Fibrodysplasia Ossificans Progressiva: A Review with Presentation of a Case with Temporomandibular Extra-Articular Ankylosis

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Abstract: Fibrodysplasia Ossificans Progressiva (FOP) is a connective tissue disorder that progressively affects tendons, ligaments, aponeurosis, fasciae and muscles which undergo fibrous cell proliferation that progresses to mature bone. It has a prevalence of 1 case per 2 million habitants, having reported less than 1000 cases worldwide. In the maxillofacial region, it might originate extra-articular temporomandibular ankylosis by ossification of ligaments, muscles of the mastication, head and a neck muscles; the most commonly affected are the masseters and sternocleidomastoids. The purpose of this article is to review the Fibrodysplasia Ossificans Progressiva (FOP) and to present the case of a 12-year-old male patient with FOP that causes extra-articular temporomandibular ankylosis. There is no effective proven treatment or prevention and the life expectancy of these patients approaches the 40 years of age, so the management of patients with FOP must be performed with a multidisciplinary approach in which the various health professionals work in a coordinated and joint way to offer a better quality of life to these patients and thus better understand the progression of the disease.

Keywords: Fibrodysplasia Ossificans Progressiva, Myositis Ossificans Systemic, Munchmeyer's Disease, Extra-Articular Temporomandibular Ankylosis, Ankylosis of the Temporomandibular Joint

1. Introduction

   The Fibrodysplasia Ossificans Progressiva (FOP), also known as Myositis Ossificans Progressive, Myositis Ossificans Systemic, Fibrodysplasia Ossificans Progressiva Idiopathic, Munchmeyer’s Disease and Stone Man’s Disease. [1, 2] Initially described by a French doctor, Guy Patin in 1648, who reported a patient “who became wood”. [7] In 1736 the English surgeon John Freke, describes the case of a child of 14 years old with “ramifications of coral do in his back”. [3] The term FOP was introduced by Bauer and Bode in 1940. [8] The dominant autosomal inheritance was first described by Simpson, but it was Eileen M. Shore and Frederick S. Kaplan in 2006 that discovered the gene and the mutation that cause it. [6, 9, 10]

   It is a genetic disease characterized by a connective tissue disorder that progressively affects tendons, ligaments, aponeurosis, fasciae and muscles, which undergo a fibrous cell proliferation that progresses to mature bone, is the most catastrophic heterotopic ossification disorder that occurs in humans. [2 - 6] Ossification follows a cephalo-caudal, proximal-distal, dorsal-ventral and axio-appendicular direction forming "a second skeleton", which progressively diminishes the patient's abilities, making difficult their basic
activities such as walking, hearing, chewing, swallowing, speech and breathing which will result in an early death. [2, 10, 14, 25, 26] Characteristically it respects smooth and cardiac musculature, oculomotor muscles, facial expression muscles and tongue. Its rhythm of progression is variable between the second and third decade of life, the onset is reported between 5 and 25 years old. It has a prevalence of 1 case for every 2,000,000 inhabitants, it is estimated that there are around 3,500 people with FOP worldwide, but only less than 1,000 cases have been reported, it has no predilection between sex, race or geographic location. [6, 10] There are less than 10 multi-generational families worldwide [3, 4] and cases have been reported with exclusive compromise of the maxillofacial region. [22]

It has an autosomal dominant pattern in 10% of the cases, but most patients (90%) correspond to a spontaneous "de novo" mutation in the gene that codes for the Activin A type I receptor like Activin- Kinase-2 (ACVR1 / ALK2) located on chromosome 2 (2q23-24). The ACVR1 / ALK2 receptor belongs to the family of bone morphogenetic protein (BMP) receptors, members of the superfamily of transforming growth factors beta (TGF-β). [2, 4, 10, 11, 12] The predisposition to develop heterotopic bone is due to the ACVR1 mutation (c.617G> A; R206H) which consists of a substitution of a "G" base (guanine) for a " A " (adenine) at position 617, which in turn causes the replacement of the nucleotide Arginine by Histidine at codon 206. This mutation causes the ACVR1 / ALK2 gene to be permanently activated, which causes an increase in the signaling in the cascade of the BMP especially at the level of BMP 1 and 4. Ossification will be favored by any type of trauma which will originate a microenvironment of hypoxia associated with the activation of proinflammatory cytokines such as prostaglandin E-2 (PG-E2) and tumor necrosis factor alpha. (TNF-α).

