Juvenile Psammomatoid Ossifying Fibroma of Fronto-Ethmoid Complex mimicking Fibrous Dysplasia: Case Report

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ABSTRACT

Juvenile Psammomatoid Ossifying Fibroma is a rare fibro-osseous tumor seen in children and adolescent and mostly arising from the cranio-facial bone. We report a case of 18-year-old boy who presented with diplopia and progressive right fronto-orbital swelling. On plain radiograph and CT, it was diagnosed as fibrous dysplasia and mucocele as differential diagnosis. The tumor was resected and histopathological examination showed psammomatoid features. Therefore, the diagnosis of Juvenile Psammomatoid Ossifying Fibroma must be based on both radiological and histopathological findings.

Keywords: Diplopia; Ethmoid Sinus; Fibroma, Ossifying

INTRODUCTION

Juvenile ossifying fibroma (JOF) is a benign fibro-osseous tumor commonly seen in young patient in cranio-facial region. It is classified under variant of ossifying fibroma.1 There are two known histological variants of JOF: Juvenile Psammomatoid Ossifying Fibroma (JPOF) and Juvenile Trabecular Ossifying Fibroma (JTOF).2 JPOF is locally aggressive tumor and can be confused for malignant lesion. It is very difficult to diagnose JPOF by only looking at the radiological findings as they may resemble other fibrous osseous lesions in similar locations. Here we present a case of JPOF involving the fronto-ethmoid complex mimicking Fibrous Dysplasia (FD).
CASE REPORT

A 18-year-old male presented to the OPD with history of diplopia and swelling on superior aspect of right eye for few months. On physical examination, patient had proptosis of the right eye. His medical and family histories were unremarkable. Plain radiograph of skull bone followed by contrast enhanced computed tomography (CT) of the brain and orbit was done. Plain radiograph showed an expansile lytic lesion of size 5×4 cm involving right frontal and ethmoidal sinuses (Fig 1).

CT scan showed a well-defined expansile lytic lesion predominately involving the superior wall of the orbit causing compression of the orbital cavity and thinning out of the inner table of right frontal bone. The lesion was predominantly of fluid attenuation with few areas of ground glass type of matrix in the lateral aspect (Fig 2). No significant enhancement was noted on post contrast study. On the basis of clinical scenario and radiological features of benign nature and few areas of ground glass matrix within, we considered fibrous dysplasia as possible diagnosis with mucocele of frontal sinus as close differential diagnosis. The patient underwent right-sided frontal craniotomy and total excision of the tumor was done and defect was covered with bone graft. The excision material was submitted as right frontal sinus mucocele with bone fragments. On gross examination, multiple bits of irregular grey-white to grey brown to blackish soft tissue was observed. Cut section showed solid grey white appearance. Representative sections examined revealed histological features consistent with benign fibro osseous lesion-Juvenile Psammomatoid Ossifying Fibroma.

DISCUSSION

Ossifying fibroma are categorized as conventional and juvenile forms.² The psammomatoid type of Juvenile Ossifying Fibroma is more common than the trabecular...
variety. It is also moreaggressive and has a high tendency to recur compared to trabecular variety. JPOF is a relatively rare fibro-osseous lesion occurring in children and young adults. In our case, the patient was in his late teen. JPOF usually occurs in orbit and paranasal sinuses as seen in our case while JTOF usually occurs in maxilla and mandible. Among facial bones, the paranasal sinuses are affected in 90% of JPOF cases among which the ethmoid sinus is the most common site. In our case, the tumor was affecting both frontal and ethmoidal sinuses. The most common complaint in patient with JPOF is proptosis. Other uncommon complaints include swelling of the face, disturbance in ocular movement, headache, and sinusitis, depending on the location of the tumor. In our case, the patient presented with facial swelling along with proptosis of right eye.

On conventional X-ray, JPOF appears as round oval expansile lytic lesion with well-defined margin. Similar findings were noted in our study. The radiographic appearance can vary ranging from cystic to sclerotic depending upon the tumor content.

On CT scan, JPOF is seen as enhancing soft tissue tumor along with multiple internal foci of calcifications and areas of fluid attenuation within due to cystic changes. These cystic changes may appear multiloculated due to thin internal septations. Some of the cystic areas may show hemorrhage within and subsequently develop secondary ABCs. JPOF mostly has well-defined margin, although some cases may show disruption of the surrounding sclerotic rim. Multiple bones may be involved along with bowing and thinning of the involved bones. JPOF may show areas of “ground glass” matrix within the lytic area as seen in our case mimicking the appearance of fibrous dysplasia.

The most important differential diagnosis of JPOF is fibrous dysplasia. JPOF is mostly unilocular, monostotic and has oval shape with well-defined margin and often lacks typical ground-glass attenuation along with multiple punctate areas of calcification within. Similarly, FD tends to be more elongated rather than ovoid appearing JPOF and the border of fibrous dysplasia is ill-defined and converges with surrounding normal bone. FD often appears as poorly circumscribed lesion causing bony expansion and thinning of outer cortex and shows variable radiolucent areas within depending on the proportion of fibrous components giving a characteristic “ground glass” appearance on CT scanning.

Other differential diagnosis of JPOF include demucocele, ABC, osteoblastoma, osteosarcoma and cemento-osseous dysplasia and cementifying fibromas. Mucocele may be associated finding with ossifying fibroma thereby causing difficulty in diagnosis. JPOF is a rare clinical entity often misdiagnosed because of its rapidly growing and aggressive nature. The present case was also misdiagnosed as fibrous dysplasia. The distinction between psammomatoid variant of juvenile ossifying fibroma, when it affects the paranasal sinuses and fibrous dysplasia, can be quite challenging especially when both present with ground glass opacity. Ossifying fibroma and fibrous dysplasia may share similar radiological features. Hence, the pathological findings of the lesion should also be evaluated side by side when radiological differentiation is uncertain.

CONCLUSION

Careful assessment of clinical, radiographic and histopathologic features is necessary for differentiation of JPOF from other fibro-osseous lesions and to arrive at the appropriate diagnosis and for necessary therapeutic intervention.

CONFLICT OF INTEREST
None

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REFERENCES


