Introduction

Odontogenic myxomas are intraosseous benign neoplasm most of them diagnosed in adults in the third decade of life [1]. Their frequency varies in different parts of the world between 3-20% of all odontogenic tumours. Two thirds occur in the mandible and one third in the maxilla, usually associated with a tooth germ. This tumour is uncommon in the paediatric population and exceptional in infants (under 2 years) with only 21 cases described in the literature [2].

Radiographically, this tumour is unilocular, with well-defined and corticated borders and most of them grow slowly. As the myxoma grows, it develops a multilocular pattern and clinically is evident a painless swelling [1]. The larger lesions have a soap-bubble and corticated borders and most of them grow slowly. As the conservative surgery (4.76%) [2].

Histologically, these lesions shows randomly orientated spindle or stellate-shaped cells set in a mucoid or myxoid matrix with a few delicate collagen fibers [1]. The neoplasm tends to permeate into the adjacent bony trabecula in a pseudo-malignant pattern.

This tendency of odontogenic myxomas to permeate into marrow spaces makes effective enucleation and curettage difficult. For small neoplasm, aggressive curettage may be adequate, but large lesions may require wide resection with free margins to prevent recurrences. Recurrence rates average about 25% and usually occur during the first 2 years after excision [1]. However, the literature review of infant odontogenic myxomas shows a low rate of recurrences after conservative surgery (4.76%) [2].

Case Report

Infant Odontogenic Myxoma: Case Report and Literature Review of a Specific Entity Recently Described

Abstract

Odontogenic myxomas are benign mesenchymal neoplasm most of them diagnosed in adults. They are uncommon in the paediatric population and exceptional in infants, with only 21 cases reported in the literature under the age of 2 years. We present a new case of infant odontogenic myxoma, that share the same clinical and radiological presentation with the cases described in the literature. They all presented with a painless paranasal swelling of short-term evolution, usually a few weeks duration (while in children or adults tumours usually develop slowly) and a well-defined, intraosseous, expansible lytic tumour of 3 cm average size in CT-scan examination. Most cases underwent enucleation and curettage with a very low rate of recurrences (4.76%). The aim of this article is to report a new case of this exceptional tumour, whose diagnosis was established at histologic examination. We focus on the importance to undergo a conservative approach in this infant population to minimize the surgical morbidity.

These tumours do not metastasize and although malignant transformation to myxosarcoma has been reported is a rare event [3].

Case Report

A healthy 21-month-old male, with no relevant personal or family medical history, presented to our hospital with a persistent swelling on the right side of his face after a minor trauma a few weeks before. The patient had persistent swelling that had no regressed. On physical examination there was an indurated, fixed 4cm mass in his right nasolabial groove, adjacent to the right anterior maxillary wall. The lesion did not enlarge with crying. The overlying skin was normal. Eye position and extra ocular motion were normal. An intraoral examination showed obliteration of the maxillary vestibule. Clinically the lesion looked like a mucocele.

In CT-scan examination there was a low-density lesion arising within the anterior medial aspect of the left maxillary bone with erosion of the maxillary sinus and the lateral nasal wall (Figure 1). There was separation of the mass from the nasal-lacrimal duct. Taking into account the history of trauma to the area, the CT scan concluded that the lesion was compatible with post-traumatic cyst.

Enucleation of the tumour was performed with curettage of the surrounding bone through a vestibular incision. The patient recovered uneventfully from surgery.

