Ameloblastic Fibroma

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History

An 18-year-old male presented for a routine dental examination with an incidental finding in the left posterior maxilla.

Radiographic Features

Imaging studies revealed a well-circumscribed, unilocular radiolucency preventing eruption of the left maxillary second molar and displacing it in a superior and posterior direction, into the maxillary sinus (Fig. 1).

Treatment

The area was treated with thorough curettage and the associated maxillary second molar was extracted.

Discussion

Ameloblastic fibromas are neoplasms of odontogenic epithelium and mesenchymal tissues, and as such are categorized as mixed odontogenic tumors. Other mixed odontogenic lesions, such as ameloblastic fibro-odontomas and odontomas share some clinical, radiographic and histologic similarities with ameloblastic fibroma. In the past, it was suggested that these lesions represented a spectrum of a single entity, with ameloblastic fibromas, the least differentiated of the tumors, maturing and developing into ameloblastic fibro-odontomas and later odontomas [1]. This theory has been refuted with the support of the tumors’ associated demographics. The least differentiated lesion, ameloblastic fibroma, actually occurs, on average, at an older age then the more differentiated ameloblastic fibro-odontoma and odontoma [4]. Further more, ameloblastic fibro-odontomas and odontomas are better categorized as hamartomas and as a result, unlike the ameloblastic fibroma, have little chance of recurrence or malignant transformation [4]. For these reasons, despite many similarities, it is essential to differentiate the ameloblastic fibroma from other mixed odontogenic lesions because it has true neoplastic qualities.
Ameloblastic fibromas are rare and comprise approximately 2% of odontogenic tumors [2, 3]. The tumors are considered a tumor of childhood and adolescence and occur almost exclusively in the first and second decades of life [3, 4]. A slight male predilection has been noted [4, 5]. The most common location for the tumor is the posterior mandible, followed by the posterior maxilla. Patients often present with painless swelling of the jaw and the lesion may affect the normal eruption of teeth in the area. An impacted tooth may be associated with the tumor in approximately three quarters of the cases [2, 5, 6]. Some lesions are asymptomatic, with up to 20% of cases initially detected upon review of routine dental radiographs [1, 3, 4, 6].

Radiographically, ameloblastic fibromas are unilocular lesions, occasionally multilocular when larger, with smooth well-demarcated borders. Cortical expansion may or may not be discernable on plane film. Because lesions are frequently associated with unerupted teeth they may initially be interpreted as dentigerous cysts [1, 3, 4, 7].

Grossly, ameloblastic fibroma appears as firm, lobular soft tissue mass with a smooth surface [3]. If a tooth is associated with the lesion it may accompany the specimen. A capsule is generally not appreciated. Microscopically, an ameloblastic fibroma is composed of a connective tissue background that appears to recapitulate dental papilla, resembling stellate reticulum [1, 3, 7]. This tissue is composed of spindled and angular cells with little collagen, imparting a myxomatous appearance. The epithelial component is made up of thin branching cords or small nests of odontogenic epithelium with little cytoplasm and basophilic nuclei. Larger nests may show a central area of stellate reticulum. Mitoses should not be a feature of ameloblastic fibroma [1, 3]. The presence of mitosis should expand the differential diagnosis to include malignant entities, to include ameloblastic fibrosarcoma. Finally, immunohistochemistry generally does not aid in differentiating ameloblastic fibroma from other mixed odontogenic tumors.

Surgical excision or thorough curettage with removal of affected teeth is the treatment of choice [1, 2]. The recurrence rate varies among sources, but is considered to be low [5, 7]. While uncommon, the possibility of malignant transformation of ameloblastic fibroma into ameloblastic fibrosarcoma is well documented [8, 9].

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