AN UNUSUAL CASE: NEUROFIBROMATOSIS TYPE 5

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Abstract

Neurofibromatosis is a disease which includes 5 types in its classification. It has been firstly described by Robert Smith in 1849, but its name comes from the first publication in 1882, in which Von Recklinghausen convinced the scientific and medical world that Neurofibromatosis (NF) was a distinct entity.

Cafe-au-lait spots, cutaneous neurofibromas and Lisch nodules are the characteristics of the most common forms. Bilateral acoustic neuromas distinguish the type 2. Type 3, Riccardi type and intestinal type are briefly described.

Neurofibromatosis type 5 (NF5) is characterized by cafe-au-lait spots restricted to one area of the body. We report oro-dental phenotype of a patient affected by NF type 5, rare per se and unreported in literature.


Keywords: Neurofibromatosis type 5, Oral Manifestation.

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Introduction

Neurofibromatosis (NF) is a relatively common disorder characterized by neurological and cutaneous lesions. Lesions in all the oral soft tissues have been noted, with a prevalence reported of less than 10%1, 26%2, and up to 72% of patients in a review that included mucosal abnormalities such as enlargement of the fungiform papillae but lacked histopathologic diagnosis3. At least one intraoral or radiographic sign of NF was found in 92% of patients. Von Recklinghausen’s disease (NF-1) and bilateraly acoustic NF (NF-2) are major forms.

Neurofibromatosis type 5 and its manifestations, especially the dental ones, are unreported. Bruxism is a common parafunctional habit, occurring both during sleep and wakefulness. The etiology is multifactorial. There is no known treatment to stop bruxism.

The role of bruxism in the process of tooth wear is unclear, but it is not considered a major cause. Grinding of teeth has long been held as one physical manifestation of stress and anxiety. Systemic conditions affecting the pulp are
seldom documented and appear to be extremely uncommon. On reviewing the literature, no reports of pulpal pathology have been found in NF4.

**Case Report**

We report a case of a 12 years old young male who came to our attention for dental pain. An accurate oral examination revealed an insufficient oral hygiene, few and not deep caries in the primary dentition, affected by evident bruxism visible at first sight.

Ortopantomography (Fig.1), which enhances dental grindings and occlusal erosion of deciduous molars and a Teleradiography (Fig.2) have been carried out. They revealed a delayed development of the root of permanent premolars.

The occlusion revealed a first skeletal class, confirmed by the examination of the cephalometry. In our case, after an instruction of oral hygiene, a follow up of 2 years has been carried out, and the decision to wait for natural exfoliation of the primary molars was taken for the first years, to control the level of bruxism.

**Discussion**

Neurofibromatosis is a diseases which mostly appears in 2 type, but a third, a fourth and a fifth are as well reported (5,6). Neurofibromatosis type 1 accounts for the 90% of the cases and is the classic form. Major features includes six or more cafe-au-lait spots, cutaneous neurofibromas and Lisch nodules. The inheritance is autosomal dominant and it's caused by a mutation in a tumour suppressor gene which maps to chromosome region 17q11.25,6. The hallmark of NF type 2 (NF2) consists of bilateral acoustic neuromas. NF2 is caused by mutation in a tumor suppressor gene which maps to chromosome region 22q12.2. Thorax and lungs are affected in various forms. The patients have dyspnea, cough and chest pain.

Thoracic manifestations of neurofibromatosis are such as upper airway, chestwall, mediastinal, thoracic neoplasms, and
lung parenchyma. Rib abnormalities are common in patients with NF. They have been described as twisted ribbon deformity. Notching of the ribs occurs secondary to erosion by intercostal neurofibromas or a primary defect in bone formation\textsuperscript{7,8}.

In NF3 (Riccardi type) cafe-au-lait-spots are usually pale, few in number and may be large\textsuperscript{4,6,9}.

In NF3 intestinal type, neurofibromatous involvement is limited to the gastrointestinal tract. NF4 is a category of different phenotypes for those patients who do not fit close into any other known type. Segmented Neurofibromatosis type 5 (NF5) is a rare disease \textsuperscript{30} time less common than Neurofibromatosis type 1. Neurofibromas and cafe-au-lait-spots have been frequently reported restricted to one area of the body\textsuperscript{10,11,14}.

Evidence is consistent with mosaicism for NF1 mutations\textsuperscript{11,13}.

Oral manifestation in segmented neurofibromatosis are unreported. Most common oral findings in neurofibromatosis are oral neurofibromas, enlarged fungiform papilla, intrabony lesions, wide inferior alveolar canals and enlarged mandibular foramina. Tumor of the tongue and tongue lesions are reported in literature as well as malposition of the teeth. No enamel or dentin deficiency are reported but increased bone density, enlarged mandibular foramen and increased dimensions of coronoid notch\textsuperscript{5,11}.

Neurofibromatosis type 5 and its manifestations, especially the dental ones, are unreported. On the other hands bruxism at young ages is becoming more and more frequent, and our hypothesis is that, since stress is the first cause of bruxism\textsuperscript{15}. Bruxism is an habit that can have a negative effect on the quality of life of affected individuals by damaging the teeth, periodontium, facial muscles, and the temporomandibular joint\textsuperscript{16-19}.

The long-term follow-up and instrumental and multidisciplinary examination may put under stress very young patients. Considering stress unusual between infancy and adolescence, we should not underestimate the role played by occlusal precontact which is present in our case. It is common for dentists to confect musclerelaxing bite plates for patient to be used while sleeping in order to protect the teeth and periodontium\textsuperscript{20-22}.

Conclusions

Moreover, the use of a bite to reduce the erosion of the surface of the teeth is not appropriate and a strict surveillance of young patients, along with a correct oral hygiene, seems to be the best choice, unless a selective reducing or totally eradicating precontact can rise a good result.

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Declarations of interest

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