Case Report

Rare cervical nerve root C2-C3 schwannoma

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INTRODUCTION

Schwannomas also known as neurilemmomas or neurinomas are benign nerve sheath tumors deriving from Schwann cells that occur in the head and neck region in 25-45% of cases.1 About 10% of schwannomas that occur in the head and neck region mostly originate from the vagus or sympathetic nervous system whereas those arising from C2 nerve root are extremely rare. Extradural schwannomas in the head and neck region are rare neoplasms. Diagnosis is established by imaging studies such as magnetic resonance imaging or computed tomography, while FNAC is used to rule out other condition. Histopathology gives definitive diagnosis. The accepted treatment for these tumors is surgical resection with preservation of the neural pathway. We present a rare case of cervical nerve (C2-C3) root schwannoma of 50 year old male who presented with right lateral neck swelling with pain radiating to right shoulder associated with right shoulder stiffness. The swelling which also had an intervertebral part was removed successfully through a posterior neck incision with no post-operative neurological symptoms.

CASE REPORT

A 50 year old male patient presented to our department with a right sided lateral neck slow growing swelling which was noticed 3 months back (Figure 1 and 2). There was no dysphagia, dyspnea or hoarseness of voice or pain, fever or trauma. It was a well-defined swelling 2×2 cms in the posterior triangle of the neck deep to the right sternocleidomastoid muscle. Carotid pulsations were displaced anteriorly. Neurological examination revealed no local or focal neurological deficits. On flexible laryngoscopy bilateral vocal cords were mobile. Ultrasound neck revealed mixed echogenic lesion mass 1.3×1.4×2.3 cms along the posterior-lateral aspect right sternocleidomastoid and superior aspect of neck muscle.

MRI cervical spine revealed well defined encapsulated pear shaped lesion in the right paraspinal region with its tail in the right C2-3 neural foramen with central cystic
component with nodular wall showing blooming on gradient images representing neoplastic etiology of nerve sheath origin like schwannoma.

Figure 1: MRI scan axial cut between C2 and C3 vertebrae showing ‘dumbbell shaped’ mass.

Figure 2: MRI scan coronal view showing mass arising from intervertebra C2-C3.

Figure 3: Intraoperative photo showing the mass relation with carotid artery.

An excision of the mass was planned by a posterior neck incision (Figure 3). Tumour was approached from posterior mid line approach to cervical spine of the neck. Right C2-C3 laminotomy and facetectomy done. A globular yellowish well defined mass visualized and the capsule (intracapsular) dissected out after taking an incision over it reaching up to the root of the swelling which was seen arising from the C2-C3 inter vertebral foramen. The tumor was identified, separated and removed. Postoperatively the patient was stable with no neurological deficits (Figure 5). Histopathological examination revealed a benign encapsulated neural tumour comprising of spindle shaped cells arranged in Antoni A and Antoni B pattern with Verocay bodies. No evidence of any atypia or malignancy was noted.

Figure 4: Specimen showing larger neck and smaller intervertebral part.

Figure 5: Benign encapsulated neural tumour comprising of spindle shaped cells arranged in Antoni A and Antoni B pattern with Verocay bodies.

DISCUSSION

Schwannomas, neurilemmomas or neurinomas are benign nerve sheath tumors deriving from Schwann cells that occur in the head and neck region in 25-45% of cases. About 10% of schwannoma that occur in the head and neck region generally originate from the vagus or sympathetic nervous system. Rarely, they arise near the vertebral foramina presenting with intraspinal and extraspinal components.  

Vagal schwannoma is typically characterized by dysphagia and hoarseness. Sympathetic schwannoma is characterized by Horner’s syndrome. In most cases, however, there are no symptoms, thus it is difficult to identify the neurological origin based on the physical examination.
As in our case though tumour was arising from C2 nerve root which had intervertebral extraspinal part, the patient had no preoperative neurological deficit. Imaging diagnostic modalities like USG and MRI offer great help in identifying the tumor and its correlations with surrounding vascular structures, muscles and nerves. Previously, to prevent the recurrence of tumors, radical dissection including the neuroprogenitor cells was performed. Even in cases in which recovery was achieved following the nerve transplantation or primary anastomosis, preservation of the neurological function was not to be expected. Most of the neuroprogenitor fibers do not run through schwannoma and they pass over the tumor capsule. Most schwannomas are encapsulated. In cases where the nerve fibers surround the surface of tumors, the intracapsular enucleation can be performed while preserving the nerve fibers. According to the study by Valentino et al, intracapsular enucleation while preserving the nerve fibers preserved its function by more than 30% when compared to tumor resection with primary anastomosis. According to Zbären et al, there was no significant difference in the recurrence rate between the total tumor resection including nerve fibers and the intracapsular enucleation. In cases where partial removal of the tumor was performed, however, the recurrence rate has been reported to rise. In our case, we found that swelling was dumbbell shaped which was successfully removed with intracapsular dissection through neck incision along with intervertebral part doing laminectomy. Postoperatively no neurological deficit was noticed. Microscopically, schwannomas are encapsulated, solid or cystic tumors. They can be composed of two cellular zones: Antony type A, densely arranged with spindle-shaped Schwann cells and areas of palisading nuclei, Verocay bodies and Antony B, characterized by a hypo cellular arrangement and a large quantity of myxoid tissue.

CONCLUSION

Extracranial schwannomas in the head and neck region are rare neoplasm. Diagnosis is established by imaging studies such as magnetic resonance imaging or computed tomography, while FNAC is used to rule out other condition. Histopathology gives definitive diagnosis. The accepted treatment for these tumors is surgical resection with preservation of the neural pathway.

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REFERENCES