CASE REPORT

Solitary neurofibroma of the palate

Neurofibroma isolado em palato

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Introduction

Neurofibromas are benign tumors arising from peripher- nal nerve sheaths that affect the head and neck region only rarely, and consist of Schwann cells, perineural cells, and a variable amount of mature collagen. They are classified as solitary when not associated with any syndrome or multiple when associated with autosomal dominant neurofibromatosis syndrome or type III multiple neoplastic syndrome; they can also be classified as myxomatous (solid, central, diffuse) or plexiform (peripheral). When not associated with syndromes, they rarely occur in the oral cavity, and then most commonly in the tongue.

Case report

A 54-year-old female patient with melanoderma without systemic involvement, was referred for evaluation of an asymptomatic intraoral pedunculated nodular lesion located on the palate, similar in color to the adjacent mucosa, that measured approximately 8 mm (Fig. 1A), exhibited an uneven surface, had a fibrous consistency, and had been present for three months. Extraoral physical examination revealed characteristic stigmata of neurofibromatosis, involving the chest. An excisional biopsy and histopathological examination were performed, and revealed mucosal fragments covered with stratified squamous epithelium (ortho or parakeratinized), with acanthosis.

The lamina propria showed spindle cell proliferation, sometimes with undulated nuclei amid marked collagen fiber deposition, arranged in variable-sized bundles in addition to mast cells (Fig. 1B-C). The report suggested a diagnosis of neurofibroma, with the analysis revealing strong immunohistochemical staining for S-100 protein (Fig. 1D). The patient is undergoing clinical follow-up and has shown no signs of recurrence (Fig. 1E).

Discussion

It is believed that the frequency of oral solitary neurofibromas not associated with neurofibromatosis-1 is low, with only a few cases located on the palate, despite a more common occurrence in the tongue. They are slow-growing, nodular, sessile, and mobile tumors, usually painless, although pain or numbness may occur due to nerve compression. Cherrick and Eversole observed a higher
Incidence in females. Chen and Miller⁵ reported that oral neurofibromas affect individuals between 9 and 72 years of age. Although some lesions require imaging tests to determine possible extension, conventional histological analysis is conclusive when proliferation of Schwann cells, perineurial cells, and fibroblasts are observed amid a stroma, sometimes myxomatous and microvacuolated.⁴

The distinction between solitary neurofibromas and those associated with neurofibromatosis is that the latter, although unencapsulated, show clear distinction between the lesion margin and adjacent tissue.¹ The treatment is surgical removal, because of an uncertain risk of recurrence or malignant transformation; the prognosis is excellent.⁴

Final considerations

Considering that isolated neurofibromas in the oral cavity are rare, the importance of a thorough clinical examination by professionals familiar with examining and treating patients with oral cavity lesions, such as dentists, maxillofacial surgeons, head and neck surgeons, otolaryngologists, as well as dermatologists, is reinforced. Neurofibroma not associated with neurofibromatosis-1 should be included in the differential diagnosis of hyperplastic lesions of the oral cavity, although they are rarely described in such location.

Conflicts of interest

The authors declare no conflicts of interest.

References