This combination of factors would induce an "Endothelial-Mesenchymal Transition" (EMT) of the vascular endothelial precursor cells, giving rise to pluripotent mesenchymal cells with the capacity to differentiate into osteoblasts and chondrocytes, these pluripotent cells will be focused on the anatomical regions where the ACVR1 / ALK2 receptor is expressed. (skeletal muscle, blood vessels and cartilage) [2, 11 - 14] Genetic factors condition alterations caused during fetal development, while environmental agents associated with lifestyle (injections, vaccines, anesthetic blockages, muscle fatigue, contusions, viral diseases such as flu, etc.) determine the intensity and rate of advancement of ossification. [2, 10, 15]

The disease is not related to specific alterations in laboratory tests, so its diagnosis is clinical and is based mainly on 2 criteria: congenital malformation of the first toes and progressive heterotopic endochondral ossification. [9, 10, 23] Within the congenital malformations, we can find shortening or malformation of phalanges, metatarsals and metacarpals, as well as synostosis and chondroactyly. Heterotopic ossification in early stages of the disease manifests with some degree of soft tissue inflammation of head, neck and upper back as well as the presence of painful subcutaneous nodules which progress to heterotopic bone plaques that can trigger calcifications at the level of the temporomandibular joint (TMJ), vertebral bodies (especially at the level of C2-C7), orthotopic ankylosis of the costovertebral joints, ossification of the intercostal and paravertebral muscles, progressive spinal deformity such as kyphoscoliosis or thoracic lordosis, evident osteochondromas especially in the medial region of the tibia and a femur with a short and wide neck. [2, 3, 7, 10, 23 - 25] A small number of patients present atypical clinical features such as osteochondromatosis of the hip, degenerative joint disease of the hip, scanty scalp, mild cognitive impairment, growth retardation, cataracts, retinal detachment, glaucoma, craniohypophyrgioma, diffuse cerebral dysfunction with seizures, cerebral and cerebellar malformations, polyostotic fibrous dysplasia, primary amenorrhea, aplastic anemia and hypospadias. [12]

The best imaging technique to establish the diagnosis is simple radiograph, which can reveal presence or absence of ossification, in most areas the lesions will appear as light radiopaque collections within a muscle (at the beginning of the disease) or as irregular masses (at the end of the disease). [2] Computed tomography (CT) is adequate to evaluate the location and degree of participation in certain muscle groups. [27, 28] Ossification can be identified through a bone scintigraphy since the uptake of the isotopes (99mTc MDP [Technetium methylene diphosphate] and Thallium 201) is markedly increased in these cases and have been used as parameters of lesional activity. [9, 13]

The biopsy should be avoided as it leads to an accelerated progression of the lesion. [29] Histologically, early FOP lesions contain intense mononuclear and perivascular infiltration of macrophages, mast cells and lymphocytes. The subsequent migration of mononuclear inflammatory cells to the affected muscle precedes the generalized death of skeletal muscle. After a rapid and destructive inflammatory stage, there is an intense phase associated with angiogenesis and abundant neovascularity. Early fibroproliferative lesions are histologically indistinguishable from aggressive juvenile fibromatosi. As the lesions mature, the tissue undergoes an avascular condensation in the cartilage followed by a revascularization stage with osteogenesis in a process characteristic of heterotopic endochondral ossification. The new heterotopic bone appears histologically normal with mature lamellar bone and often contains marrow elements. Mast cells have been identified in all histological phases of the formation of FOP lesions and are found in a much higher abundance compared to normal skeletal muscle. All stages of histological development are present in an active FOP lesion, which suggests that different regions within the lesion mature at different rates. [3, 10, 27, 30]

The lack of diagnosis or a misdiagnosis is reported in 90% of cases worldwide. In the early stage, it should be differentiated from aggressive juvenile fibromatosi, lymphedema and soft tissue sarcoma. The limitation of movement and the presence of ossifications / calcifications at the radiographic level may indicate myositis ossificans traumatic, myositis ossificans neurogenic, osteoblastomas,
osteosarcomas, chondrosarcomas, as well as mature venous hemangiomias with multiple phleboliths. FOP should be differentiated from other genetic diseases such as progressive bone heteroplasia, Albright's hereditary osteodystrophy, osteoma cutis, ankylosing spondylitis, Still's disease, rigid spine syndrome and Klippel Feil syndrome. [3, 5, 9, 10, 12, 14, 16 - 21]

