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Keywords
Fibrous dysplasia; McCune-Albright syndrome; dental anomalies; gene mutation; oral health

Disciplines
Dentistry

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Dental Perspectives in Fibrous Dysplasia and McCune-Albright Syndrome

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Abstract

McCune-Albright syndrome (MAS) is a rare multisystem disorder characterized by the triad of polyostotic fibrous dysplasia (PFD), endocrine disorders and café-au-lait skin pigmentation. Ninety percent of MAS patients have FD lesions in the craniofacial area, resulting in significant orofacial deformity, dental disorders, bone pain and compromised oral health. Maxillo-mandibular FD is also associated with dental developmental disorders, malocclusion, and high caries index. There is limited data on the outcomes of dental treatments in maxillo-mandibular FD/MAS patients, because clinicians and researchers have limited access to patients, and there are concerns that dental surgery may activate quiescent jaw FD lesions to grow aggressively. This report highlights current perspectives on dental management issues associated with maxillo-mandibular FD within the context of MAS.

Keywords

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INTRODUCTION

Fibrous dysplasia (FD) is a skeletal disorder characterized by replacement of normal bone and marrow by fibrous tissue, leading to fracture, deformity, and pain (Figure 1).1 FD may affect a single bone (monostotic FD) or multiple bones (polyostotic FD), and may occur in association with café-au-lait skin pigmentation and hyperfunctioning endocrinopathies,
including precocious puberty, hyperthyroidism, hypercortisolism, growth hormone excess, and fibroblast growth factor (FGF)23-mediated hypophosphatemia. When FD occurs in combination with one or more extra-skeletal features, it is termed McCune-Albright syndrome (MAS) 2–4. MAS is relatively rare with an estimated prevalence ranging from 1/100,000 to 1/1,000,000, however FD is more common, accounting for approximately 7% of all benign bone tumors 4, 5.

ETIOLOGY AND PATHOGENESIS

MAS arises from somatic activating mutations in GNAS gene, which encodes the Gsα subunit of the heterotrimeric G protein complex6, 7. This results in constitutive activation of adenylyl cyclase and overproduction of 3′,5′-cyclic adenosine monophosphate (cAMP) 7, 8. The severity of the disease phenotype depends on when the mutation occurs during embryogenesis, and the locations where mutated progeny cells subsequently migrate. If the mutation occurs during formation of the inner cell mass, all three germ cell layers will be affected and the individual will develop MAS. Mutations occurring later in embryogenesis will have a more limited phenotype, such as isolated FD, or endocrinopathies without associated bone disease.

At the cellular level, FD is considered a disease of the skeletal stem cell/osteoblastic lineage in which excess cAMP impairs the ability of bone skeletal stem cells to differentiate to mature functioning osteoblasts 9. In normal bone, remodeling is a coordinated cycle of sequential osteoclastic bone resorption followed by osteoblastic bone matrix deposition. In FD bone, remodeling is altered by replacement of normal bone and hematopoietic marrow by abnormal osteogenic tissue and bone trabeculae. The increased bone resorption and fibrous tissue deposition in FD/MAS is associated with increased secretion of interleukin (IL)-6 and RANKL, which contain a cAMP response elements in their promoters 10, 11, 12. Thus, the abnormal patterns of bone formation and resorption that characterize FD/MAS lesions are both related to mutation of Gsα and overproduction of cAMP.

CLINICAL PRESENTATIONS

Clinical signs and symptoms of the skeletal, endocrine and cutaneous components of MAS may present at varying stages of development. Café-au-lait skin hyperpigmentation in MAS is often observed at birth. It has characteristic irregular borders analogous to the ‘coast of Maine’, unlike the smooth borders of the hyperpigmentation in neurofibromatosis, analogous to the ‘coast of California’. MAS associated café-au-lait hyperpigmentation is typically described as “respecting” the midline, either by terminating at or reflecting around the midline (Figure 1). This distribution pattern reflects patterns of early embryonic cell migration. There is a wide clinical spectrum of FD severity, depending on the amount and areas of the affected skeleton (Figure 1). As FD patients grow older, features of FD in the appendicular skeleton may manifest with fracture, limp or bone pain, which in small children may manifest as complaints of tiredness and easy fatigability 4. Also, long bones affected by FD are prone to fracture and progressive deformities such as coxa vara of the proximal femur, commonly termed the “shepherd’s crook” deformity (Figure 1). Interestingly, facial asymmetry may be the first sign of craniofacial FD, and it may be associated with hearing and more rarely vision loss 13. Precocious puberty is common in girls, and occurs due to spontaneous development of functioning ovarian cysts. This presents as episodic periods of estrogen excess, early breast development, growth acceleration, and vaginal bleeding. Precocious puberty is less common in boys, however they frequently present with macro-orchidism and testicular ultrasound abnormalities consistent with Leydig and Sertoli cell hyperplasia (Figure 2)14. Hyperthyroidism typically develops during infancy or early childhood, and may be diagnosed by abnormalities seen on thyroid function tests or...
radiographic abnormalities on thyroid ultrasound (Figure 2)\textsuperscript{15}. Patients with growth hormone excess from MAS-associated pituitary disease (Figure 2) initially present with growth acceleration. If untreated, growth hormone excess causes craniofacial expansion and optic neuropathy, so early diagnosis and treatment is a critical component of management\textsuperscript{14, 16}.

