ABSTRACT – Schwannoma is a benign encapsulated nerve sheath tumor composed of Schwann cells, that may arise anywhere along the course of a nerve. Cutaneous schwannomas, associated with peripheral nerves, usually present as a slow-growing well-circumscribed nodule, localized in the dermis or subcutaneous tissue. They are generally asymptomatic; however, they may become painful by the nervous compression. Since the clinical signs are non-specific, histology is required for a definitive diagnosis. Herein, we present a case of a solitary cutaneous ancient schwannoma of the ear, a histopathological variant of schwannoma with distinctive morphological characteristics.

KEYWORDS – Neurilemmoma; Skin Neoplasms.

RESUMO – O schwannoma é um tumor benigno encapsulado da bainha nervosa composto por células de Schwann, que pode surgir em qualquer lugar ao longo do curso de um nervo. Schwannomas cutâneos, associados a nervos periféricos, apresentam-se geralmente como nódulos solitários bem delimitados de crescimento lento, localizados na derme ou tecido celular subcutâneo. Apesar de geralmente assintomáticos, estes tumores podem tornar-se dolorosos pela compressão nervosa. Os seus sinais clínicos são inespecíficos, pelo que a histologia é necessária para o diagnóstico definitivo. Apresentamos um caso de schwannoma cutâneo antigo solitário do pavilhão auricular, uma variante histopatológica de schwannoma com características morfológicas distintivas.

PALAVRAS-CHAVE – Neoplasias da Pele; Neurilemmoma.

INTRODUCTION

Schwannoma, also known as neurilemmoma, is a benign encapsulated neoplasm composed of Schwann cells that originates from nerve sheaths.1 Commonly found in association with the cranial nerves (especially the acoustic nerve), schwannomas may arise anywhere along the course of a nerve, including peripheral nerves.2 Cutaneous schwannoma usually presents as an asymptomatic well-circumscribed skin-colored nodule of slow growth, located in the dermis or in the subcutaneous tissue.1,2 The most common sites are the head and neck and the flexor surfaces of the extremities.3,4 Although they commonly occur as single lesions (90%), they can be associated with several central neurological tumors (usually meningiomas, 5%), neurofibromatosis type 2 (3%), or appear as multiple lesions (schwannomatosis, 2%).3 Herein, we present a case of a solitary cutaneous ancient schwannoma of the ear, a histopathological variant of schwannoma with distinctive morphological characteristics.

CASE REPORT

An otherwise healthy 69-year-old caucasian woman was observed for a painless skin nodule on her left ear, that had developed slowly over a period of more than eight years. The patient denied previous local trauma or any other similar lesions. Physical examination revealed a well-defined, flesh-colored, cutaneous nodule, 1 cm in diameter, with elastic consistency, non-painful on palpation, located on the patient’s concha of the left ear (Fig. 1). There were no other relevant skin lesions.

Surgical excision with clear margins was performed. Histological examination revealed an encapsulated tumor located in the upper dermis composed of mostly dense cellular areas (Antoni type A), alternating with hypocellular myxoid areas (Antoni type B) (Fig. 2 A,B). On high-power magnification, the tumor was composed of spindle cells with back to back nuclei arranged in parallel rows without mitoses. There were some enlarged, hyperchromatic, and bizarre nuclei (Fig. 2 C). Conspicuous small blood vessels were seen. Immunohistochemically, the tumor cells were strongly and diffusely immunorreactive for S-100 protein (Fig. 2 D), and the tumor capsule cells stained for EMA (Fig. 2 E). Alcian blue showed staining of the myxoid matrix (Fig. 2 F). Based on histological examination, a diagnosis of ancient schwannoma was made.

DISCUSSION

Verocay first described schwannomas in 1908.3 They represent approximately 5% of all benign soft-tissue tumors and they are the most common peripheral nerve tumor.4 Schwannomas occur most commonly during the fourth and fifth decade of life, without significant evidence of sex predilection.3,5
In general, cutaneous schwannomas are asymptomatic. However, as they grow larger, pain, paresthesias, and tenderness can occur, associated with the compression of the nerve of origin or adjacent structures.

The clinical diagnosis is challenging due to the non-specific presentation, becoming necessary to exclude other well-circumscribed skin masses such as lipoma, angiolipomas, leiomyomas, cysts, neurofibromas, and neuromas.

Histology is required to confirm the diagnosis. The tumor is typically well-encapsulated by perineurium and is classically characterized by a biphasic pattern: areas of high cellularity (Antoni A) that alternate with loosely myxoid edematous areas (Antoni B) (Fig. 3). In Antoni A areas, uniform spindle cells are clustered in stacks and arranged back to back, creating palisades of stacked nuclei between which the cell processes are fused into eosinophilic masses, called Verocay bodies (Fig. 3 A, B). In Antoni B areas, cells are more loosely arranged, separated by spaces filled with mucin. (Fig. 3 C, D). Most schwannomas display both areas, and although intermingled, both region types usually appear fairly demarcated from one another.

The immunohistochemical profiles of schwannomas are typically positive for S100 protein and collagen type IV, while the fibroblast in the perineurium sheath is positive for epithelial membrane antigen (EMA). As most of schwannomas are devoid of axons, they are characteristically negative for neurofilaments.

Ancient schwannoma is a rare subtype of schwannoma, which is usually found in long-standing lesions, characterized by pronounced degenerative changes. In addition to the classical features exhibited in schwannoma, ancient type displays the presence of enlarged, hyperchromatic and bizarre nuclei and the absence of mitosis. Cystic formation, calcification, haemorrhage and hyalinization may also be demonstrated. The characteristic histologic appearance seen in ancient schwannoma can make pathologic diagnosis difficult and lead to an erroneous diagnosis of malignancy.

Although ancient schwannomas are a well-known entity that can involve the head and neck region, with reports in a variety of locations, such as the orbit, infratemporal fossa, submandibular gland and parotid gland, but it is extremely rare for them to be found in the skin of the ear.

The treatment of choice for cutaneous schwannoma is surgical excision, indicated in the presence of symptoms or for cosmetic reasons. Local recurrence is infrequent as the tumor is usually well-encapsulated and thus easily resectable. Malignant
transformation of solitary schwannomas is exceedingly rare. The prognosis of ancient schwannoma is similar to the classic variant so atypia should not raise alarm for malignant transformation.

In conclusion, cutaneous schwannomas are benign skin tumors with a very characteristic histology but non-specific clinical presentation. We highlight the importance of recognizing the histopathological variant of ancient schwannoma to avoid an erroneous diagnosis of malignancy and unnecessarily aggressive treatment.

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