The central odontogenic fibroma: How difficult can be making a preliminary diagnosis

Roberto Pippi 1, Marcello Santoro 1, Romeo Patini 2

1 Department of Oral and Maxillofacial Sciences, “Sapienza” - University of Rome, Caserta street, 6 - 00161 Rome, Italy
2 Department of Surgical sciences for head and neck diseases, School of dentistry, Catholic University of Sacred Heart, Largo A. Gemelli, 1 - 00168 Rome, Italy

Correspondence: Department of Surgical sciences for head and neck diseases School of dentistry, Catholic University of Sacred Heart Largo A. Gemelli, 1 - 00168 Rome, Italy romeopatini@hotmail.it

Received: 05/10/2015 Accepted: 09/12/2015

Abstract
Central odontogenic fibroma (COF) is a rare benign odontogenic tumor derived from the dental ectomesenchymal tissues. A 16-year-old Caucasian female patient was referred by her dentist for a radiolucent asymptomatic area associated with the crown of the impacted lower right third molar. A preliminary diagnosis of a follicular cyst was supposed. The lesion was surgically removed under general anesthesia together with the impacted tooth. The microscopic diagnosis of the excised tissue revealed an odontogenic fibroma. No clinical or radiographic signs of recurrence were found five years after surgical excision. Despite the various differential diagnoses of homogeneous unilocular and well delimited radiolucencies of the jaws, enucleation with peripheral curettage, without any other pre-operative imaging exams or biopsies, can be considered as the treatment of choice.

Key words: Differential diagnosis, impacted third molar, radiographic imaging, microscopic diagnosis, odontogenic fibroma.

Introduction
The central odontogenic fibroma (COF) is an uncommon odontogenic tumor whose typing is controversial both clinically and histologically (1). COF is reported to occur in the 4-80 year range of age (mean age: 40 years) with a 2.2:1 female predilection. It accounts for 0.1-1.5% of all odontogenic tumors and 6.1% if odontoma is excluded. Approximately 55% of the cases occur in the mandible, half of which posterior to the first molar and up to one-third in conjunction with an un-erupted third molar (2). From a topographic point of view two different kinds of COF exist, an intra-osseous or central form and an extra-osseous or peripheral form (3). A recent report demonstrated that age distribution among central lesions showed a shallow curve, with all decades represented whereas the peripherally located lesions showed a predilection for the 2nd to 4th decades of life and that intraosseous tumors were relatively evenly distributed in the anterior, premolar and molar regions whilst peripheral lesions tended to arise in the anterior sextants of the jaws (4).

In 2005, WHO (5) defined two different histological types of COF, the simplex type, with only few epithelial isles, and the complex type, rich of epithelial cells. Be-
cause the presence of epithelium is a requisite to con-
firm the diagnosis, immunohistochemistry has been de-
monstrated to be helpful in epithelium-poor cases (6).
The clinical and radiographic diagnosis of COF has been
reported not to be always easy and the tumor should be
differentiated from many other lesions of the jaws such
as keratocystic odontogenic tumor, ameloblastoma,
odontogenic myxoma, ameloblastic fibroma, calcifying
odontogenic cyst, dentigerous cyst since it could appear
as a well-defined radio-transparent area associated with
the crown of an impacted tooth.
The aim of this case report is to describe a case of COF
in which the preliminary diagnosis was particularly diffi-
cult because the lesion mimicked a dentigerous cyst.

Case Report
A 16-year-old woman was referred by her dentist to
the Oral Surgery Unit – Department of Oral and Maxi-
llofacial Sciences, Faculty of Medicine and Dentistry,
“Sapienza” University of Rome for a radiolucent area
associated with the impacted right third molar whose
radiographic characteristics mimicked those of a
dentigerous cyst. The intraoral examination revealed
no alterations, the gingival surface distal to the lower
right second molar was healthy and smooth. The ortho-
pantomography showed a radiolucent lesion located in
the right mandibular angle containing an impacted right
lower third molar (Fig. 1). A preliminary diagnosis of a
dentigerous cyst was therefore supposed.
The surgery was conducted under general anesthesia.
After a muco-periosteal flap was incised and raised, the
ostectomy was performed to reach the pathologic tissue
which was not adherent to the impacted tooth and there-
fore easily separated from it. Due to the fibrous aspect of
the lesion, an odontogenic fibroma was considered as a
probable diagnosis so that a thorough 1-2 mm curettage
of the residual bone cavity was therefore performed after
the complete excision of the pathological tissue and the
impacted tooth removal.
The excised lesion was constituted by multiple fragments
not exceeding 2.5 cm in diameter. All fragments were
fixed with 10% buffered formalin and microscopically
analyzed. Microscopic examination showed a cellular
loose connective tissue with epithelial cell aggregates
showing the typical morphological characteristics of
the enamel organ. These aspects were compatible with
the diagnosis of the simple type of odontogenic fibroma
(Fig. 2). No recurrence was seen at the 5 years clinical
and radiographic follow-up (Fig. 3).

Discussion
The central odontogenic fibroma is an uncommon tu-
mor which clinically appears as an asymptomatic well-
deefined osteolytic lesion and which rarely can be loca-
llly aggressive, with dental displacement and rhizolysis.
Radiologically, COF appears as an uni- or multi-locular
radiolucent area and it can be indistinguishable from
other radio-transparent lesions making the pre-operative
diagnosis more difficult. Actually, in the present case a
dentigerous cyst was suspected from the two-dimensio-
nal x-ray performed.
Nevertheless the use of 3D x-ray exams, such as tradi-
tional CT or Cone Beam CT and high resolution nuclear
magnetic resonance (HRNMR), could be considered to
better investigate all jaw radiolucencies and correctly
plan the surgery (7,8), thus resulting in a lower risk of
contiguous anatomic structure surgical injury.
A correct pre-operative evaluation deeply influences the surgical approach in that a previous biopsy is highly indicated for tumors but not for cysts. If a benign tumor is diagnosed, a more extensive ostectomy and a more accurate curettage of the residual bone cavity should be performed in order to completely excise the lesion and to avoid recurrences (2). In this light, HRMR or contrast-CT are nowadays indicated for pre-operative evaluation of multi-locular or/and ill-defined radiolucent lesions while direct enucleation with peripheral curettage, without any pre-operative imaging exams or biopsies, can be considered for uni-locular homogeneous radiotransparent and well-delimited lesions.

Due to the very low incidence of recurrence and the benign biological behavior of the COF, the surgical approach is usually conservative but, if a microscopic diagnosis has not been performed from a pre-operative biopsy, a more aggressive lesion, such as a keratocystic tumor or a unicystic ameloblastoma, that have a higher risk of recurrence in relation to their different pattern of local aggressiveness resulting in a longer follow-up required, cannot be excluded from the definitive diagnosis (2).

Conclusions
The great variability in radiological appearance of the COF means that it should be considered in the differential diagnosis of all jaw radioluencies. The case presented here shows how difficult can be making a preliminary diagnosis of COF although the intra-operative appearance of the lesion is suggestive and the prognosis is anyhow good, provided that a peripheral 1-2 mm curettage is performed.

References

Conflict of Interest
The authors received no financial support and declare no potential conflicts of interest.