Adulthood Benign Triton Tumor Developed in the Orbit

Dong Hyeon Bae, M.D., Choong Hyun Kim, M.D., Jin Hwan Cheong, M.D., Jae Min Kim, M.D.

Department of Neurosurgery, Hanyang University Guri Hospital, Guri, Korea

Benign triton tumor (BTT) or neuromuscular hamartoma is an uncommon tumor composed of mature neural and well-differentiated striated muscular elements. Its development is exceptionally rare in the adult and head region. This report describes a case of adulthood BTT that occurred in the orbit. The patient was a 53-year-old woman who presented with right periorbital swelling and pain in eyeball over 2 months. Magnetic resonance imaging revealed a well-enhancing mass surrounding optic nerve and ocular muscles in the right retrobulbar area. The tumor was subtotally removed via transcranial approach. Its pathological diagnosis was confirmed to be a neuromuscular hamartoma. She developed diplopia postoperatively. Adulthood BTT should be considered in the differential diagnosis of head and neck tumors. It is also important to make adequate therapeutic strategy to avoid postoperative neural dysfunction.

Key Words: Adult · Benign triton tumor · Head · Neuromuscular hamartoma · Surgery.
was improved, but she developed diplopia possibly by the injury of medial rectus muscle. Even though she has been undergone reconstructive surgery for the medial rectus muscle by ophthalmology department, her diplopia was persisted. There was no recurrence of tumor about 10 years after surgery.

DISCUSSION

BTTs are extremely rare reported that less than 20 cases have been reported in the English language literatures. The name triton has been originated from the amphibian triton salamander, in which the normal nerve seems to induce regeneration of skeleton muscle. These tumors commonly occur in infants and children, however, it has been rarely reported in adults. The most common sites of invasion are the large peripheral nerves, especially the brachial and sciatic nerves. A few cases have been reported in the head and neck region. These tumors have about 2.4 : 1 female predominance in literature reviews.

Daley et al. have analyzed previous 9 cases developed in the head and neck of children and adults, and they suggested that BTTs have been categorized into two groups by their location: 1) a more aggressive central type and 2) a non-aggressive peripheral type. A more aggressive central type involves large intracranial nerve or infratemporal fossa nerve trunks with extension into multiple adjacent spaces, sometimes associated with muscle atrophy and weakness, and usually associated with destructive growth. Involvement of the fifth cranial nerve in or near the middle cranial fossa (specifically Meckel’s cave) with extension through the foramen ovale into the infratemporal fossa is most common. These cases typically appear infants and children, and require extensive surgical treatment.

A non-aggressive peripheral lesions are subcutaneous or submucosal, non-encapsulated lumps that are asymptomatic, non-destructive, and enlarge slowly, if at all. They are found at a later age, including adulthood. Surgical excision is simple and curative. Daley et al. also reviewed 9 cases with BTTs in the head and neck, and residual or recurrent cases have been developed in one case of 5 central BTTs associated with trigeminal nerve and only case of 4 peripheral BTT, respectively. Our case may be categorized into the former type. Four theories have been suggested to explain the origin of this unusual tumor; 1) Hamartomatous growth of muscle spindles: Markel and Enzinger suggested that these lesions may represent only muscle spindle hamartomas because of its structural resemblance to the neuromuscular spindle organ. 2) Entrapment of skeletal muscle fibers within nerves during embryogenesis: This theory was initially postulated by Markel and Enzinger after treating a patient with a tumor composed of mature muscle cells and fibrous tissue in the vicinity of the sciatic nerve. Louhim and Rapola suggested that mesenchymal tissue might become incorporated by the extending peripheral nerves and ensheathed by the neurilemma. 3) Differentiation of neuroectoderm into skeletal muscle: This theory was proposed initially by Masson. He suggested that neuroectodermal cells could undergo mesenchymal differentiation. This is based on his observation of the histological development of the cranial meninges and intraocular muscles, both of which are derivatives of the ectodermal cells, the ectomesenchyme. 4) Epigenetic or acquired alteration of the motor end-plate: This recent theory was suggested by Daley et al. They propose that there is an alteration, either epigenetic (infants) or acquired (adults), in which there is a motor end plate/synaptic junction defect. Thereafter, this leads to reactive proliferation of neural and skeletal muscle tissue in an organoid pattern. They indicated two meaningful points: 1) There would be no contraction of the tumor mass despite the abundant highly innervated skeletal muscle. 2) The nerve involved would be a motor nerve. No single theory adequately explains all of reported neuromuscular hamartoma.