Some authors recommend surgery procedure in late stages of the disease when ossification results in disability. [32] The current trend is to avoid surgical excision of the ossified parts to increase the movement of the affected region, since surgical excision areas have been observed to re-ossify and further limit mobility with a more aggressive progression. [2, 3, 8, 14, 25, 31]

Attempts to slow the progression of the disease with drugs such as isotretinoin (13-cis-retnoic acid), corticosteroids, indomethacin and disodium etidronate have been ineffective. [21, 33] It is believed that radiotherapy at low doses may be useful for its anti-inflammatory action. [34] but the International Clinical Consortium on Fibrodysplasia Ossificans Progressiva contraindicates the surgical, radiotherapy and chemotherapy procedures for its treatment. [10] Currently, there is no proven effective prevention or treatment for FOP. [3, 10, 11]

Their life expectancy is approaching 40 years of age, longevity is reduced by the involvement of the intercostal muscles and the diaphragm which leads to respiratory failure. [9, 12]

2. Case Description

12-year-old male who is referred to the Maxillofacial Surgery Department of Pediatric Hospital of the Mexican Institute of Social Security from its General Hospital of Zone (GHZ) for assessment due to limitation of oral opening with 9 months of evolution. In his antecedents of importance, he has the diagnosis of Fibrodysplasia Ossificans Progressiva since he was 5 years old, being subjected to multiple surgical interventions in his GHZ, due to the presence of exostosis in the neck, back and pelvic limbs without beneficial result, evolving in a torpid way. He reported a subcutaneous nodule biopsy in the cervical region 2 years ago with subsequent induration and calcification of soft tissues of the surgical site.

Physical examination revealed a calm, collaborative, normocephalic patient with severe limitation of mandibular movements (1mm) (opening / closing, laterality, protrusion / retraction) with malocclusion, dental malposition and multiple septic foci, as well as poor oral hygiene. Neck with movement restriction at 0°. (Flexion / Extension, Rotation) It presents an increase in the volume of indurated, stony consistency at the level of the sternocleidomastoid muscle (SCM), which compromises its entire extension, with surgical scar at level of zone II. (Figure 1) Kyphosis and cervical and thoracic scoliosis. Multiple dorsal exostosis with pain and limited movement of the elbows, shoulders, thorax, back, hips and knees. Bilateral deformity in the first toes (hallux valgus). (Figure 2) Laboratory tests do not show relevant results. In the CT-3D of the head and neck with sternal extension shows total ossification of the right sternocleidomastoid muscle (SCM), merging it with the mandibular angle, base of the skull, clavicle and ipsilateral sternum, extending to the pectoralis major muscle, supraclavicular, serratus muscles and compromising regions of scapula and humerus. Ossification of trapezius muscle is also observed in its cervical and dorsal portion, as well as the left paravertebral muscles. These calcifications are seen without an apparent vascular compromise. (Figure 3 and 4) Due to the patient's background and its clinical and radiological findings, multidisciplinary management was initiated, being referred to physical rehabilitation, pediatric dentistry, psychology, and nutrition for the integral management of the case. On our part, it was decided to provide a conservative management with periodic reviews.

Figure 1. Intraoral photographs 1a and 1b: right and left laterals, shows the limitation of oral opening (1mm) with bad occlusion and poor dental position is appreciated. Extraroral photographs 1c, 1d and 1e: Restriction of neck movements with an increased volume of indurated, stony consistency at the level of right SCM that compromises its entire length, with surgical scar at the level of zone II.
Figure 2. 2a Multiple dorsal exostosis known as "coral branches". There is evidence of a scar from previous surgical treatment. 2b and 2c Simple x-ray of the thorax and hip that reveals multiple bony bars corresponding to heterotopic ossification at the level of the neck, thorax, spine, extremities and hips, which condition limitation of movements at the back, hip, thoracic and pelvic limbs. 2d and 2e Photograph and simple x-rays of both feet showing bilateral deformity on the first toes. (hallux valgus).