**DIAGNOSIS**

FD/MAS is a clinical diagnosis based on a combination of clinical, biochemical, and radiographic findings. Mutation analysis of FD tissue is problematic due to the mosaic nature of the disease, and is rarely helpful for clinical management. All patients suspected of having FD/MAS should undergo screening for thyroid and pituitary involvement that include thyroid function tests, thyroid ultrasound, and insulin growth factor (IGF)-1 level. Although testicular ultrasound is a helpful diagnostic tool in boys, routine pelvic ultrasound is rarely informative in girls without clinical signs of precocious puberty due to the episodic nature of ovarian cysts. It is essential to monitor growth velocity in children, because all of the endocrine manifestations of MAS can cause growth alterations. In children ages 5 years and older, technetium (99mTc-methylene diphosphonate) bone scintigraphy is useful to identify areas of FD, which can be further evaluated with plain radiographs (Figure 1)\textsuperscript{17, 18}. The radiographic appearance of FD is affected by both location and disease activity. During childhood, appendicular and craniofacial FD demonstrates a homogenous ‘ground glass’ appearance. The disease activity typically declines during adulthood, at which point FD may adopt a more sclerotic, heterogeneous appearance.

This is illustrated in Figure 3, where panoramic radiographs and computed tomography (CT) of the jaws demonstrate ground glass trabeculation that also includes mixed radiolucent/radio-opaque lesions and thinning of the cortical margin\textsuperscript{19}. FD in the axial skeleton (including the spine, ribs, and pelvis) may be difficult to visualize on plain films, so bone scintigraphy is useful to detect lesions in these areas\textsuperscript{17}. Histologically, FD demonstrates unique site-specific features\textsuperscript{1, 20}. In the cranial bones, it displays a dense, sclerotic trabecular bone pattern with an interconnected network similar to the pattern observed in Paget’s disease. In the jaws, the pattern is characterized by the presence of significant amounts of sclerotic bone, however, bone trabeculae are discontinuous, and in the axial/appendicular bones, they display a ‘Chinese character pattern’ with the fibrous tissue predominating over the abnormal bone trabeculae (Figure 4)\textsuperscript{1}. The lamellation pattern seen in normal bone is conspicuously absent in the\textit{de novo} bone formed within areas of FD. Rather FD exhibits a pattern consistent with woven bone.

**MEDICAL MANAGEMENT**

The multi-organ clinical pattern makes dental and medical management of patients with FD/MAS complex and challenging. Therefore, coordination of comprehensive care involving interactions between dentist, endocrinologist, primary care provider, orthopedic surgeon, physical therapist and social workers is essential. The endocrine features of FD/MAS are treated either by surgical removal of affected glands or by medications directed at hormonal inhibition or blockade\textsuperscript{21}. However, there is still no satisfactory treatment to alter disease progression of FD. Surgical treatment of FD involves orthopedic hardware to stabilize bones and mediate deformity. Discouragingly, these existing methods of surgical treatments still have unsatisfactory outcomes\textsuperscript{13, 22, 23}.

In the craniofacial region, surgery is indicated when the lesion causes loss of function of vital structures such as optic nerves, or for cosmetic recontouring to improve appearance. Screening for and treatment of FGF23-mediated hypophosphatemia is a critical component...
of FD management, as phosphate wasting may worsen bone pain and place patients at higher risk for fracture and deformity. Bisphosphonates have been moderately effective in relieving FD-related bone pain, but are ineffective in altering the disease course. The bisphosphonate dosing regimen in FD/MAS is similar to that of Paget’s disease because the drug is cycled ‘on and off’ so patients are often off bisphosphonate for several months until pain recurs. FD/MAS patients do not routinely undergo treatment for skin hyperpigmentation (Figure 1), although there has been one report of successful laser therapy for a facial café-au-lait macule.

