Case Report Article

Prader-Willi Syndrome: clinical case report

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Received for publication: December 1, 2013. Accepted for publication: December 13, 2013.

Abstract

Introduction: Prader-Willi syndrome (PWS) is a neurobehavioral genetic disease whose cause is failure on chromosome 15. It is considered the primary genetic cause of obesity, since it is characterized by hyperphagia. Although the scientific literature will produce articles on Prader-Willi syndrome, few reported oral conditions of these patients. Objective: This study aimed to describe the oral health status of a patient diagnosed with PWS. Case report: A boy aged 10 years-old, leucoderma, attended the Discipline of Dentistry for Special Care Patients, Pontifical Catholic University of Paraná (PUCPR), with all the inherent PWS characteristics such as hyperphagia and obesity. Clinical, radiographic and laboratory tests were performed to verify the oral health conditions which showed the presence of biofilm accumulation, gingivitis and a high DMFT index. It was necessary to adequate oral environment through extractions, restorations, and prophylactic therapy. Conclusion: Considering the information obtained, it was concluded that the motivation to maintain oral health should be constant for this patient and involve family, since hyperphagia, which is a determinant for obesity, decisively contributes to the evolution of oral diseases.
Introduction

Prader-Willi syndrome (PWS), described by Prader, Labhart and Willi in 1956, is considered a neurobehavioral disease currently indicated as one of the most frequent cause of chromosome microdeletions [9]. PWS is the most common phenotype of genetic obesity. Additionally, Prader-Willi and Angelman syndromes are the main recognized human diseases determined by mechanisms of genomic imprinting, that is, a genetic phenomenon in which certain genes are expressed by only one allele [7]. In this case, deletion of the proximal portion of the long arm of the paternal chromosome 15 (15 q11-13) or, more rarely translocation of chromosome 15 occurs [8]. The prevalence is of approximately 1 for each 15,000 births on both genders [6].

PWS is characterized by two clinical phases. At first phase, the symptoms are neonatal hypotonia, difficulties to be fed, lethargy, weak cry and hyporeflexia. At second phase, starting from six months of age, a gradual improvement of hypotonia, hyperphagia, weight gain and obesity occur [7, 15]. Obesity is the major cause of the morbidity and mortality increasing among patients exhibiting the syndrome [3]. Clinical features comprised also respiratory problems during sleep, mild to severe mental retardation learning disability and short stature [7, 13, 15].

The syndrome diagnosis is clinical based on physical and behavioral data which can be confirmed by the analysis of chromosome 15 segment (q11-q13) through methylation or in situ hybridization [10].

With regard to oral health of PWS individuals, few studies have been conducted. Some authors reported an increasing in carious teeth and salivary flow reduction [4, 13], in addition to enamel hypoplasia [5, 6]. Although scientific literature is reporting on PWS, little studies focus on oral health conditions of these individuals. Therefore, the aim of this study was to describe the oral condition of a PWS individual treated at the Pontifical Catholic University of Paraná (PUCPR).

Clinical case report

This study was conducted on a leucoderma male patient born by normal delivery. Pre- and post-natal tests were normal indicating a healthy fetus/newborn. Until 6 months of age, the infant was formula-fed exclusively, period after which he started to be fed by solid food. The patient was diagnosed with PWS at three years-old by the medical team of the Clinical Hospital of the Federal University of Parana (UFPR), due to the presence of muscle hypotonia, delay in common psychomotor development phases, difficulty in sucking/swallowing, weak cry, and somnolence.

At 10 years-old, the patient started dental treatment at the Discipline for Special Care Patients (PUCPR). At anamnesis, marked physical alterations of the syndrome were noted, such as: myopia, strabismus, obesity, short stature (figure 1), maxillary lip and labial commissures facing down, small hands, and hypopigmentation of the hair, skin and eyes. The patient’s mother reported during anamnesis that her pregnancy was uneventful and no other cases have been seen in the family. The mother still informed that her son had episodes of nervousness and anxiety due to dietary restrictions. The patient has been treated by a multidisciplinary team composed by nutritionist, psychologist, and endocrinologist. During his free time, the patient liked to play videogame, listen to music and sing.

