Adenomatoid odontogenic tumor associated with odontoma: a case report and critical review of the literature

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Abstract
We describe a case of adenomatoid odontogenic tumor (AOT) associated with odontoma occurring in the posterior mandible of a 32-year-old man. Although calcifications are commonly found in the AOT, the presence of rudimentary dental structures is a very rare phenomenon. Cases with similar aspects have been described as ameloblastic dentinoma, ameloblastic odontoma, adenoameloblastic odontoma and AOT associated with odontoma. After a careful analysis of the literature we describe the clinical aspects of this tumor. Further case reports and surveys of odontogenic tumors are necessary to define whether AOT associated with odontoma is a variant of AOT or a distinct clinicopathologic condition.

Odontogenic tumors and hamartomas encompass a large variety of rare lesions that originate from odontogenic tissue and present with variable levels of differentiation. Determination of their precise nature (i.e., a hamartoma or a neoplasm) is difficult and sometimes inconclusive. This, in turn, makes it difficult to devise a nomenclature for this group of lesions.

In 1998, Allen and co-workers [1] described a rare odontogenic lesion showing a distinct proliferative component of epithelial cells arranged in cords, strands, swirls, and duct-like structures associated with a prominent dentin formation lined by ameloblast-like cells [1]. The prominence of dentin and adenomatoid structures in this lesion led the authors to describe it as an “adenomatoid dentinoma”. This lesion showed overlapping features of an adenomatoid odontogenic tumor (AOT) and an odontoma, with histological features similar to those reported by Dunlap et al., [2] and was described as an adenoameloblastic odontoma. Subsequently, different terms were used to describe similar lesions and, up to now, there is no consensus on to the nature, histopathological spectrum, and clinical significance of this lesion. Here, we report a similar case that exemplifies the odontogenic potential of an AOT and we present a critical analysis of the literature regarding this diagnosis.

Case presentation
A 32-year-old man presented with an asymptomatic, well-delimitated, 2.6 x 2.1 x 1.3 cm unilocular hypodense tumor in the left posterior mandible (Figure 1). On examination of the oral cavity, a mild expansion of the retromolar trigone was observed. An incisional biopsy was performed, revealing a cystic capsule lined with stratified squamous, non-keratinized epithelium. The lesion was completely enucleated and was found to contain a cystic capsule adhering to a thin hard tissue resembling a tooth. Microscopic observation showed a cystic cavity lined by flattened, stratified non-keratinized squamous epithelium. These findings suggested a benign odontogenic cystic lesion. The lesion was completely enucleated and was found to contain a cystic capsule adhering to a thin hard tissue resembling a tooth. Microscopic observation showed a cystic cavity lined by flattened, stratified non-keratinized squamous epithelium. In some areas, we observed nodules, cords and strands in the epithelial lining forming swirls of fusiform cells and ribbons of ameloblast-like cells (Figure 2). We also observed numerous islands of odontogenic epithelium in the fibrous capsule and solid areas formed mainly by dentin with lacunar borders externally lined by ameloblast-like cells (Figure 2). Some areas had a thin layer with an appearance similar to that of an enamel matrix deposit (Figure 3C). The diagnosis

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was of AOT associated with odontoma. No recurrence was found upon an 8 months follow-up after enucleation of the lesion.

Discussion
The present report describes a cystic lesion characterized histologically by the presence of an AOT associated with an odontoma. Previous reports of lesions with AOT-like epithelial structures and rudimentary dental formation have variously described them as adenoameloblastic odontomas [2], cystic complex compound odontomas [3], adenomatoid dentinomas [1,4], AOT associated with odontoma [5], and AOT [6]. One can conclude from the variable nomenclature used that no consensus has been reached regarding the precise nature and definition of histological characteristics of such lesions.

After extensive review of the literature on lesions with similar histological aspects, we retrieved those case reports showing overlapping features of AOT and odontoma or dental tissue presenting with rudimentary

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**Figure 1** Computerized tomography scan in sagittal (A and B) and axial (C) planes showing unilocular well defined hypodense mandibular tumor.

**Figure 2** Histological aspects of the epithelial component of the tumor. Cystic cavity lined by flattened, stratified non-keratinized squamous epithelium (A). Epithelial cords and strands forming swirls and nodules of fusiform cells and ribbons of ameloblast-like cells (B, C and D) (hematoxylin-eosin, original magnification X 200 and X 400).
odontogenesis described under the following nomenclature: ameloblastic dentinoma, adenoameloblastic odontoma, AOT associated with odontoma and adenomatoid odontoma [1-7]. Only cases with published histological pictures of these findings were included. The clinical and radiographic features together with the clinical behavior of the AOT–odontoma variants reported in the literature are listed Table 1. This variant of AOT affects patients aged 4 to 46 years (mean 24.4 years). Five of the 11 patients were younger than 18 years. Instead of being located in the anterior maxilla, which is more typical of AOT [8], this odontoma-associated variant was mainly present in the posterior mandible (9 of 11 cases) and was located in the maxilla in only one case. Radiographically, the tumor typically results in a well-defined radiolucent image, sometimes exhibiting subtle calcifications or radiopaque deposits within the lesion [1,4]. A radiopaque image was noted in 1 case [6].

