Neurofibromatosis type I with an Rare Oral Manifestation

Case Report

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Abstract

Neurofibromatosis type I is one of the most common genetic diseases and it may have oral manifestation.

Objective: To report a case about a 33 years old female patient diagnosed with neurofibromatosis type I in the skin.

Case Report: In the extraoral clinical examination were observed subcutaneous nodules and café-au-lait spots, in different sizes around the body. In the intraoral examination was diagnosed the presence of rounded nodule in the hard palate. The excisional biopsy of the lesion was performed and the histopathological examination suggested a diagnosis of neurofibroma.

Conclusion: The case report emphasizes the importance of a detailed anamnesis and a thorough physical examination, so that systemic diseases can be considered in oral diagnosis.

Keywords: Neurofibromatosis; Neurofibroma; Mouth.

Introduction

Neurofibromatosis type I (NF1), classically described by Friedrich Daniel von Recklinghausen in 1882, is the most common autosomal genetic disease with dominant inheritance of the human species. The prevalence of NF1 is one case per 3,000 births, and may be appear at birth or late, there is no evidence of predilection for gender or ethnicity [1].

This neuro-skin disorder is caused by a mutation of the NF1 gene located on the long arm of the chromosome 17q11.12, causing the loss of function of the protein neurobromin, thus leading to multiple manifestations of the phenotype of NF1 [2].

Clinically, there are seven major components of the syndrome, including the presence of cafe-au-lait spots, two or more neurofibromas of any type or a plexiform neurofibroma, freckles in the axillary or inguinal region, optic gliomas, two or more Lish nodules, bone lesions, these are the primary degree relative with NF1 according to the preceded criteria. If the individual has at least two of these characteristics, he can be diagnosed with NF1 [1-3].

Despite being more prevalent in skin, NF1 may be manifested in the oral cavity in about 10% of the cases, particularly in the tongue [3, 4]. However, changes such as macroglossia, increased of fungiform papilla and mandibular foramen, unilateral or bilateral intraosseous lesion and delayed tooth eruption can also be found in patients with NF1 [5].

The prognosis of NF1 is unpredictable, showing the possibility of various complications [2]. The malignant transformation of NF1 has been reported, mainly evolving to malignant neoplasm of peripheral nerve sheath and neurofibrosarcoma [6, 7]. However, the malignant transformation is poorly understood, and in particular the mutation in the p53 gene on chromosome 17p is associated with this progression [7].

Thus, the aim of this article is to present one case of a classic neurofibromatosis type I, with clinical evidence similar to the literature, but showing an unusual oral manifestation.
Case Report

A 33 years old female patient, melanoderma, sought the Dentistry Clinic of Integrated Studies XIV at the State University of Feira de Santana - Bahia, complaining about "the presence of lumps in the roof of the mouth."

In the anamnesis the patient reported the presence of nodules around the body since birth, and a surgery to remove a nodule on the right side of the face about 15 years ago, which removed the ear (Figure 1). The histopathological diagnosis of the lesion localized in her face was neurofibroma. Also, the patient reported the appearance of nodule in the mouth for a year and two months ago (Figure 2). In the medical history, the patient denied having health problems and said that there is no other case of neurofibromatosis type 1 in her family.

In the extraoral clinical examination was observed facial asymmetry (Figure 3), scoliosis, short stature, subcutaneous nodules of different sizes around the body, the plexiform neurofibroma in the arms (Figure 4) and café-au-lait spots on the limbs lower (Figure 5). The intraoral examination showed macroGLOSSIA, an increase of fungiform papillae and the presence of rounded nodules in the hard palate measuring about 5 mm, showing regular contour, pink color, soft consistency and exophytic growth.

An excisional biopsy of the lesion and the histopathologic examination was performed and it showed mesenchymal tissue fragments with fusiform cells and corrugated core arranged in interlocking beams, permeated by connective tissue stroma composed of delicate collagen bundles (Figure 6), suggesting the diagnosis of neurofibroma.

This case has the approval of the ethics committee with CAAE: 0086.059.000-08 the information in this article has permission from the patient to divulgation.

Discussion

NF1 is an autosomal genetic disease with dominant inheritance; it is common to have more than one case in the same family [1]. In this case report, the patient is unaware of another family member carrier of the disease.

The initial suspected diagnosis was based on clinical manifesta-
The clinical findings in the patient corroborate with those mentioned by the literature, once she has the presence of cafe-au-lait spots in the lower limbs, as well as plexiform neurofibroma in the upper limbs, which is a pathognomonic characteristic for this disease [1-3]. It was also noticed a hemi facial hypertrophy, scoliosis and short stature [9].

The first diagnosis of NF1 occurred when the patient was between the first and second decade of life after a surgical removal of a lesion in the face. The diagnosis of this disease in common at this age group, because during puberty occurs an exacerbation in the number and size of neurofibromas, which leads the patient to seek a professional follow-up [3].

The oral manifestation of NF1 is controversial. Some authors admit this is a common expression of the disease, which occurs in 72% of cases [6], others authors believe this is rare, affecting only 10% to 26% of cases [1]. In this case report the oral manifestations occurred three decades after the appearance of the characteristics in the skin, showing neurofibromas on the palate, which is an unusual site [4]. In the intraoral examination was also observed macroglossia, which is in accordance with the aspects commonly found for the disease in the mouth [9].

Currently, the patient is being followed in a close preservation, since some authors admit the possibility of malignant transformation of NF1. This transformation remains controversial in the literature, ranging from 3 to 5% (9) or affecting up to 26% of cases [10]. This disparity suggests a certain degree of unaware about the malignant mechanism of NF1, being essential that more studies be conducted about this topic.

The treatment of choice is limited to provide a better quality of life for the individual affected by NF1. The surgical excision of the lesions is indicated in the most cases [9], which was the proposed treatment for this case. Since the aesthetic deformity that the neurofibromas can provide, it may result in a psychosocial impairment of the patient, so that it is necessary a multidisciplinary follow-up [3]. In this case, the patient showed up quite shy, reporting she never did any kind of work throughout life, so it was proposed to her a psychotherapeutic therapy as a treatment.
Conclusion

As it has been seen, the characteristics findings in this case are similar to those reported in the literature, differing only by the rare anatomical location on the palate. Also, it is important to highlight the importance of a detailed anamnesis and physical examination, once systemic diseases may have oral manifestations. It is important to emphasize the necessity of a follow up in these patients diagnosed with oral NF1, because the malignant transformation is possible and the rates of recurrence of the lesions in skin and mouth are high.

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