Parry-Romberg syndrome in a pediatric patient.
A case report.

Abstract: The Parry-Romberg syndrome is a rare degenerative disease of unknown etiology that has dental implications. It is characterized by a progressive hemifacial atrophy that appears in the early stages of life. It causes aesthetic, functional and psychological alterations, and has social implications for the patient. There is no definitive treatment for the Parry-Romberg syndrome. Systemic and immune alterations produce oral and maxillofacial manifestations, which need to be managed by specialized dental professionals. The aim of this paper is to do a literature review of the Parry-Romberg syndrome and describe the oral and clinical characteristics of this condition in a 12-year-old male pediatric patient, who had a history of neurological disorders and facial asymmetry on the left side. Dentists require an adequate knowledge of the clinical and dental characteristics of this syndrome. With early diagnosis and appropriate surgical and orthodontic treatment at an early age, they can improve the quality of life of patients and minimize invasive long-term effects.

Keywords: Parry-Romberg syndrome, Facial asymmetry, Progressive facial hemiatrophy, Oral manifestations.

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INTRODUCTION.

Described in 1825 by Caleb Parry and in 1846 by Moritz Romberg, the Parry-Romberg syndrome or Progressive Hemifacial Atrophy, as Eulemberg coined the term, is a rare degenerative condition characterized by a generally slow and progressive unilateral atrophy of facial tissues, affecting the subcutaneous fatty tissue, connective tissue, skin, muscle and bone.

It has a higher prevalence in women (3:1) and occurs during the first two decades of life. The syndrome progresses for several years and then it reaches a period of stability. Its prevalence is estimated at 1/700,000 individuals. Its etiology is unknown and several hypotheses have been proposed: a manifestation of localized scleroderma; an imbalance in fat metabolism; trauma; viral infections; endocrine disturbances, autoimmunity, and genetic factors.

Clinically, it is characterized by a marked unilateral facial asymmetry; however, 5% and 10% of cases being bilateral have also been reported. The asymmetry is preceded by a change in epidermal hue, by hyperpigmentation or depigmentation. The affected area usually becomes tense and compact, extending to the neck and to the side of the body, accompanied by other manifestations such as: alopecia, loss of eyelashes and eyebrows.

There are other symptoms such as trigeminal neuralgia, facial paresthesia, headaches, contralateral epilepsy and enophthalmos (neurological and ophthalmological manifestations), which can occur frequently in patients with this syndrome. The intraoral soft tissues and muscles of mastication may not be affected, so their basic functions are not affected either. However, dental exposure caused by problems on the lips, deviation of dental midline and lingual atrophy may
occur. Facial muscles are thinned, but keep their function. Manifestations in bones or cartilages can be observed, especially when the syndrome occurs at an early age.

Between 36.6% and 53.6% of patients with Parry-Romberg syndrome have linear scleroderma or “coup de sabre” located in the frontal region, from the eyebrows to the scalp. They are considered a clinical-pathological variant of the same condition. Besides the fact that sometimes the two diagnoses coexist, they also coincide with the age of onset and the course of the syndrome. Histologically a fatty tissue atrophy and fibrosis of the skin are observed.

In the case of scleroderma it has been reported a massive lymphocytic inflammatory infiltration around surface vessels and deep plexuses of the skin, with a hardening of the skin caused by excessive collagen formation.

Pathophysiologically, some authors have proposed that the syndrome may be caused by an inflammatory meningoencephalic condition associated with a vasculitis; others, a chronic disorder related to vasomotor hyperactivity of the sympathetic nervous system.

Diagnosis can be made clinically if the patient has facial asymmetry, based on skin characteristics and soft tissue findings. If there are neurological symptoms a MRI and a CT scan should be performed as diagnostic aids. The Parry Romberg syndrome and scleroderma require a similar treatment, based on antibiotics and antimalarials (chloroquine and hydroxychloroquine); for severe cases, cyclophosphamide, cyclosporine and corticosteroids are suggested.

The aesthetic treatment is not recommended until the syndrome is stable. Surgical-aesthetic procedures include silicone implants, fat, skin and bone/cartilage grafting.

Regarding the dental aspects, the teeth on the affected side are smaller and have shorter roots.

The aim of this paper is to describe the clinical and dental characteristics of the Parry-Romberg syndrome in a male pediatric patient.

**CASE REPORT**

12-year-old male patient treated at the pediatric unit at Hospital General “Luz Gonzalez Cosío” in Zacatecas, Mexico, showing changes of symmetry in his skull and face with six months of evolution. The mother says his son suffers from oppressive headache.

