Case report

Case report of eyelid schwannoma: A rare presentation in a child

Mantapon Ittarat\(^a,\)b, Pakamat Srihacha\(^a\), Sunee Chansangpetch\(^b,\)c,\(^*\)

\(^a\) Department of Ophthalmology, Prasat Hospital, Surin, Thailand
\(^b\) Faculty of Medicine, Chulalongkorn University and King Chulalongkorn Memorial Hospital, Bangkok, Thailand
\(^c\) Glaucoma Research Unit, Chulalongkorn University, Bangkok, Thailand

ABSTRACT

Purpose: To report a rare case of solitary eyelid schwannoma in a Thai child without clinical manifestation of neurofibromatosis.

Observations: A 9-year-old Thai boy presented with an isolated painless mass on the left lower eyelid. The mass had gradually grown for 2 years. He denied a history of trauma or previous eyelid surgery. A clinical examination revealed no clinical features of neurofibromatosis. The provisional diagnosis at that time was a sebaceous cyst. However, excisional biopsy showed an encapsulated tumor characterized by interlaced spindle-celled fasciculi with focal palisading of nuclei arranging in Antoni A and Antoni B patterns. The immunocytochemistry was strongly positive for S-100 protein reaction. The diagnosis of schwannoma was made, with no recurrence at.

Conclusions and importance: Schwannoma of the eyelid is extremely rare in children, accounting for less than 0.1% of all eyelid tumors. Combined with clinical features, histopathologic and immunohistochemical analyses emphasize the disease entities. These findings may extend the knowledge on experiences of schwannoma in children.

1. Introduction

Schwannoma is uncommonly seen in ocular tissue. The eyelid is an extremely rare location for schwannoma, especially in children. The purpose of this article is to report a rare case of an isolated eyelid schwannoma in a Thai child.

2. Case report

A 9-year-old Thai boy presented at the hospital in February 2016 with a painless solitary mass on the left lower eyelid. The patient reported a two-year history of a slowly growing mass. The tumor produced no discharge or redness. He denied a history of previous trauma or surgical excision. The child was otherwise normal, with no clinical finding suggestive of systemic tumor. He had no family history of neurofibromatosis. On examination, there was a round, non-erythematous mass located at the lateral third of lower eyelid margin. The mass was firm, smooth surfaced, and non-tender on palpation. The lesion laid superficially and was movable under the skin. The approximate size on examination was 1 cm in diameter, with minimally visible protrusion from the skin. Other aspects of the ophthalmic exam were within normal limits. The presumptive diagnosis at that time was sebaceous cyst. An excisional biopsy was performed via anterior dissection in the skin under local anesthesia. Gross histopathological examination showed well encapsulated, smooth consistency, tan tissue. The mass measured 0.8 × 0.5 × 0.5 cm. Microscopic section revealed a well-defined mass with a thin capsule consisting of the spindle to oval nuclei with eosinophilic fibrillar cytoplasm. The histology also showed a dense hyalinized and fibrillary area (Antoni A pattern) alternated with a loose hyalinized and fibrillary area (Antoni B pattern). A mitotic figure was not seen. Hyalinized vascular walls were noted. A nerve of origin was not discernible. The immunocytochemistry for S100 protein showed a strongly positive reaction (Fig. 1). The final diagnosis was a benign schwannoma of the eyelid. The post-operative wound healed completely without any postoperative complication. No recurrence was observed at 18-months follow up.
3. Discussion

Schwannomas are Schwann-cell tumors that arise from peripheral or cranial nerves. Solitary lesions can occur sporadically; multiple schwannomas can be associated with neurofibromatosis 2. Schwannoma that involves the eye most commonly develops in the orbit, accounting for around 1% of orbital tumors. Less frequent involvement has been found in uveal tract, conjunctiva, and sclera.

Eyelid schwannoma is extremely rare. Siddiqui et al. presented 5 cases of eyelid schwannoma. Four out of the five cases were in late adulthood (age 53–63). One case was in young adulthood (age 19). Lopez-Tizon et al. reported another 2 adult cases, which they noted as the only 2 cases of schwannoma out of 2400 eyelid neoplasms observed in their 11 years of experience (1995–2005), indicating a prevalence of 0.1% among eyelid tumors.

Eyelid schwannomas in children are rarer than in adults. Only 3 cases have been published so far. None were from Southeast Asian region. The clinical presentations of all 3 cases and our current case are summarized in Table 1. The diagnosis was confirmed with a strong reactivity to S100 protein, which is considered a hallmark feature of schwannomas.

The histological finding in our case also followed a previous pattern occurring on eyelid margins reported by Stagner and Jakobiec in 2016. It is important to note that, in eyelid peripheral nerve sheath tumors, S100 positivity in immunostaining can be demonstrated, not only in schwannoma, but also in solitary circumscribed neuroma (SCN), another eyelid tumor that has been reported in adults. Schwannoma and SCN share the common characteristic of densely packed spindle-cell and S-100 positivity. However, the SCN histologically presents a more prominent axon, less palisading, and insubstantial capsule; the schwannoma, as in this present report, has a distinct pattern of fascicular cells alternating myxoid areas in conjunction with a complete, thick, and hyalinized capsule. Immunostaining for neurofilament antibody and immunostaining for perineurial cells (i.e., epithelial membrane antigen staining or glucose transporter-1 staining) might aid in differentiating these two eyelid nerve sheath tumors.

Since eyelid schwannomas are rare and without characteristic clinical features, definitive diagnosis depends on histopathology. Fortunately, schwannomas rarely undergo malignant transformation.

4. Conclusions

Schwannoma of the eyelid in children is extremely rare, accounting for less than 0.1% of all eyelid tumors. Combined with clinical features, histopathologic and immunohistochemical analysis emphasize the disease entities. These findings may extend the knowledge on experiences of schwannoma in children.

Patient consent

The patient’s assent for publication was obtained from a patient. Written informed consent was also obtained from the parents.

Acknowledgments and disclosures

Funding

No funding support.

Conflicts of interest

The following authors have no financial disclosure.
Authorship

All authors attest that they meet the current ICMJE criteria for Authorship.

Acknowledgements

Phanchanut Mahantassannapong; Pathologist.

References