ABSTRACT
Psammomatoid juvenile ossifying fibroma (PsJOF) is a well defined clinical and histological entity with earlier onset at (childhood or adolescence) classified under the broad category of fibro-osseous lesion. Histological PsJOF is one of the entity of juvenile ossifying fibroma (JOF) characterized by small spherical ossicles resembling psammoma bodies and the other with trabecular or fibrillar osteoid and woven bone which is termed as trabecular juvenile ossifying fibroma (TrJOF). This case of PsJOF involved maxilla, premaxilla, antrum, lateral wall of nose on left side of face. The size, extent and aggressive behavior of the tumor which has lead to facial disfigurement and difficulty in breathing, inability to speak, eat and drink is rare in literature. The treatment done by resection with safe margins is adequate with no recurrence as patient is under follow up for more than one year.

KEY WORDS: Psammomatoid juvenile ossifying fibroma; Juvenile ossifying fibroma; Cement- ossifying fibroma; Ossifying fibroma; Fibro-osseous tumors.

INTRODUCTION
Juvenile ossifying fibroma is a well defined clinical and histological entity which is a subtype of ossifying fibroma, belonging to a group of Fibro-osseous lesion (FOL). The term fibro-osseous lesion is a generic designation of a group of jaw disorders characterized by fibrous tissue containing mineralized structures that may resemble bone or cementum. This group include developmental, reactive or dysplastic lesions as well as neoplasm. When ossifying fibroma are diagnosed in young people the term “Juvenile” is used but cases up to 70 years have been reported. The group of JOF describes two distinct clinic patholgical variant i.e psammomatoid juvenile ossifying fibroma [PsJOF] and trabecular juvenile ossifying fibroma [TJOF] of cranio facial skeleton.

Psammomatoid juvenile ossifying fibroma is rare fibro-osseous neoplasm benign in nature and aggressive in growth. Most of the PsJOF simulating malignancy can be a worrying factor. The lesion is encapsulated and well demarcated from surrounding bone.

Benjamins was first to describe this lesion as osteoid fibroma and later Gogi in 1949 was first to designate it as psammomatoid (juvenile) ossifying fibroma of nose and paranasal sinuses.

Case Report
A 14 years old girl was referred to the department of oral and maxillofacial surgery for the treatment of maxillary tumor which was noticed in her mouth when she was 12 years of age. Patient had taken consultations for the same, but did not undergo definitive treatment due to low socio-economic status. The tumor mass was massive seen protruding from the mouth due to which patient complained of difficulty in breathing, eating and drinking. She stated that she could not close her lips since last 6 months (Fig.1). All her vitals and laboratory investigations were within normal limits except hemoglobin (7.8g/dl).

Clinical examination revealed a tumor protruding out of the mouth with open mouth appearance, displaced anterior teeth, with mucosa over the swelling appearing to be blanched and stretched. Tumor mass was non ulcerated, non tender, firm to bony hard in consistency. Intra oral examination was not possible as there was hardly any space for inspection or palpation. Left side submandibular lymph nodes were palpable and tender.

Computerized tomogram (Fig.2) revealed tumor in whole maxilla with definite margins eroding
Fig.1. Prooperative view of the lesion

Fig.2. Coronal section C.T. Scan

Fig.3. Pathological section of the excised mass

Fig.4. Matured osteoid tissue surrounded by fibrous tissue.

Fig.5. Dense fibrous tissue with focal Spherical Basophils.

Fig.6. Post operative view
maxillary sinus, zygomatic bone and lateral wall of nose. It shows a mixture of radiodensity and radiolucency. Based up on aggressive behaviour and clinico-radiographic interpretation the differential diagnosis of osteogenic sarcoma, chondrosarcoma, giant cell tumor, ossifying fibroma, fibrous dysplasia ,cento-ossifying fibroma was made.Histopathology of Incisional biopsy specimen revealed the lesion as Juvenile Ossifying Fibroma (Fig.3 & 4). The tumor was resected after securing airway by performing tracheostomy with safe margins as planned with Weber-Ferguson incision (Fig.5). Posteriory debulking was done. After hemoastasis was achieved and the surgical site was closed by mucocutaneous and mucosal flap and some area left to granulate. After complete healing, patient was referred to the department of prosthetics for prosthetic rehabilitation. An obturator was fabricated which aided in improving esthetic profile, phonetics and mastication .The tumor mass which weighed 1 kg was sent to histopathological examination to study under different sections.Finally the lesion was reported as Psammamotoid juvenile ossifying fibroma.

Discussion

As per WHO classification, Psammomatoid Juvenile Ossifying Fibroma (PsJOF) is grouped under category of Ossifying Fibroma (Table 1). The literature regarding its incidence, behaviour, location, treatment options and recurrence vary.

The diagnosis is confusing but should be based on age of onset, clinical behaviour, radiographic interpretation and detail histopathological review. When initial histopathology reports as Ossifying Fibroma, it is advised to go for deeper sections of the specimen to study the other three variants i.e., Conventional ossifying fibroma, Juvenile trabecular ossifying fibroma and Juvenile psammomatoid ossifying fibroma (Table 2).