Physical examination showed asymmetry on the left side with an increase in the depth of the nasolabial folds, appa-

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**Figure 1.** Hemifacial atrophy on the left side with apparent sinking of structures on the affected hemiface.

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**Figure 2.** Patient does not show hyperpigmentation or depigmentation in the neck.
rent sinking of the structures on the left side of the face, thinning of the skin from the temporary area to the lower border of the mandible, without altering the function of the muscles in the affected area. (Figure 1) A depressed groove at the middle frontoparietal area extending toward the scalp, causing hair loss is observed.

The patient showed symmetrical tendon reflexes. Cogni-
tive functions and language are normal. Enophtalmos with prominent brows and deviation of the outer corner of the eye and of the lip were observed.

Hyperpigmentation or depigmentation in the neck as described by various authors was not observed. (Figure 2). The patient was referred to the dental and maxillofacial surgery unit to undergo clinical examination and imaging studies of the facial bones with 3D reconstruction.

Intraoral examination revealed pigmentation of left attached gingiva, maxillary and mandibular depression on the left side, with loss of continuity of the ramus.

Anterior walls showed no evidence of alterations, atro-
phy or deviations of the soft tissues; mixed dentition with 24 teeth, dental midline deviated 4mm to the left, uni-
lateral class II canine and molar relationship, horizontal overbite of 6.5mm and vertical overbite greater than 50%. The curve of Spee is mild on the right side and severe on the left, which causes the left temporomandibular joint to dislocate when opening the mouth.

The orthopantomography shows short tooth roots on the affected side (Figure 3).

The scan of the skull and facial bones with 3D re-
construction confirms the deformation of the left bone, affecting bone and showing a decrease in the posterior re-
gion of the eye; a reduction of the size of the facial bones and skull on the left side, affecting frontal and temporal bones on the same side, cheek bone and jaw.(Figure 4) Midline of frontal bone shows a highly marked "coupe de sabre".

Deviation of the bones of the nose to the left side, mar-
ked depression at the level of the canine eminence and malocclusion of teeth 21 and 22 were also observed.

The maxillofacial surgery unit refers the patient to specialists in pediatric dentistry to treat the condition coordinately together.

The main treatment will be in charge of the maxillofacial surgery team at a later stage. Gonadotropin tests are not re-
quested because the patient is in pubertal development.

**DISCUSSION.**

The Parry-Romberg syndrome is a rare disorder, with a higher prevalence in females, and a long evolution during
the first 20 years of life\(^{14}\). The patient presented in this study had a unilateral disorder involving the abovementioned tissues, bone and intraoral soft tissues, especially in the tongue and lips.

There are cases of late-onset of the disorder, some with significant differences in the severity of the symptoms, with a serious affectation of facial tissues. It has been suggested that the degree of deformity may depend on the duration of the syndrome\(^ {15}\). In this particular case and up to the date of this report, the alteration on the left side of the patient’s face has been mild, with bone and intraoral tissue atrophy, and dental malocclusion.

Neurologic abnormalities associated with Parry-Romberg syndrome have been well documented\(^ {16}\). Despite the fact that the patient’s exams did not show any kind of brain injuries, he did suffer from oppressive headache, and is currently under treatment for that condition.

The patient also suffers from enophthalmos on the left side of his face. Enophthalmos are caused by the loss of peripheral adipose tissue of the eyeball\(^ {17}\).

In order to restore function and avoid potential damage to the temporomandibular joint, malocclusion (crowding, pronounced curves of Spee, and deviation of the dental midline) caused by this pathology should be corrected. The patient received basic dental treatment before undergoing treatment to correct malocclusion.

The reduction in soft tissues can be observed on the CT. It helps to distinguish between Parry-Romberg syndrome and other disorders such as atrophy of the salivary glands or facial and chewing muscles paralysis, which could also result in facial asymmetries. Changes in bone tissue on the left side of the patient’s face were observed\(^ {18}\).

The replacement of lost tissue can be performed surgically. Grafts are preferred instead of implants, because they have a lower rate of recession of tissues, and produce a lower local and systemic inflammatory response\(^ {19}\).

In this case it was decided to refer the patient to the pediatric dental service to receive basic dental treatment because he was in pubertal development\(^ {20}\). Subsequently he will be referred to the orthodontic and maxillofacial unit to continue with his treatment.

**CONCLUSION.**

The Parry-Romberg syndrome when detected at an early age should be treated by a multidisciplinary pediatric team, supported by methods of differential diagnosis and aimed at minimizing the effects produced in the skull and dentofacial area. In this case, the patient was referred to a pediatric dental service to receive basic treatment. The use of 3D imaging allows specialists to fully identify the characteristics of the disease and provide early intervention, reducing invasive long-term effects.
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