**ORAL AND DENTAL IMPLICATIONS**

Approximately 90% of patients with FD have lesions in the craniofacial bones including the maxilla and mandible. FD can cause massive expansion of the craniofacial complex, severe malocclusion and facial disfigurement. FD lesions can grow rapidly, leading to bone expansion and displacement of adjacent structures such as the orbit and teeth. The metabolic dysfunctions and disordered bone architecture in FD/MAS can potentially affect tooth development and eruption. Although FD is a disease of the skeletal stem cell/osteoblastic lineage in which excess cAMP impairs the ability of the stem cell to differentiate to mature functioning osteoblasts, it is unclear whether excess cAMP affects the developing tooth, either directly or indirectly.

There is clinical concern that FD/MAS can disrupt the coordinated sequential events involved in replacement of primary dentition with permanent dentition, but it is also unclear if the presence of FD in the jaws has any effects on tooth development and function. FD is associated with dental disorders such as enamel hypoplasia, dentin dysplasia, taurodontic pulp, odontoma, tooth displacement, malocclusion, and high caries index. These anomalies are seen in approximately 28% of patients with craniofacial FD, and have significant impact on their oral health status and management. Pain is common in craniofacial FD, affecting approximately 40% of patients. Rapid growth of jaw FD may also be associated with pathological lesions such as aneurysmal bone cysts, or more rarely malignant transformation to osteosarcoma or other forms of sarcoma. Taurodontism, a condition characterized by enlargement of pulp chamber, is often associated with presence of endocrine disorders in FD/MAS, suggesting that FD, endocrinopathies, or a combination of these may impact tooth development in FD/MAS patients.

**DENTAL MANAGEMENT ISSUES**

Dental management of FD/MAS is medically complex because treatment of dental disorders must be balanced with multiple factors, including skeletal disease burden, endocrine disorders, multiple medications and general debility. Due to the presence of often-complex medical comorbidities, the dental aspects of FD/MAS are frequently overlooked, and dental needs are often underserved. Furthermore, the variable clinical, radiological and histological presentations of FD, as well as apparent risk of malignant transformation, cause some dental practitioners to delay or avoid dental surgical procedures in FD. Also, some dental healthcare providers may also feel uneasy about treating FD/MAS patients because of previous subjective reports that dental surgery might exacerbate jaw FD, transforming a quiescent lesion to an aggressively growing lesion. There are also concerns regarding bisphosphonate-associated osteonecrosis of the jaw (ONJ). However, ONJ is rare in FD/MAS, possibly due to the relatively lower bisphosphonate dosing regimens compared to doses used in cancer patients, or perhaps due to the relatively hypervascular nature of FD. While bisphosphonate doses used in FD/MAS are higher than...
those used in osteoporosis therapy, risk of ONJ does not appear to be greater than that seen in osteoporosis patients.

Due to the severity of dental malocclusion and high caries index in patients with maxilla-mandibular FD, more frequent recalls may be required for scaling and root planning to control dental plaque accumulation. The use of electric toothbrush and application of topical fluoride may be helpful to control dental caries. When caries extends to the pulp chamber in a tooth with a taurodontic pulp, root canal therapy may be challenging. This is an indication for prompt referral to an endodontist for management. Severe malocclusion may also require orthodontic intervention. When orthodontic therapy is clinically indicated, the timing should be carefully coordinated among the dental healthcare providers. Literature reports on outcomes of orthodontic therapies in FD/MAS are still unclear. One theory is that orthodontic tooth movement tends to be rapid in FD jaw. Our experience is that orthodontic therapy takes much longer than in normal patient population; and relapse is more common because teeth tend to return to their original position after removal of orthodontic appliances. It may be advisable to delay orthodontic therapy till after the age of skeletal maturity based on individual patients’ needs and outcomes of orthodontic evaluation. The advantage is that FD disease activity tends to decrease after skeletal maturity, which is reflected in a decline in bone turnover markers, a decrease in the number of mutated lesional skeletal stem/progenitor cells, and a tendency of FD histology to improve over time. While root canal therapy in a taurodontic tooth and orthodontic tooth movement in FD-related malocclusion are some clinical challenges faced by FD/MAS patients, it is still unclear if other dental anomalies would pose any treatment challenges. Disorders of the liver, heart, and spleen have been associated with MAS, so it is imperative to consult with the patient’s physician on medical stability for dental therapy before embarking on extensive dental surgery.

