Odontogenic Myxoma of the Maxilla Infiltrating the Maxillary Sinus: A Rare Case Report

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Abstract: Odontogenic myxomas are tumors derived from embryonic mesenchymal elements of dental anlage. Although benign, odontogenic myxomas draw great attention because of their invasive nature into the surrounding tissues, which make the tumour amenable to recur especially after conservative treatment. We present a case of odontogenic myxoma with a displaced maxillary molar in the sinus, an unusual radiographic finding reported in literature.

Key words: Odontogenic, Myxoma, Mesenchymal, Tumor, Benign

I. Introduction

Odontogenic myxoma (OM) is an uncommon, mucoid neoplasm of oral cavity. As it resembles the dental pulp microscopically, in 2003, the World Health Organization (WHO) classified it as a benign neoplasm arising from odontogenic ectomesenchyme with or without odontogenic epithelium, ultrastructural features suggesting many lesional cells to be very similar to fibroblast and myofibroblasts. It is regarded as a locally invasive tumour displaying aggressive infiltration, that does not metastasize and exhibits slow and asymptomatic expansion, sometimes resulting in perforation of the cortical borders of the affected bone.

The radiographic appearance of odontogenic myxoma of the jaws varies considerably. Large lesions may exhibit characteristic radiological signs of a slowly growing lesion. However, discrete displacement of teeth associated with a small osteolytic zone of the alveolar process between two teeth can be an OM. Careful interpretation of conventional radiographs is a must in identifying early lesions.

The tumor is rich in extracellular matrix (ECM) represented by type I collagen, fibronectin, tenasin, chondroitin sulfate and especially abundant hyaluronic acid. This excessive ECM production has been implicated in the invasive behavior of the tumor. However, the exact mechanism involved in the local invasiveness of the tumor have not been established yet.

In view of its rarity, a case of odontogenic myxoma of the maxilla infiltrating the maxillary antrum and adjacent tissues is herewith reported.

II. Case report

A 24 year old female patient reported with a chief complaint of painless, gradually progressive swelling in upper left side of face since one and half years. There was no history of trauma. Past medical and dental histories were non-contributory to the swelling.

Extra-oral examination revealed a diffuse, non-tender, firm swelling in the mid-face region approximately 5cm X 3cm in size, extending superoinferiorly 1.5 cm from the infraorbital ridge to 2 cm above the inferior border of the mandible and anteroposteriorly 1cm from the left corner of the mouth to 2.5 cm anterior to the tragus (figure 1). There was no local rise in temperature or change in color of the overlying skin. Lymph nodes were not palpable.

Intra oral examination showed ovoid swelling of about 5cm × 3 cm. extending anteroposteriorly from left maxillary second premolar till tuberosity of the same side. Buccal vestibule appeared obliterated. The swelling was firm, non-fluctuant and pale pink in colour with smooth surface. There was no bleeding or pus discharge from the lesion (figure 2).

Radiographically, it appeared to be a large osteolytic lesion involving the left posterior maxilla extending from the alveolus to the infraorbital margin and from nasal septum to the zygomatic arch. The lesion had a mixed density with calcified mass seen within the radiolucency. The margins were relatively well defined. 27 was displaced in the left maxillary sinus, whereas, 28 appeared to be floating in space. Displacement of adjacent teeth was also evident (figure 3).

Histopathological examination of an incisal biopsy revealed loosely arranged stellate, spindle-shaped, rounded and angular cells in an abundant loose myxoid stroma, relatively avascular containing few collagen fibrils. Inconspicuous island of odontogenic epithelium was also seen (figure 4). Stellate tumor cells showed anastomosing, long tapering, cytoplasmic processes (figure 5). On the basis of clinical, radiographical and histopathological features, diagnosis of “Odontogenic Myxoma” was made.
III. Discussion

Myxoma is Latin word originates from the Greek word ‘muxa’, which means ‘mucous’. There is a great controversy about the origin of myxomatous tumors. Bryant, in 1802, introduced the term myxosarcoma, which he described as a mucous transformation of round cell sarcoma, malignant and of large volume, usually attacking the omentum and skin. Furthermore, myxomas were described under the name of collenemas by Johannes Miller in 1838. In an article titled cellularpathologie, German pathologist Rudolph Virchow (1863) coined the term ‘myxoma’ for a group of tumors that had histologic resemblance to the mucinous substance of the umbilical cord. In 1947, Thoma and Goldman were the first to mention myxomas of the jaws.

The prevalence is principally quoted between 0.04% and 3.7%. On the basis of prospective study carried out in Tanzania, an annual incidence of 0.07 per million has been ascertained.

The classification of Odontogenic Myxoma as an odontogenic tumor has been justified by its (1) almost exclusive occurrence in the tooth-bearing areas of the jaws, (2) its frequent occurrence in young individuals, (3) its common association with an unerupted tooth or a developmentally absent tooth, (4) its histologic resemblance to the primitive mesenchymal portion of the developing tooth germ (dental follicle, dental papilla, periodontal ligament), (5) the occasional presence of sparse amounts odontogenic epithelium. Adekeye et al, however, expressed the view that the frequency and significance of these features may have been overstated. The authors supposed that the rarity of OMs in any extragnathic bone could be the only firm reason for suggesting an odontogenic origin.

Recently, Miyagi et al analyzed the expression of two proteins related to OM invasiveness (MMP-2 and hyaluronic acid) in human immature dental pulp stem cells and their results strongly suggested that dental pulp stem cells can be the precursors of OM.

