Humanized dental care for a patient with neurofibromatosis type 1 in a specialized children’s dental clinic: a case report

Atención odontológica humanizada a un paciente con neurofibromatosis tipo 1 atendido en una clínica de odontología de especialización infantil: reporte de Caso

Atendimento odontológico humanizado a paciente com neurofibromatose tipo 1 atendido em ambulatório especializado de odontologia infantil: relato de caso

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Abstract

This paper aims to report an individualized, inclusive, and humanized dental approach, emphasizing the importance of treating a child aged four years and three months with neurofibromatosis type 1. The patient was referred and treated at the Clinic of the Specialization Course in Pediatric Dentistry of the Graduate School, School of Dentistry, Universidad de la República, Montevideo, Uruguay. The clinical examination revealed scars from the surgical treatment of the plexiform neurofibroma and neck and oral muscle hypotonicity. The intraoral clinical examination showed large edentulous areas as 15 deciduous teeth were missing. The patient had active carious lesions, generalized microbial biofilm on the tooth surfaces, and gingival bleeding. A comprehensive treatment was provided, as well as restorative therapy with upper and lower removable partial dentures. We conclude that the dental educational, preventive, restorative, and rehabilitation treatment was satisfactory. Furthermore, empathy, as a dynamic process, involved cognitive, affective, and behavioral mechanisms. It also helped enhance the patient’s oral health.

Keywords: Recklinghause’s disease of the nervous system, mouth, child, maintenance of space, aesthetics, dental prosthesis.

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Neurocutaneous syndromes comprise a heterogeneous group of idiopathic pathologies characterized by alterations of the central nervous system (CNS), and integuments. One of these syndromes is neurofibromatosis (NF), an autosomal dominant disease with variable expression and a progressive multisystem pathology. \(^{(1,2)}\) This condition includes three autosomal dominant diseases: Neurofibromatosis type 1 (NF1), Neurofibromatosis type 2 (NF2), and Schwannomatosis or Neurofibromatosis type 3. These last two are rare diseases, while Neurofibromatosis type 3 is very rare. \(^{(3)}\) NF2 is known as bilateral acoustic NF and is characterized by multiple tumors and brain and spinal cord lesions. \(^{(4)}\) The first symptom is usually the growth of tumors in the auditory nerves resulting in hearing loss. \(^{(5)}\) NF1 is the most common human dominant genetic disease of the diseases caused by single-gene...
defects. Its prevalence is 1:3000 children born alive, which is more frequent than other diseases such as type 1 diabetes mellitus (1:13000) or cystic fibrosis (1:10,000). The first confirmed case of NF was reported in a 1768 publication, which included the description of a patient with cutaneous neurofibromas, a condition the patient had been inherited from his father. In 1882, the physician Fiedrich Von Recklinhausen described the pathology for the first time, which is why NF1 bears his name. The disease is also known as the "Elephant Man" disease, but this is a historical error since it has been proven that the "Elephant Man" had Proteus syndrome.

The NF1 gene is located on chromosome 17, band q11.2 and encodes a GTPase-activating protein called "neurofibromin," with a high expression level in ectodermal tissue (skin, nerves, and bone tissue). Over 500 mutations of this gene have been described. Although diagnosis is based on clinical criteria, molecular tests for the NF1 gene may be useful in some cases. The disease has complete penetrance, and its expression is variable, so the degree of involvement of each individual is difficult to predict even in the same family and in cases of identical twins.

NF1 diagnosis is clinical, and in 1988 the specific features were defined by the US National Institute of Health as six or more light brown spots equal to or larger than 5 mm in children; two or more neurofibromas of any type, or one plexiform neurofibroma; freckling in the armpits or the groin; optic nerve glioma; lisch nodules (iris hamartomas); bone lesion defined as sphenoid dysplasia; and a first-degree relative with the disease. Finding two or more of these criteria justifies the diagnosis. Although not all patients have the most severe clinical or cosmetic complications of the disease, most NF1 patients and their families feel uncertainty about the course and evolution of the disease. This includes new tumors, aesthetic compromise, and potential transmission to their children, which greatly impacts the patients' quality of life.

