Tibial nerve schwannoma: short review of surgical management

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ABSTRACT
Schwannoma is a benign, solitary nerve sheath tumour and accounting for about 5% of soft tissue tumours. It can occur along the peripheral nervous system in any part of body. It presents as a painless, swelling. We report an adult male presented with tibial nerve schwannoma underwent successful surgical excision. However, differentiation with neurofibroma is very important as surgical planning and prognosis is quite different. In lower limb usually incidence of neurofibroma is higher in contrast of upper limb. Pertinent literature and management are briefly discussed.

INTRODUCTION
Schwannoma represents a benign peripheral nerve tumour, originating from Schwann cells [1], [2]. It usually presents as a solitary, slow growing mass. It accounting for about 5% of all soft tissue tumours. It can present with pain, paraesthesia or rarely with neurological deficit [3], [4], [5].

CASE ILLUSTRATION
An adult male reported presented with complaints of painless mass with paraesthesia and difficulty in sitting on chair for two-years. Local examination showed presence of mass in the popliteal fossa, about size of 4cm X 5 cm, no compressible, nonpulsatile being mobile along transverse axis but no mobility in craniocaudal axis. A magnetic resonance imaging was carried out to ascertain the nature, revealed presence of a mass lesion causing expansion of tibial nerve. (Fig-1) A provisional diagnosis of peripheral nerve sheath tumour was made and planned for surgical excision. He underwent micro-surgical total excision, intraoperative expansion of the nerve was observed, and nerve fascicles were carefully separated from mass lesion, with electrophysiological nerve monitoring. He had relief in pain and...
paraesthesia in post-operative period. Histopathology of resected specimen was suggestive of schwannoma.

**DISCUSSION**

Tibia nerve is continuations of larger terminal branch of the sciatic nerve with root values of L4, L5, S1, S2, and S3 [1], [5], [6], [7]. Tibial nerve usually lies superficial to the popliteal vessels, extending from the superior angle to the inferior angle of the popliteal fossa, crossing the popliteal vessels from lateral to medial side [8].

Peripheral nerve sheath schwannoma symptoms are related to alteration in the function of nerve and surrounding muscle and neurovascular bundles, and mostly commonly present with paraesthesia or pain of insidious onset and progresses slowly [2]-[4]. Pain is a much more common symptom than focal motor or sensory deficits. Physical examination may reveal the presence of a mass along the course of the nerve, tender, usually mobile along the transverse axis but limited along the longitudinal course of the nerve, and positive Tinel sign [6], [9].

However, pre-operative confirmatory diagnosis of schwannoma usually not possible in most cases but can help in delineating shape, size, location, extent and relation with parent nerve and adjacent neurovascular structures and muscle. Imaging plays a limited role in distinguishing among various types of peripheral nerve sheath tumours. Magnetic resonance imaging may show presence of fusiform mass with characteristic tapering cephalad and distal ends, fasciculation sign and split fat signs [3], [8], [9]. The mass is well-circumscribed and eccentrically placed, and showing isointense signal on T1-weighted images and T2 weighted images shows hyperintense signal and peripheral rim demonstrate hypo-intensity signal representing capsule [3], [5].

After confirmation of diagnosis management of peripheral nerve schwannoma is usually surgical except when the mass is very small and not causing any physical disfigurement.

Treatment of epineurium encapsulated tumour is microsurgical excision with careful preservation of the nerve fascicles. Histopathological examination of specimen provides definitive diagnosis [4], [5]. Kim et al. analysed 397 cases of peripheral nerve sheath tumour, out of which 91% were benign and the rest were malignant. A total of 251 were located in the brachial plexus region or upper limb. The peripheral nerve sheath tumor involving lower-limbs included 53 cases of neurofibroma and 32 cases of schwannomas [5]. Typically showing the incidence of schwannoma is less than neurofibroma.

Recurrence is uncommon following total surgical excision. Usually surgical excision provides good outcome in view of its benign biological nature and malignant transformation is extremely rare.

**CONCLUSION**

Tibial nerve schwannoma is rarer entity compared to neurofibroma and prognosis and surgical planning should be discussed and prognosticated to patient as imaging may also not definitely distinguish between neurofibroma, schwannoma. However electrophysiological monitoring is an important aid in preserving neurological outcome. Hence, every surgical team member should be always considering the possibility of neurofibroma, schwannoma.
References