RESEARCH GROUP IN ORAL PATHOLOGY

SYNDROMIC FORM OF OROFACIAL CLEFTING:
VAN DER WOUDE SYNDROME

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Abstract  
The Van der Woude Syndrome (VWS) is a rare congenital malformation, clinically characterized by the presence of lower lip pits, cleft lip and/or palate and occasionally hypodontia. The present report presents a clinical case of VWS, and emphasizes the role of the dentist as the first person to diagnose this syndrome.

Keywords: Van der Woude Syndrome; Cleft lip and/ or palate; Lower lip pits.

INTRODUCTION

The Van der Woude (VWS) Syndrome is a congenital malformation, genetically inherited by an autosomal dominant trait, of incomplete penetrance and clinically characterized by the presence of lower lip pits, cleft lip and/or palate, and occasionally hypodontia. In spite of Van der Woude having described this pathology in 1954, and hav-
ing studied it intensely, the first reports were made in 1845 by Demarquaz and in 1860 by Murray.\(^1\)

As it concerns a rather uncommon anomaly that affects approximately 2 in every 40,000 to 100,000 live births,\(^2\) the aim of this report is to present a clinical case of VWS with emphasis on its main clinical characteristics.

**CLINICAL CASE**

The patient, a 7-year-old girl with bilateral bottom lip pits, bilateral cleft lip and palate – operated, and clinical absence of teeth 11 and 21. The panoramic radiograph showed supernumerary teeth in the region of the left and right maxillary lateral incisors, denominated precanines, one of them impacted and agenesis of teeth 12 and 22. (Figure 1) The patient had been submitted to two cheioplasty procedures and surgical closure of the cleft lip and palate. The first surgery was performed ten months after birth and the second at five years of age. The family history showed mother and sister with bilateral cleft lip, with absence of lower lip pits. The presence of other cases of cleft lip and/or palate in the family was related, among them the father and paternal grandfather. The patient affirmed having good performance at school and no otalgia, otorrhea was observed. The patient denied having difficulties with chewing and swallowing. According to the clinical findings associated with the family, the diagnosis of van der Woude Syndrome was established. At present dental and medical follow-up of the patient is being performed.

Figure 1 – Van der Woude syndrome

A. Bilateral cleft. B. Symmetric lower lip pits. C. Panoramic radiograph: precanine supernumerary teeth (arrows).
DISCUSSION

The van der Woude Syndrome has a highly variable expression, with different phenotypes that show, for example, a hardly evident depression through to congenital pits in the bottom lip. The presence of orofacial clefts vary from bifid uvula to complete clefts.\(^{(5)}\)

In this disease, other less common anomalies may be observed, such as maxillary and mandibular adherence, absence of the maxillary lateral incisors, ankyloglossia and bifid uvula.\(^{(6)}\) The dental anomalies are considered a cardinal sign of VWS.\(^{(5)}\)

The patient in this study, in addition to the classical clinical presentation of VWS, also has dental anomalies such as giroversion, “precanine” supernumerary teeth and hypodontia of the maxillary lateral incisors. In the literature, these characteristics in the same patient are not common. In the cases described, when there is presence of supernumerary teeth, hypodontias are not shown.

The family occurrence of VWS ranges from 50% to 60%, and its variability of expression for members of the family that have the same mutation suggests multifactorial influences, similar to those of non-syndromic clefts. The remainder of the cases occur with new forms of mutation.\(^{(5,6)}\) According to Burdick et al (1985),\(^{(7)}\) 15% of those with the syndrome do not present lip pits, and these individual are clinically indistinguishable from those who present isolated non syndromic clefts. In the case under study, various members of the patient’s family with VWS have cleft lips, but none of them presented lower lip pits. One understands that these individuals may have the van der Woude Syndrome, in spite of the clinical characteristics not being sufficient to define the diagnosis. It is known that over 300 syndromes exhibit cleft lip and palate as one of the characteristics.\(^{(8)}\)

In the literature, the van der Woude Syndrome is considered a Mendelian disorder. Its genetic origin is attributed to microdeletions in chromosomal bands 1q32-q41, which belong to gene IRF6. Sertie et al.\(^{(9)}\) have suggested the association of another gene, 17p11.2-11.1, especially in those patients who have the syndrome with a cleft palate, and have affirmed that the Mendelian concept of monogenic disease for VWS appears to be increasingly unsustainable.

The popliteal pterygium syndrome (PPS) is frequently referred to in the differential diagnosis of VWS. It is characterized by the same clinical signs as those of VWS, in addition to abnormalities of the lower limbs and nails, adhesion of the popliteal membrane, ankyloglossia and genital anomalies, including cryptorchidism, bifid scrotum and uterine hypoplasia.\(^{(10)}\) It is associated with mutations in gene IRF6, also attributed to VWS. In the patient under study, who has VWS, the clinical alterations described were not observed.
The dentist should recognize and diagnose this syndrome, especially at the beginning of life, and establish adequate treatment. The participation of various medical and dental specialties will provide individuals with VWS good esthetic-functional conditions. The patient in this study was submitted to various corrective surgical procedures and is being followed up by a phonoaudiologist. In addition, she is undergoing orthodontic, clinical dental and surgical treatment for the correction of malocclusion, supernumerary teeth, and replacement of absent teeth.

REFERENCES