In summary, NF1 is a multisystemic pathology that can potentially affect the eyes, the musculoskeletal, cardiovascular, endocrine systems, the CNS, and the peripheral system. There is no cure for the condition. It is treated by managing symptoms and potential complications, especially those associated with tumor growth. It is estimated that these patients’ life expectancy is about eight years less than that of the general population. Between 50% and 80% of patients with the disease may have learning difficulties and behavioral disorders. Thirty percent of children with NF1 may have an ASD, and about 40% may have attention deficit or hyperactivity disorder. Voice and oral motor disorders have also been described in children.

The health personnel should know these oral and maxillofacial manifestations since dentists can be essential to diagnose the disease correctly. Oral manifestations in patients with neurofibromatosis range from 3.8% to 20%. Any part of the oral cavity can be affected. The tongue is the site most frequently affected by neurofibromas, followed by the palate, gums, oral mucosa, floor of the mouth, and labial mucosa more rarely. Although not all patients have the most severe clinical or cosmetic complications of the disease, most NF1 patients and their families feel uncertainty about the course and evolution of the disease. This includes new tumors, aesthetic compromise, and potential transmission to their children, which greatly impacts the patients’ quality of life.

Rezende et al. believe that the patient’s well-being is more important than striking a balance between meeting expectations and pain. That is to say, the patient’s well-being should be the starting point of any treatment plan. This case report describes an individualized, inclusive, and humanized dental approach emphasizing the importance of rehabilitative treatment with preventive maintenance. The patient had NF1 and was treated in 2017 at the Clinic of the Specialization Course in Pediatric Dentistry.
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Clinical case

A male patient aged four years and three months was taken to the Clinic of the Specialization Course in Pediatric Dentistry of the School of Dentistry, University of the Republic by his parents. His usual health center had referred him to assess a potential rehabilitation treatment. The patient’s caretakers signed an informed consent to authorize all the necessary dental procedures, photographs, and the dissemination of the case.

In the medical history, both parents reported that the child was diagnosed with NF1, a disease inherited from his father, who also had the condition. The interview with the parents was not easy because the family’s socioeconomic and education levels are low: they are a vulnerable population, they do not own their house, they are tenants and have electricity and drinking water but no sanitation. Both report dental caries, gum diseases, and missing teeth in their dental histories. The mother has removable dentures.

As for personal medical history, the patient is being treated by physicians to control his condition. Her mother had a normal pregnancy, but the delivery was premature: 34 weeks. The patients’ birth weight was 2 kg 898 g, his height 46 cm, and he was breastfed exclusively until the age of 6 months. Psychomotor maturation was very slow, as he began to sit up at one and walk at two.

Regarding the dental eruption process, his mother reported that a lower incisor erupted at eight months of age. The child was breastfed until the age of two and a half, and he had the deforming habit of lip-sucking.

In terms of oral health habits, the child uses table salt and topical fluoride toothpaste as systemic fluoride. The child brushed his teeth three times a day without adult supervision and followed a cariogenic diet: frequent sugary drinks, foods, and carbohydrates that he consumed between four and up to six times during the day.

The area’s examination showed that the patient presented facial asymmetry, dark circles under his eyes, an everted and large lower lip, reduced muscle tone, and both labial commissures cracked. Upon conducting a functional exam, we detected mixed respiration, altered phonation and mastication due to missing teeth, and atypical swallowing.

Regarding the clinical signs of NF1, the child had scarring from the surgical treatment of his plexiform neurofibroma, affecting the neck and oral muscular hypotonicity. His cognitive and communication skills were intact. The child was quite shy and had great difficulty relating to others but was cooperative in the clinical visits (Figure 1).

Figure 1: Frontal extraoral examination.