Clinical manifestations are variable ranged from asymptomatic to pain or development of neurologic deficits. Most BTTs do not cause symptom, however, depending on the site of involvement, pain, neurologic dysfunction in the form of muscle weakness, paraesthesia, and headache including oculomotor ophthalmoplegic migraine, could be occurred. Radiographic features of BTT show a wide spectrum. Most tumors enhance after intravenous contrast infusion on the computed tomography scans and MR images. Histopathologically, these tumors are multi-nodular, subdivided by collagenous connective tissue into smaller nodules and fascicles. The fascicles are comprised of mature striated muscle fibers intimately associated with nerve fibers, and enclosed within the same perimysial-like fibrous sheath.

Fig. 2. Photomicrographs of the resected lesion. A: H & E staining shows nerve fibers admixed with skeletal muscle (×200). B: Immunohistochemical staining reveals diffuse immunoreactivity to S-100 protein suggesting neural elements (×200). C: Luxol-fast blue staining demonstrates dark bluish colored muscle fibers in the striated fashion (×200).
Treatment options are excision with more aggressive central type and conservative for the peripheral tumors. Incomplete resection could relieve symptoms. Post-resection nerve palsies are always crucial, thus, preservation of neural function is essential. In our case, patient developed diplopia postoperatively. We must consider whether aggressive treatment or not for patients satisfaction and favorable prognosis. Although prognosis is good in the most cases, recurrence has been reported. Therefore, we should be kept in mind the close follow-up of patients.

CONCLUSION

Although this tumor rarely occurs, its frequent connection with large nerves or cranial nerves often makes complete resection difficult or impossible. Moreover, surgical treatment of these tumors may develop residual neural dysfunction. In consideration of their extremely low recurrence and benign lesion, BTT should be removed subtotally and save the involved nerve to avoid neurological deficits.

References
1. Akimoto J, Fukami S, Hashimoto R, Haraoka J: Neuromuscular hamartoma is a possible primary pathology of oculomotor ophthalmoplegic migraine. Cephalalgia 32:171-174, 2012
2. Amita K, Shankar SV, Nischal KC, Basavaraj HB: Benign triton tumor: a rare entity in head and neck region. Korean J Pathol 47:74-76, 2013
3. Bonneau R, Brochu P: Neuromuscular choristoma. A clinicopathologic study of two cases. Am J Surg Pathol 7:521-528, 1983
4. Castro DE, Raghuram K, Phillips CD: Benign triton tumor of the trigeminal nerve. AJNR Am J Neuroradiol 26:967-969, 2005
5. Daley TD, Darling MR, Wehri B: Benign Triton tumor of the tongue. Oral Surg Oral Med Oral Pathol Oral Radiol Endod 105:763-766, 2008
6. Demir Y, Uluoglu O, Ozmen S, Boyacioglu ZM, Atabay K: Neuromuscular hamartoma in the mental region. J Oral Maxillofac Surg 61:397-400, 2003
7. Louhim I, Rapola J: Intraneural muscular hamartoma: report of two cases in small children. J Pediatr Surg 7:696-699, 1972
8. Markel SF, Enzinger FM: Neuromuscular hamartoma--a benign "triton tumor" composed of mature neural and striated muscle elements. Cancer 49:140-144, 1982
9. Masson P: Tumeurs Humaines: Histologie, Diagnostics et Techniques, ed 2. Paris: Maloine, 1956, pp973-975
10. Moore KL, Persaud TVN: The Developing Human: Clinically Oriented Embryology, ed 5. Philadelphia, PA: Saunders, 1993, pp375-423
11. O’Connell JX, Rosenberg AE: Multiple cutaneous neuromuscular choristomas. Report of a case and a review of the literature. Am J Surg Pathol 14:93-96, 1990
12. Oppen RS, Harden SP, Argent JD: Neuromuscular hamartoma: imaging features of a rare paediatric craniofacial tumour. Pediatr Radiol 33:50-52, 2003
13. Tiffee JC, Barnes EL: Neuromuscular hamartomas of the head and neck. Arch Otolaryngol Head Neck Surg 124:212-216, 1998
14. Tobias S, Kim CH, Sade B, Staugaitis SM, Lee JH: Neuromuscular hamartoma of the trigeminal nerve in an adult. Acta Neurochir (Wien) 148:83-87; discussion 87, 2006
15. Vajramani G, Devi IC, Santosh V, Hegde T, Das BS, Das S, et al.: Benign triton tumor of the trigeminal nerve. Childs Nerv Syst 15:140-144, 1999
16. Weiss SW, Goldblum JR, Enzinger FM: Enzinger and Weiss’ Soft Tissue Tumors, ed 5. Philadelphia, PA: Mosby Elsevier, 2008, pp834, 917-919