The epithelial component of AOT associated with odontoma exhibits nodules of cuboidal and fusiform cells forming nests (condensations) or rosette-like structures and a variable number of duct-like formations lined by cuboidal or columnar cells. Interconnecting strands and ribbons with two or more cells are present throughout the lesion. In our case we did not find duct-like structures typically reported in AOT. However, due to the overall morphologic aspect of the tumor, this diagnosis can be made without the presence of these structures [9]. The presence of tubular dentin, with or without a lining of ameloblast-like cells, and an amorphous and/or reticulated enamel matrix complete the histological features of the lesion. These characteristics are essentially identical to those of lesions previously identified as AOT, in which histological elements of odontogenesis were also described. Until a better definition is established in the literature, our opinion is that this group should be defined as AOT associated with odontoma.

AOT associated with odontoma should not be confused with adenomatoid odontogenic hamartoma (AOH). The AOH is characterized by the deposition of dentin in a ring-like pattern with odontoblasts, dental papilla, lined on the outer surface by enamel, ameloblasts and stellate reticulum-like cells [10,11]. Small duct-like structures are seen on the dental organ-like area and they are lined by flattened or cuboidal cells [10]. This finding is in contrast to AOT, in which ducts are lined by columnar ameloblast-like cells. In addition, no rosette-like structures are identified in AOH. It appears that these formations represent a morphological variation similar to that of the external epithelium of the dental organ. On the basis of this observation and in agreement with the opinion expressed by Kemp et al. [4] we believe that AOH simply represents a failed attempt at tooth development, more specifically the third molar, and could be classified as hamartomatous in nature. Takeda et al. [12] showed similar morphological variations in the odontogenic epithelium of compound odontomas, which are different from those present in AOT.
| Authors                  | Nomenclature                              | Gender | Age | Site                   | Radiographic features                                           | Size (cm)       | Treatment                        | Follow-up (Recurrence) |
|-------------------------|-------------------------------------------|--------|-----|------------------------|-----------------------------------------------------------------|-----------------|----------------------------------|------------------------|
| Miles [3]               | Cystic complex composite odontoma         | M      | 18  | Posterior mandible     | Well-circumscribed radiolucent lesion-like dentigerous cyst     | NA              | Enucleation                      | NA                     |
| Dunlap & Fritzlen [2]   | Adenoameloblastic odontoma                | F      | 4   | Posterior mandible     | Well-defined radiolucent lesion                                | NA              | Enucleation                      | NA                     |
| Tajima et al. [6]       | AOT arising in an odontogenic cyst        | M      | 15  | Maxillary sinus        | Well-defined radiopaque mass                                    | 4.0 × 4.0       | Enucleation                      | 5 years (No)           |
| Allen et al. [1]        | Adenomatoid dentinoma                     | M      | 37  | Mandibular third molar region | Unilocular radiolucent                                          | 1.5 × 2         | Enucleation                      | 2 years (No)           |
|                         | Adenomatoid dentinoma                     | M      | 29  | Mandibular third molar region | Well-circumscribed radioluency with subtle linear radiopacity within the lesion | 0.75 × 0.75     | Enucleation                      | 8 years (No)           |
|                         | Adenomatoid dentinoma                     | M      | 35  | Left posterior mandible | Unilocular radiolucent                                          | 1 × 0.8         | Curettage                        | 3 years (No)           |
|                         | Adenomatoid dentinoma                     | F      | 29  | Mandibular third molar region | Radioluency with subtle focal calcifications                     | 2.0 × 2.75      | Surgery, not specified           | 8 years (No)           |
| Cudney et al. [7]       | AOT associated with odontoma              | M      | 13  | Mandibular left canine | Pericoronal radioluency with faint snowflake calcifications     | 1.4 × 0.7       | Excisional biopsy                | 1 year (No) 1          |
| Kemp et al. [4]         | Adenomatoid dentinoma                     | M      | 46  | Right mandibular body and ramus | Well-circumscribed with focal internal radiopacity              | 4.0 × 2.5       | Enucleation and curettage        | 6 months (No)          |
| Martínez et al. [5]     | AOT concomitant with cystic complex odontoma | F    | 10  | Left posterior mandible | Well-circumscribed radioluency with radiopaque mass inside     | 3.0 × 1.5       | Enucleation with posterior curettage | NA                    |
| Gomez et al. (present report) | AOT associated with odontoma               | M      | 33  | Left mandibular third molar region | Well-delimited unilocular radioluency                           | 3.0 × 2.0       | Enucleation                      | 8 months (No)          |

NA: not available.
In the case we report, we found a cystic cavity lined by stratified squamous, non-keratinized epithelium. The same microscopic appearance was reported by Tajima et al. [6], indicating that odontogenic cysts should be included in the differential diagnosis of this tumor.

Although some AOT with odontoma are large lesions [4,6], their clinical behavior seems similar to conventional AOT. None of the cases showed recurrence, suggesting that the tumor can be treated conservatively.

Conclusion
In the present report we review the literature and describe the clinical and histological aspects of AOT associated with odontoma. Although this tumor differs in some aspects to classic AOT, further case reports and surveys of odontogenic tumors are necessary to define whether it is a variant of this odontogenic tumor or a distinct clinicopathological condition.

Consent
The authors declare that the patient had given consent for the case report to be published.

Competing interest
The authors declare that they have no competing interests.

Authors’ contribution
WHC performed surgery and drafted the manuscript. AML, RSG and CCG performed histopathological examination, conducted a review and analysis of the literature, and drafted the manuscript. All authors read and approved the final manuscript.

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