On gross pathology, the largest dimension of the lesion was 4cm and it was gelatinous with pale brown colour. Histology revealed a myxoid tumour with haphazardly arranged stellate to spindle-shaped cells in a mucoid-rich intercellular matrix (Figure 2). The cells had an eosinophilic cytoplasm, small round hyperchromatic nuclei and fine chromatin. There was not cellular pleomorphism or nuclear atypia. Immunohistochemical staining was positive for vimentin and...
International literature [2]. All these patients, as well as our case, share
to our knowledge, 21 cases have been described in infants in the
unit or multilocular radiolucency with well-defined margins [1].
Expansion and radiographically, the lesion frequently appears as a
of young adults. They usually present as a slow, painless bony
growing tumours that occur most commonly in the mandible
lytic tumour developed from the maxilla. 71.43 % underwent
weeks evolution showing in the TC-scan a unilocular, homogeneous
mass include a wide variety of benign and malignant tumours:
immediate follow-up period of 3.068 years.
The histological appearance of our case is similar to those
described in the literature. These lesions resemble the mesenchymal
portion of a developing tooth. Macroscopically they are gelatinous
masses with well-defined borders but a true capsule is absent.
Microscopic examinations show a myxoid background with randomly
oriented stellate-shaped fibroblastic cells and a few delicate collagen
fibres. Some myxomas may show areas with a little more collagen
production and they are termed fibromyxomas or myxofibromas.
Binucleated cells, mild pleomorphism and mitotic figures may occur,
but these lesions do not behave differently. Immunohistochemically
tumoral cells usually react with antibodies to vimentin and muscle-
specific actin [1].

Myxomas are commonly associated with unerupted teeth and
probably arise from the mesenchymal portion of the tooth germ. The
histology of immature dental tissues (dental papillae and follicles)
and dental pulp is very similar to myxomas, so these structures are
the most commonly mistaken histopathologically for this tumor [4].
Dental papilla is composed of stellate and fusiform cells set in a myxoid
matrix with delicate collagen fibres. However, this tissue is always
lined, at least focally, by a rim of odontoblasts. A dental follicle can
also show a myxoid appearance. This tissue is lined along one margin
by reduced enamel epithelium (Figure 3). This features, together with
a clinical-radiologic correlation, help to distinguish these normal
dental tissues from odontogenic myxoma. Others central entities
that enters into the histological differential diagnosis are tumours
with a myxoid background: myxoid neurofibroma, chondromyxoid
fibroma, myxoid chondrosarcoma and the myxoid variant of desmoid
fibromatosis. Myxoid neurofibroma presents numerous mast cells,
a positive S-100 immunohistochemical reaction, and zones with
organization of the collagen and lesional cells into broad fascicles.
Chondromyxoid fibroma and myxoid chondrosarcoma should show
some area with chondroid differentiation and, in the latter one,
cellular atypia. The myxoid variant of desmoid fibromatosis presents
focal areas with dense collagen bundles. Cranial fasciitis can also have
a prominent myxoid background, but this is a extraosseous lesion and
express smooth muscle actin. Three cases described in the literature
were initially misdiagnosed histopathology as nodular fasciitis [2,5,6].

Treatment of this lesion consists of surgical resection, but the
extent of the resection is controversial. Some authors support a
radical surgery with wide clear margins [7,8]. However, there may
be the doubt whether to perform a wide surgical excision in children
or not, given this is a benign disease. So others authors suggest a
conservative excision or curettage [9,10]. In the series of infant
patients treated with conservative excision there is no clinical or
radiological evidence of recurrence over a median follow-up period of
3 years. These results suggest that in the infants conservative surgical
treatment is an effective therapy. The tumour is neither radiosensitive
nor chemosensitive.

As the neoplasm is located in or near the maxilla, initial clinical
misdiagnosis is not rare. Clinically differential diagnosis of a midline
mass include a wide variety of benign and malignant tumours:
cephalocoele, dermoid, glioma, hamartoma, lymphangioma,
rhabdomyosarcoma, neurofibroma and nasolacrimal duct cyst. In our case, the diagnosis of myxoma was not initially suggested clinically or radiologically and a diagnosis of mucocele and post-traumatic cyst were suggested.

This case highlights the importance of including odontogenic myxoma in the differential diagnosis of lesions in the maxilla of infants. The unusual clinical behaviour of this tumour as a rapidly expansile lesion and the different entities that simule odontogenic myxoma underline the difficulty in making a correct diagnosis, that requires interaction among pathologist, radiologist and clinician. As we have already discussed, review of the literature suggests that a conservative surgery should be the initial surgical approach in infants.

References