A Case Report of Juvenile Aggressive Ossifying Fibroma [Trabacular Type] in a young Patient

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Abstract
Juvenile aggressive ossifying fibroma is a rare variant of fibro-osseous lesion occurring primarily in children and young adults. It is a benign, but osteogenic neoplasm of bony origin with a potentially aggressive nature. On the account of age of onset, site of occurrence, clinical presentation, histological feature, aggressive nature and increased tendency for recurrence, it is differentiated from the larger group of ossifying fibroma as a separate entity. This article reports a case of juvenile aggressive ossifying fibroma of trabecular pattern in the anterior maxilla of a 6 year old male patient.

Keywords: Trabecular juvenile ossifying fibroma, Fibro-osseous lesions, Maxilla, Aggressive.

Introduction
Juvenile ossifying fibroma (JOF) is described in literature as a rare variant of ossifying fibroma, occurring in the craniofacial skeletal of young individuals[1]. Based on the type of mineralized component, two histological variants are recognized namely, Psammomatoid juvenile ossifying fibroma (PsJOF) and Trabecular juvenile ossifying fibroma (TrJOF, WHO (World Health Organisation) Type)[1]. The most significant clinical feature that helps to differentiate TrJOF from PsJOF is the site of involvement, with PsJOF occurring mainly in the paranasal sinuses and TrJOF occurring mainly in the maxilla[2]. Here one such occurrence of this rare lesion is reported in the maxilla of a 6 year old patient.

Clinical history
A 6-year-old male child reported to the Department of Pedodontics and Preventive Dentistry, Rajah Muthiah Dental College and Hospital, with the chief complaint of rapidly enlarging mass for past 30 days over the upper front region of jaw. The history of presenting illness revealed that, the swelling started off as a small painless mass that gradually increased in size over a period of one month with mild discomfort along with bleeding while brushing. The past history revealed that, the patient had undergone an excision for a similar lesion at the same site, by a private practitioner elsewhere 6 months back, which recurred. The patient had no contributing family and medical history. No clinically relevant finding could be detected on extra-oral examination. Intraoral examination...
revealed an asymptomatic ovoid swelling measuring approximately 1.5 cm x 1 cm extending mesiodistally from mesial half of upper right primary central incisor to the mesial half of upper left primary lateral incisor; superiorly extending up to the marginal gingiva and inferiorly to the level of palatal mucosa with labially displaced upper left primary central incisor. Secondary changes such as redness and mild ulceration were seen over the swelling. There was no evidence of abscess, draining fistulae or sinuses associated with the swelling. On palpation, the swelling was uniformly firm in consistency, non tender and non fluctuant. Hard tissue examination showed Grade I mobility in deciduous right and left central incisors. (51 & 61)

**Fig. 1** Preoperative photograph showing the lesion

Radiological investigation included an intraoral periapical radiograph of the lesion which showed evidence of root resorption of primary teeth (51&61). Routine hematological evaluation ascertained values within normal limits. Based on the clinical findings and presentation, the differential diagnosis established were fibroma, osteofibrous dysplasia and fibrous dysplasia.

**Fig. 2** Intraoral periapical radiograph of 51, 61, 62 showing root resorption of 51, 61

Excisional biopsy was performed under local anesthesia along with the extraction of both the upper central incisors. The excised specimen was fixed in 10% formalin and sent for histopathological examination. A follow-up examination of the patient after nine months showed no sign of recurrence or relapse. The patient is under regular clinical and radiological follow-ups due to the recurrent nature of the lesion.

**Fig. 4** Postoperative photograph after 6 months

**Investigation**

Histopathological examination revealed numerous trabeculae of forming bone [‘a’ in fig.5] in a highly cellular connective tissue. The connective tissue showed monomorphic fibroblast with spindle to plump nuclei [‘b’ in fig.5] and indistinct cytoplasm. The bony trabeculae showed osteoid matrix in the periphery [‘c’ in fig.5]. Few chronic inflammatory cells were also noted all suggestive of juvenile ossifying fibroma of trabecular pattern. All of the above finding pointed towards a definite diagnosis of trabecular variant of juvenile ossifying fibroma.

**Fig. 5** Photomicrograph showing epithelium and underlying connective tissue with calcifications. (Original magnification ×400)

**Discussion**

According to WHO (World Health Organisation) classification (2005), ossifying fibroma most commonly occurs in the 2nd to 4th decades with a female predilection. The histological subtypes...
varies in accordance with mean age of the patients. The subtypes includes, JPOF (Juvenile psammomatoid ossifying fibroma) occurring in patients around 20 years, while that of conventional ossifying fibroma occurring in patients around 35 years of age. JTOF (Juvenile trabecular ossifying fibroma) has a still lower mean age range of 8.5-12 years\(^3\).

JTOF also known as trabecular desmo-osteoblastoma, affects mainly the jaws of children and adolescents. Only 20% of the patients are over 15 years of age. In a review of a number of case series the mean age range was found to be 8.5-12 years\(^{4,5,2}\).

This present case report attempts at the documentation of a rare entity of juvenile ossifying fibroma occurring in a 6 year old male child. The patient reported with a chief complaint of swelling in upper jaw with a previous history of treatment for the same. As the lesion recurred and grew rapidly in the present case, a more aggressive behavior of juvenile ossifying fibroma was provisionally diagnosed which probably occurred due to incomplete initial removal of the lesion from the prior treatment or persistence of some local irritants. The site of occurrence and the histo-pathological picture further confirmed the final diagnosis as a trabecular pattern of juvenile aggressive ossifying fibroma.

The few commonly suggested etiology for the occurrence of aggressive pattern of juvenile ossifying fibroma in young patient are the high levels of periodontal ligament activity (e.g., formation and degradation) and the constant irritation associated with both primary tooth exfoliation and permanent tooth eruption in children\(^{6,7}\). In the present case, the pre-shedding mobility associated with 51, 61 along with the eruption of 11, 21 could be suggested as a possible etiological factor contributing to the recurrence of the lesion after the initial treatment. Juvenile ossifying fibromas are clinically and histologically distinct from other fibro-osseous lesions, in that it has a recurrence rate of 30 to 56%\(^1\), thereby necessitating a more radical approach rather than conservative curettage\(^{8,9}\).

Accordingly, in the present case, the line of treatment performed was surgical resection, which was highly effective with no signs of recurrence even after a follow up of 6 months [Figure 4].

The aggressive local behavior of juvenile ossifying fibroma and the rarity of this condition along with the unpredictable recurrence at varied intervals calls for an early diagnosis, prompt treatment, and especially, long-term follow up of the patient. Elimination of etiological factors and complete surgical excision along with involved periodontal ligament and periosteum minimize the possibility of recurrence. The case presented here is an unique variant of juvenile ossifying fibroma of trabecular pattern that has occurred in a very younger age in comparison to the expected age of occurrence and this necessitates the need for further studies for the better understanding of juvenile ossifying fibroma.

References