Most of the authors considers Cemento-ossifying fibroma [COF] and Juvenile Ossifying fibroma to be a single entity. Samir El-Mofty clarifies COF as a separate lesion which is well encapsulated, predominantly seen in women in third and fourth decades, affects tooth bearing areas of jaws composed of mature and immature bone trabeculae and lobulated basophilic masses of cementum like material. Many a times concentrically laminated particles, called cementicles are seen. In contrast, the microscopic features of PsJOF are distinct and include a cell rich fibrous stroma containing bands of cellular osteoid without osteoblastic lining, osteoid strands and trabeculae of woven bone characterised by innumerable round cement-like or psammoma-like particles called ossicles. PsJOF is usually reported as asymptomatic, although facial asymmetry, nasal obstruction, epistaxis and proptosis were common findings.

The common sites for PsJOF were paranasal sinuses,orbital region, maxilla, mandible and other bones of craniofacial region. This case primarily involves premaxilla, maxillary sinus and zygomatic bone.

Juvenile ossifying fibroma has varied radiological features like radiolucent, mixed and radiopaque depending on the degree of calcification by tumor and sometimes ‘Ground Glass’ appearance . The other lesions which radiographically mimic Juvenile ossifying fibroma are Fibrous dysplasia and Cemento-ossifying fibroma. Fibrous dysplasia blends with surrounding bone and the borders are not well defined. Cemento-ossifying fibroma has well defined sclerotic border, most of the times with tooth bearing areas, representing a fibrous capsule which may separate from the surrounding bone. Maxillary Chondrosarcoma typically occurs in the anterior region where cartilagenous tissue may be present. Radiologically it gives a mixed radiolucent and radiopacity with calcification in the center, sometimes described as ‘flocculent’ implies snow like features. Careful examination of the areas of flocculent may reveal a central nidus, which is probably cartilage surrounded by calcification. The other clinical entity which was also included in the differential diagnosis i.e., Osteosarcoma is usually radiolucent with no peripheral sclerosis or capsulation. If lesion involves peristomeum, the typical ‘Sun Ray Spicules’ or ‘Hair on End’ trabeculae may be seen.

Table 1. Classification of fibro-osseous lesions of the maxillofacial region

<table>
<thead>
<tr>
<th>Fibrous dysplasias</th>
<th>Monostotic fibrous dysplasia</th>
<th>Polystotic fibrous dysplasia</th>
<th>Craniofacial fibrous dysplasia</th>
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</thead>
<tbody>
<tr>
<td>Osseous dysplasias</td>
<td>Peri apical osseous dysplasia</td>
<td>Focal osseous dysplasia</td>
<td>Florid osseous dysplasia</td>
</tr>
<tr>
<td>Familial qianqiform dysplasia</td>
<td></td>
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</tr>
<tr>
<td>Ossifying fibromas</td>
<td>Conventional ossifying fibroma</td>
<td>Juvenile trabecular ossifying fibroma</td>
<td>Juvenile Psammomatoid ossifying fibroma</td>
</tr>
</tbody>
</table>

#Based on the WHO classification and also from Waldron, Slootweg, Bannon, Fowler and Elmofty.
Table 2. A working classification of the three clinicopathological variants of ossifying fibroma

<table>
<thead>
<tr>
<th>Ossifying fibroma</th>
<th>Juvenile trabecular Ossifying fibroma</th>
<th>Juvenile Psammomatoid Ossifying fibroma</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>10-50 years</td>
<td>20-30 years</td>
</tr>
<tr>
<td>Female: Male</td>
<td>2:1</td>
<td>2:1</td>
</tr>
<tr>
<td>Site</td>
<td>Tooth bearing areas</td>
<td>Mandible-50%, Maxilla-44%</td>
</tr>
<tr>
<td>Radiology</td>
<td>well circumscribed</td>
<td>well circumscribed, Spotted calcification</td>
</tr>
<tr>
<td>Histology</td>
<td>Cellular fibrous tissue, storiform pattern, trabeculae of woven bone, occasional spherical Psammomatoid calcification.</td>
<td>Densely cellular immature osteoid and bony in trabecular pattern. Osteo blast rimming and cellular osteoid</td>
</tr>
<tr>
<td>Synonyms</td>
<td>Cemento-ossifying</td>
<td>Juvenile active ossifying</td>
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This case which is reported deserves attention due to size and presentation leading to facial disfigurement and asymmetry. Due to the massive tumor size, dysphagia and difficulty in breathing were associated symptoms which has not been documented before in literature review for PsJOF. Surgical resection with safe margins was proved to be adequate if done properly without recurrence. A high recurrence rate of 30-56% is more likely due to incomplete excision and infiltrative nature of the tumor. To conclude, it can be stated that JOF and its variants despite different histological patterns share clinical and radiological features. The management is also same which depends on size, site and nature of growth. These forms are uncommon but there is a need to recognise them separately and manage appropriately. However, the case reported here is followed up for more than one year without recurrence. The importance of close monitoring and long term follow up cannot be ruled out.

References

Corresponding Author
Dr.Syed Ahmed Mohiuddin M.D.S
Principal, Professor and Hod
AL-Badar Dental college and Hospital
SY.NO 12, Daryapur Village, Nagarahalli,
Near koranti hanuman temple,
Gulbarga -585103
Email ID : dsyedahmed28@yahoo.com