Source: Department of Pediatric Dentistry, Universidad de la República.
The clinical examination of the oral cavity (Figures 2, 3, and 4) showed no mucosal lesions, 15 temporary teeth missing (55-54-52-51-61-62-63-63-64-65-73-73-74-75-83-84-85) and 5 temporary teeth with caries lesions (53-71-72-81-81-82). We also detected biofilm on the tooth surfaces, gingival bleeding on probing, abundant saliva, a voluminous, overflowing hypotonic tongue, and adhesions and frenulum without peculiarities.

**Figures 2, 3, 4:** Clinical appearance of the child’s oral cavity at the start of treatment.

Source: Pediatric Dentistry Department, Universidad de la República.

The initial panoramic radiograph (Figure 5) showed many temporary teeth missing, the germ of permanent teeth, and delayed eruption of some teeth, which required follow-up treatment.

**Figure 5:** Initial panoramic radiograph.

Based on the examinations and the diagnosis of the patient’s condition, it was determined that he was a high-risk patient likely to develop new carious lesions. First, an introduction was made to explain prevention and motivational strategies for health education aimed at the child, on an individual basis, and at the entire family. We explained oral hygiene education, the importance of supervising oral brushing in children, provided food advice (a food plan was requested), and explained the use and dosage of fluoride toothpaste.

A dental visit was scheduled to perform the necessary outpatient dental procedures. The teeth with active carious lesions (53-palatal face, 71, 72, 81, and 82-mesial and distal faces) were rehabilitated by removing caries selectively, and the upper and lower removable dentures were made. In each patient visit, biofilm was controlled, and professional brushing was performed.
with 1450 ppm fluoride toothpaste, emphasizing the importance of brushing in the evening and after main meals to achieve a healthy behavior change. The child and his family were instructed to brush their teeth at least twice a day with 1100 ppm or over fluoride toothpaste. The patient received a hygiene kit with a toothbrush and 1450 ppm fluoride toothpaste. with 1450 ppm fluoride toothpaste, emphasizing the importance of brushing in the evening and after main meals to achieve a healthy behavior change. The child and his family were instructed to brush their teeth at least twice a day with 1100 ppm or over fluoride toothpaste. The patient received a hygiene kit with 1450 ppm fluoride toothpaste.

The dental restorations were made with light-curing glass-ionomer cement (FUJI IX GC Corporation) in teeth 53, 71, 72, 81, and 82 (Figure 6). The upper and lower removable dentures for space maintenance were made following the necessary steps: a) taking one-step impressions with heavy and light silicone and casting to obtain plaster models, b) registration plate and impellers, c) occlusal registration and restoration of the vertical dimension, d) dental alignment, e) tooth try-in and f) upper and lower removable dentures (Figures 7, 8, 9, 10, 11, 12, and 13).

**Figure 6:** Restoration with light-curing glass-ionomer cement 71, 72, 81, 82.

![Figure 6: Restoration with light-curing glass-ionomer cement 71, 72, 81, 82.](image)

**Source:** Pediatric Dentistry Department. Universidad de la República.

**Figures 7 and 8:** Taking silicone impressions and making models and plates of registration and impellers.

![Figures 7 and 8: Taking silicone impressions and making models and plates of registration and impellers.](image)

**Source:** Pediatric Dentistry Department. Universidad de la República.
**Figures 9 and 10:** Vertical dimension registration and alignment test.

**Source:** Pediatric Dentistry Department. Universidad de la República.

**Figures 11, 12, 13:** Clinical aspects of the oral cavity with the upper and lower removable dentures in the mouth. Installation.

**Source:** Pediatric Dentistry Department. Universidad de la República.

**Figure 14:** The child’s smile when he returned three months later for evaluation and dental examination.

**Source:** Pediatric Dentistry Department. Universidad de la República.
The patient returned for preventive dental check-ups periodically and three months after denture installation to control the infection and the space maintainers (Figure 14).