During current medical history, patient exhibited cognitive deficiency characterized by learning difficulties, mainly related to writing. During past dental history, the mother reported that his first dental appointment was at seven years-old, at school, and the behavior was satisfactory.

Intraoral clinical examination revealed significant biofilm accumulation, gingivitis, dry mouth and angular chelitis. Occlusion examination evidenced ogival palate, anterior crowding, and Class II malocclusion. Panoramic and periapical radiographic examination indicated delay in tooth eruption and tooth crowding at maxillary and mandibular anterior teeth (figure 2).

Figure 1 – Patient’s physical characteristics. At 10 years-old, the boy showed obesity, short stature, small hands and feet and hair/skin hypopigmentation.
Figure 2 - Panoramic radiographic examination showing maxillary and mandibular anterior tooth crowding and giroversion of some teeth; presence of residual roots of teeth #53, #83, and #85; and loss of tooth #63. Radiographic examination also revealed caries in teeth #16, #26, #46, #73, #74 and #75.

Discussion

In this present study, a boy aged 10 years-old was diagnosed with PWS and treated at PUCPR’s dental clinic. The clinical characteristics of this syndrome reported in the literature, such as: neonatal hypotonia, difficulties in feeding, lethargy weak cry, hyporeflexia, hyperphagia, and obesity [7, 11, 15], were confirmed by the patient’s mother during anamnesis. Short stature, small hands and feet, skin/hair/retina hypopigmentation, thin lips, labial commissures facing down, myopia, and strabismus also described in prior studies were also observed [4, 13, 14].

Randomized clinical trials have demonstrated the favorable effect of growth hormone (GH) reposition to reduce fat mass and increase lean body mass, complementing nutritional guidance. Not with standing, GH seems not to have significant effect on controlling hyperphagia in these patients [13].

Corroborating Cortés et al. [7], Olczak-Kowalczyk et al. [13], and Vargas et al. [15], the patient of this case report showed learning disorder related to writing; but with mild cognitive deficiency. PWS patients also presented bone mineralization decrease, fact explaining the high osteoporosis incidence associated with the syndrome [5].

According to the patient’s mother, episodes of irritability were rare, mainly related to dietary restriction. Prior studies also reported this situation [1, 11]. Generally, this study’s patient had a sociable and friendly behavior with all multidisciplinary team.

The peculiar facial characteristics of PWS patients were also observed in this case report: almond eyes, thin maxillary lips, labial commissures facing down, and dysmorphic face. Among the oral manifestations reported by Carvalho et al. [5, 6], this case report found the presence of caries, enamel hypoplasia, and malocclusion. On the other hand, ogival palate, delay in tooth eruption, supernumerary teeth, microdontia, micrognathia, taurodontism and candidiasis were not seen.

Hyperphagia, the most important PWS feature [7], can explain the high rate of carious teeth observed in this patient who exhibited a DMTF index greater than that of children at same age. Six teeth had caries, three were residual roots and one tooth was lost. The treatment executed comprised the extractions of teeth #53, #83 and #85; resin composite restorations of teeth #16, #26, #46, and #74; and light-cured glass ionomer cement restoration of teeth #73 and #75. Next, preventive measures were instituted with sealants on many teeth and fluoride varnish application.

Other important factor contributing to oral health deterioration is salivary flow decrease [4, 13], a feature clinically observed through dry mucosae. To evaluate the amount of produced saliva, we used the method proposed by Navazesh et al. [12] and Banderas-Tarabay [2]. The produced volume was very low, characterizing hyposalivation.

The next step was to instruct the family to adopt measures promoting oral health maintenance. The relatives were instructed regarding to proper oral hygiene, to make them also responsible by the treatment. The patient still continued being followed-up at the Discipline of Dentistry for Special Care Patients of the institution to preserve oral health.

Conclusion

Based on the information obtained after clinical and radiographic examination, it can be concluded that both PWS patient and the family should be constantly motivated to maintain oral health, because hyperphagia which is determinant for obesity, decisively contributed to increase DMTF index. Additionally, hyposalivation can contribute to develop bacterial biofilm and gingivitis.
References