OMs are slow-growing, painless, and site-aggressive tumors. Growth may be rapid and infiltration of neighboring soft tissues structures may occur. Since pain and hypoesthesia are not common, the lesions may reach a considerable size before patient perceives its existence and seeks treatment. Larger lesions may cause tooth displacement and cortical bone expansion, as seen in our patient.

When the maxillary sinus is involved, OMs often fills the antrum. In severe cases, nasal obstruction or exophthalmus may be the leading symptoms. Association with unerupted teeth has been reported in only 5% of OM cases reviewed. No specific predilection for any ethnic group has been reported.

The usual occurrence in 2nd to 4th decades of life and male to female ratio is 1:1.5, although cases in 3 month baby and 64 years old patients have also been reported. The occurrence of OM in the maxilla is rare when compared to the mandible (1:2). Farman et al. suggested that the mean age at the time of diagnosis of maxillary odontogenic myxomas in men was 29.2 years and in women was 35.3 years, while the mandibular odontogenic myxomas in men occur at a mean age of 25.8 years and in women at 29.3 years. Our case involved a 24 year old female with a large lesion of the left maxilla infiltrating the maxillary sinus of the same side.

Radiological diagnosis of odontogenic myxoma is difficult because of overlapping features with other benign and malignant bone lesions. Appearance of lesion may vary from a unilocular radiolucency to a multicystic lesion with well-defined or diffused margins with fine, bony trabeculae within its interior structure expressing a “honey combed,” “soap bubble,” or “tennis racket” appearance. A unilocular appearance may be seen more commonly in children and in anterior parts of the jaws.

Zhang et al in 2007 divided radiographical findings of OMs into six types: Type I-unilocular; Type II-multilocular (including honeycomb, soap bubble and tennis racquet patterns); Type III-involvement of local alveolar bone; Type IV-involvement of the maxillary sinus; Type V-osteolytic destruction and Type VI-a mix of osteolytic destruction and osteogenesis. A study done by Naffke CE, et al. on odontogenic myxoma revealed tennis racket appearance to be the most common radiographic finding.

Because of mixed radiopaque-radiolucent appearance, ascribed to the presence of foci of calcification, the diagnosis should be considered in mixed radiolucent-radiopaque lesions.

Macroscopically, OM has a whitish, translucent, mucinous appearance. It is un-encapsulated and poorly demarcated, thus permeating the surrounding bone and soft tissue by expansion rather than as a result of cellular growth.

Microscopically, odontogenic myxomas are composed of loosely arranged, evenly distributed spindle-shaped, rounded, and stellate cells with a lightly eosinophilic cytoplasm in a mucoid-rich (myxoid) intercellular matrix. Many stellate tumor cells have anastomosing, long tapering cytoplasmic processes. Cellular and nuclear polymorphism is rare as is mitotic activity. The tumor is relatively avascular or may exhibit delicate capillaries. Remnants of odontogenic epithelium may or may not present.

The histopathological examination in the present case showed presence of loose, spindle, rounded, stellate cells with long branching cytoplasmic processes in loose myxoid stroma.

OM has to be differentiated from other pathologies with similar histologic presentations like myxoid neurofibroma, myxoid liposarcoma, myxoid chondrosarcoma by enzymatic reactions (these being low in...
alkaline phosphatase and lactate dehydrogenase activity and high acid phosphatase, G6Pase dehydrogenase and isocitrate dehydrogenase activity while the reverse is true for odontogenic myxoma.24

Farman et al (1977) histochemical findings of ground substance of OM showed to consists of about 80% hyaluronic acid and 20% chondroitin sulphate.14,22 Garcia-Muñoz et al., stated that osromucoid protein is immunomodulatory and angiogenic and therefore results in invasive behavior of OM. It is anti-inflammatory, anti-neutrophil, anti-complement. Presence of osromucoid protein justifies classical mucoid appearance of the tumor.7

Odontogenic myxosarcoma, malignant variant of OM is exceedingly rare. It is a controversial issue, and several authors have addressed it. Many of these cases have had a malignant histologic appearance or exhibited an aggressive clinical course with death by local extension but without documented metastases. Lamberg MA et al reported findings on a maxillary myxoma in which histologic and cytogenetic findings supported a malignant interpretation.25

The tumor is not radiosensitive. Hence, surgical excision is the recommended treatment ranging from conservative curettage to radical excision. Owing to its potential for local infiltration, simple enucleation and curettage alone have been associated with a high recurrence rate of 10% to 33%. Thus, radical excision with burring of the cavity borders with a drill should be performed with maximum preservation of surrounding structures.26

IV. Conclusion

Myxoma is one of the least common lesions of maxillary bone, representing 3 to 8% of odontogenic tumors and cysts. The dental origin cannot be absolutely validated but is most probable as these tumors are almost exclusive to facial skeleton and universally absent in other bones of the skeleton. Our case underlines the difficulty in establishing a correct diagnosis, which requires interaction between radiologist, surgeon and pathologist.

References:

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Fig 1: Showing extra oral swelling

Fig 2: Showing intaroral swelling
Fig 3: PNS view showing maxillary sinus infiltration with displaced teeth

Fig 4: Photograph shows collection of stellate, spindle-shaped, rounded and angular fibroblasts cells in an abundant loose myxoid stroma, with inconspicuous strands of odontogenic epithelium in a myxoid stroma; H&E(x100).

Fig 5: Stellate tumor cells showing anastomosing, long tapering, cytoplasmic processes; H&E(x400).