Unfortunately, there is limited data on effectiveness and outcomes of dental therapies in maxillo-mandibular FD/MAS patients because clinicians and researchers have access to very few and small patient pools. To address the healthcare needs of FD/MAS patient populations, the National Institutes of Health (NIH) Office of Rare Diseases Research (ORDR), the National Institute of Dental and Craniofacial Research (NIDCR), and the Fibrous Dysplasia Foundation organized an International meeting in October 2010 in Bethesda, MD to determine best clinical practice and future research on FD/MAS. One clear outcome was that clinical and basic scientists must share research data and tissue samples to foster new research that will expand our understanding of FD/MAS and improve dental and medical treatment outcomes.

CONCLUSION

The broad clinical spectrum of FD/MAS makes dental care challenging and medically complex. Therefore, healthcare providers often refer FD/MAS patients to dentists equipped with special patient care facilities. Literature reports so far indicate that routine dental care can be safely and successfully carried out in FD/MAS patients with minimal complications. However, there are still several unanswered questions: Is healing delayed in FD bone after tooth extraction? Does dental surgery aggravate FD lesion? Are FD/MAS patients more prone to endodontic therapy, since the variable radiographic FD patterns in the alveolar region may mimic pulpally induced periapical lesions? Is orthodontic therapy unusually protracted in FD/MAS patients? Are FD/MAS patients easily prone to orthodontic treatment relapse due to poor quality of FD bone with teeth moving away from where they were orthodontically positioned? These are areas that additional research can address to expand our understanding of the outcomes of dental extractions, dental implants,
root canal therapy and orthodontic therapy in FD/MAS patients with maxillo-mandibular FD.

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References


FIGURE 1. Images of a 12-year-old girl with fibrous dysplasia/McCune-Albright syndrome (FD/MAS)
A) Photograph showing café-au-lait pigmentation on the posterior trunk (black arrow), and expansion in FD of the left femur leading to deformity and limb-length discrepancy. Shortening of the left lower extremity has resulted in an altered stance with pelvic obliquity (white arrowhead). B) Technetium-99 bone scintigraphy shows patchy areas of increased radiotracer uptake in FD in the skull, humeri, radii, femurs and fibulas (black arrows). FD in the proximal femurs has led to bilateral coxa varus (“shepherd’s crook” deformities) (black arrowheads). C) A characteristic facial café-au-lait macule is limited to one side without crossing the midline (‘respecting the midline’) and exhibits typical ragged “coast of Maine” borders. D). Radiographs show bilateral femoral FD and “shepherd’s crook” deformities (white arrowhead) that is more severe on left.
FIGURE 2. Radiographic images of endocrine organs in McCune-Albright syndrome

A) Testicular ultrasound of a patient with McCune-Albright syndrome (MAS) shows a heterogeneous, mixed cystic and solid lesion characteristic of Leydig and Sertoli cell hyperplasia (white arrows). B) Thyroid ultrasound in a patient with MAS-associated hyperthyroidism demonstrates characteristic heterogeneity with a cystic, “Swiss cheese”-like appearance (white arrow). C) Magnetic resonance imaging of the brain in a patient with MAS-associated growth hormone excess shows a large, dumbbell-shaped growth hormone-and prolactin-secreting pituitary macroadenoma (white arrows); the bright spot (white arrowhead) indicates the posterior pituitary.
FIGURE 3. Images of an adult woman with McCune-Albright syndrome and craniofacial fibrous dysplasia
A) Facial asymmetry resulting from expansion of a fibrous dysplasia (FD) lesion in the right mandible. B) Panoramic radiograph shows a mixed radiolucent/radio-opaque pattern characteristic of FD. C) Computed tomography (CT) of the head shows expansion of FD in the right mandible (arrow). D) Three dimensional volume rendering of CT images shows expansion of FD in the right mandible (white arrow), as well as multiple smaller areas of FD (black arrowheads).
FIGURE 4. Histological features of fibrous dysplasia
Hematoxylin/eosin stained sections of maxillary fibrous dysplasia [FD] (A, low power and B high power) demonstrate haphazard trabecular bone matrix (‘Chinese character’ pattern, black arrowheads) within a fibrous stroma. Goldner’s trichrome stained sections from calvarial FD (C, low power and D, high power) demonstrate similar haphazard bone matrix deposition (back arrowheads) and unmineralized regions (white clear arrows) that indicate osteomalacia [ft = fibrous tissue].