end-to-end anastomosis using microsurgical techniques. Invariably, this type of procedure will result in vocal cord paralysis or paresis. Therefore, aggressive voice and speech rehabilitation should entail after the surgery.

Morbidities are not uncommon after surgical resection of extracranial nerve sheath tumours. Biswas et al.\textsuperscript{16} reported high complications rates of surgery on benign extracranial nerve sheath tumours. Complications like dysphagia, dysphonia, Horner’s syndrome, facial myotonia, hypoglossal palsy, facial palsy and keloid formations have been reported.\textsuperscript{14-15} Nonetheless, neural complications could occur as a result of tumour compression. Therefore there is a need for relevant head and neck neural assessment prior to the surgery so as to correctly identify post-operative neural morbidity. Vascular complications have
CASE REPORTS

also been reported and are common after intracranial schwannoma resection but rare for extracranial schwannomas. Rapidly forming hematoma in a neck surgical wound is a dreadful complication and should merit immediate intervention. Tracheostomy may be needed if the airway is compromised. Post surgery atelectases of the lung is usually the main cause of pneumonia and aggressive chest physiotherapy should be instituted both pre and post operation. Pneumonia may also occur as a result of aspiration secondary to vocal cord palsy possibly due to direct trauma to vagal trunk, traction injury or even compression from hematoma. Because initial laryngeal functions were noted to be normal after the tumour resection in this case, the subsequent wound hematoma could have compressed on the preserved vagal trunk and precipitated aspiration pneumonia.

In conclusion, a patient presenting with solitary mass in the neck should raise the alarm of possible nerve sheath tumour and in the setting of parapharyngeal space involvement, the possibility of arising from cranial nerves, brachial plexus, cervical sympathetic chains or major peripheral nerves. Every effort should be made to preserve the nerve of origin and the patient should be counseled preoperatively on the risk of morbidity especially in the elderly group. Intervention was needed in this case in view of the rapid enlargement of the cervical swelling that could signify malignant transformation or anticipated compressive symptoms. Surgical extirpation is still the treatment of choice for nerve sheath tumours and tumour recurrence is uncommon. Nonetheless, in view of the possible surgical morbidities, close follow up and aggressive rehabilitation should be instituted following surgical treatment.

REFERENCES