Figure 3. 3a, 3b and 3c CT of head and neck with bony window with sagittal, axial and coronal slices in which the heterotopic ossification of SCM is seen from its origin in the mastoid portion of the temporal bone until its insertion in the right clavicle and sternum, as well as its fusion with the right mandibular angle. A heterogeneous image was seen at the level of the middle third of the corresponding SCM with the area where the patient was biopsied. The anatomy of the TMJ is appreciated without apparent alteration. 3d, 3e, 3f and 3g CT-3D in which the ossification of SCM and the formation of heterotopic bone bars or bridges are observed at the zygomatic apophysis of the temporal bone, mandibular angle, clavicle, sternum, scapula, humerus on the right side, as well as in the dorsal region.
must be addressed in a priority manner due to its potential to cause dyspnea and dysphagia that can lead to the death of the patient. The increase in volume at this level can be confused mainly with cervico-facial infectious processes, as well as with mumps, angioneurotic edema, infectious mononucleosis or neoplasms. (Lasry et al., 2005) [24], (Sellami et al., 2015) [37], (Janoff et al., 1996) [44], (Leavitt et al., 2009) [45], (Awais et al., 2015) [46] Effective treatment includes early identification of submandibular volume increase caused by FOP, nutritional support, glucocorticoid therapy, monitoring, and airway protection. Surgical manipulation of the submandibular region should be avoided because this will worsen the clinical problems of inflammation and accelerate the formation of heterotopic bone. (Janoff et al., 1996) [44]

In the presence of an increase in soft tissue volume at the submandibular level, the International Clinical Consortium on FOP recommends the following: [10]

1. All physicians treating patients with FOP should be aware that an acute volume increase at the submandibular level may be a manifestation of the disease at any age.

2. The diagnosis of FOP should be communicated to the attending physician so that the submandibular lesion can be managed in the context of the patient’s underlying disease.

3. Injury handling should be avoided because even minor trauma can lead to a catastrophic exacerbation with compromised airway.

4. Patients should sleep semi-fowler to reduce the risk of airway obstruction.

5. Patients should be monitored during the acute inflammation phase, they should be hospitalized immediately in case of airway obstruction.

6. The food must be mashed or semi-solid. Fluids often cause episodes of suffocation due to the compromise of the muscles of the floor of the mouth.

7. Patients should be encouraged to eat frequently to minimize weight loss. High-calorie food supplements should be considered.

9. Precautions should be taken to prevent food aspiration.

10. High-dose oral glucocorticoids should be considered in the early treatment of acute submandibular out-brakes. The glucocorticoid of choice is prednisone, 2 mg / kg body weight (up to 100 mg) orally once a day for four days. If it is used a second course of four days of high doses of glucocorticoids, these should be slowly lowered during the next two to three weeks. After discontinuation of glucocorticoid therapy, NSAIDs can be considered for the next 6-8 weeks.

4. Conclusions

The case we present is considered a sequel due to an exacerbation of FOP by the surgical procedure performed 2 years previously, which resulted in a temporo mandibular extra-articular ankylosis with severe limitation of jaw movements, which caused problems of nutrition, oral hygiene, as well as dental problems (bad occlusion, crowding, dental caries among others). Multidisciplinary management was...
initiated by physical rehabilitation, pediatric dentistry, psychology, as well as nutrition for its integral management. On our part, conservative management with periodic reviews was decided.

The maxillofacial surgeon must know the risks of a surgical treatment in patients with FOP and it is necessary to guide the patient in the care of their temporomandibular joints, oral health care and periodic dental check-ups to avoid later major complications.

Currently, there is no effective therapy to prevent or treat FOP, that is why the management must be carried out with a multidisciplinary approach in which diverse health professionals work in a coordinated and joint way to provide a better quality of life for these patients and have a better understanding in the progression of the disease.

Compliance with Ethical Standards

The authors declare that they have not received any type of monetary support for the realization of this case report.

Conflict of Interests

The authors declare that they have no conflicts of interest.

Ethical Approval

The authors declare that for this case report no experiments have been conducted on humans or animals. The authors declare that patient data does not appear in this article.

Informed Consent

Informed consent of the patient was obtained for the publication of this case report and any attached image. A copy of the written consent is available, at any time, for review by the editor of this journal.

References