The boy, who had no intellectual disabilities, showed a positive attitude in all the appointments. The professionals responsible for his dental care used the tell-show-do technique and positive reinforcement. There was no crying or need for physical restraint.

**Discussion**

Neurofibromatosis types are a group of genetic disorders that cause nerve tissue tumors. These tumors can develop anywhere in the nervous system: the brain, the spinal cord, or the nerves.\(^1,2\)

Figueiredo et al.\(^14\) wrote a clinical case of a girl who had NF1, more specifically plexiform neurofibroma (involvement of several nerve groups). The condition affected the upper eyelid of her right eye and caused ptosis, loss of vision, and facial asymmetry. In addition, she had a younger sister with the same condition. In the case of our patient, the child’s father also had clinical signs of NF1.

Working with the patient’s diet and oral hygiene was challenging. Something noteworthy is the patient’s vulnerability due to his low self-esteem. Brêtas, 2010,\(^15\) states that low self-esteem may be due to the individual’s lack of motivation regarding his personal care and thus not finding any reason to take care of himself, even if he is a 4-year-old boy. Improving motivation in the child and his relatives is essential in the care strategy to help identify their basic dental needs and alleviate their suffering.

The treatment proposed included prosthetic rehabilitation to restore function and aesthetics, improve communication and social integration with other people, as this can also be associated with improved self-esteem, confidence, and quality of life. According to Antoniazzi et al., 2018,\(^16\) quality of life is related to oral health and includes several dimensions, as it involves health status and social and behavior factors.

In addition, removable partial dentures are essential in these pediatric patients to restore their altered functions and optimize the craniofacial skeletal growth and development of the dentition. This improves mastication, swallowing, phonation, and digestion. However, Prabhakar et al.\(^17\) and Brelaz\(^18\) reported the disadvantages of using removable partial dentures and, in some cases, the fact that it may be difficult to adjust them in young patients. This was not the case in this clinical case, as the patient was very cooperative.

As stated by Figueiredo et al.\(^19\) and Gonçalves et al.,\(^20\) it is necessary to create a bridge between dentistry and the medical specialties involved in the care of these patients to enable treatment progress. The smile is a cause for aesthetic concern and also a way to recover one’s self-esteem.\(^19,20\) In this case, the patient’s smile was restored by making removable partial dentures.

Finally, the dentist providing care to patients with severe NF1 must empathize with them, engage with their clinical history to assess their cognitive profile, and perform procedures according to their ability.\(^14\) According to Brazilian Law no. 13.146\(^21\) on the inclusion of people with disabilities, health professionals must provide humanized dental care for people with any syndrome and base their work on psychosocial and family interaction.

They must also demonstrate proficiency in the clinical dental procedures, patience, and the enjoyment of caring for a patient with special needs, as these are essential attributes for the care of children in the clinic and to have a successful treatment, as presented in this work.

**Clinical relevance and application**

This clinical case presents the characteristics of a patient with NF1, in a clear, descriptive, and illustrative manner, without limitations, thus facilitating diagnosis. In addition, it shows adequate preventive, restorative, and rehabilitation treatment, highlighting the importance of the
professional's bond with and commitment to the patient and his family while providing care.

Conclusions

The main oral and general clinical characteristics of the patient reported agreed with what is described in the literature for NF1: poor muscle tone, poor oral hygiene, and active dental carious lesions. This showed the need for dental educational, preventive, restorative, and rehabilitation treatment. Dentists play an essential role in the rehabilitation and well-being of these patients and should be included in a multidisciplinary care team.

It is essential to know the comorbidities of patients with NF1 to manage their dental treatment.

References


2. Gonzalo M. Neurofibromatosis tipo 1: análisis de características clínicas y tipo de complicaciones en pacientes asistidos en un Hospital de referencia nacional [Monografía de posgrado en Pediatría]. [Montevideo]: Universidad de la Republica, UdelaR; 2015. 20p.


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