Diagnostic Imaging of Recurred Mandibular Ameloblastoma with Large Soft Tissue Involvement

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ABSTRACT

An uncommon case of a patient with recurrent mandibular ameloblastoma involving various adjacent soft tissues is presented with plain film radiography, computed tomography and magnetic resonance imaging (MRI) and bone scan. The tumor involved mandible and eroded several bones including foramen ovale. This case shows that although an ameloblastoma primarily affect mandible, exact radiographic evaluation of tumor extent will assist the surgeon in the progress of rational approaches to the management of ameloblastoma. (Korean J Oral Maxillofac Radiol 2000; 30: 281-285)

KEY WORDS: ameloblastoma, recurrence, mandible, skull base

Ameloblastoma is the second most frequent odontogenic tumor and represents about 1% of all tumors and cysts of the jaws. It is usually benign but invades locally and has high recurrence rate and occasionally metastasizes. Histopathologically, it is divided into 4 categories-follicular, plexiform, acanthomatous and granular cell type. It is most common in the third and fourth decade and shows no sexual predilection and occurs predominantly in the mandible.

In plain film radiography, ameloblastoma in the mandible shows a multilocular (so called “soap-bubble” or “honeycomb”) expansile radiolucency or a unilocular radiolucency. It frequently shows bucco-lingual cortical expansion especially in axial CT scan. Ameloblastoma in the maxilla can show hazi-ness and mucosal thickening in the maxillary sinus, and also show multilocular or unilocular radiolucency. CT and MRI are essential for diagnosis and treatment planning of ameloblastoma, because they can show tumor extension, border invasion, shell-like bulging of the bony cortex, papillary projection and so on.

The treatment of ameloblastoma includes conservative or radical resection. In case of multicyclic type, radical resection is preferably performed and the tumor is removed with a safety margin of at least 1 cm in healthy tissue. The recurrence rate of ameloblastoma has been reported variously by numerous authors. Some authors have found the overall recurrence rate to be 55-90%. However, the incidence of recurrence following radical resection including bone and soft tissues has been reported to be 5-15%.

We present a case of recurrent mandibular ameloblastoma which extensively involves adjacent soft tissues and is suspicious of extension into the skull base.

Case report

A 40-year-old woman visited Seoul National University Dental Hospital (SNUDH) for discomfort of right bucal mucosa. The history revealed that she had undergone surgery in 1982 for an ameloblastoma of right mandibular body. The procedure included partial mandibulectomy with bone grafting. Three years later, she underwent an operation of graft removal because of infection on graft site. On her visit-14 years after graft removal, she had a right facial swelling, firm mass palpated on right cheek, hyperplastic tissue on right bucal mucosa, and redness on lower right molar area.

In the panoramic and occlusal radiography, honeycomb and soap-bubble appearances of mandible were seen from lateral side alveolar bone of #42 tooth to resected border of right mandibular body, the bony cortex showed thinning and expansion and cystic radiolucent lesion was also seen on mandibular body (Fig. 1). Lateral wall of the right maxillary sinus was depressed and alveolar bone of the maxillary tuberosity was eroded.

The CT scan showed the mass primarily involving right
masticator space with extension to upper masticator space, atrophy of upper portion of right temporal muscle, depression and erosion of lateral wall of right maxillary sinus and right sphenoid sinus. The mass was heterogenously enhanced and the lesion similar to described above was seen in the right mandibular body (Fig. 2).

Fig. 1. Panoramic (A) and Occlusal radiograph (B). Honey-comb and soap-bubble appearances of mandible are seen from lateral side alveolar bone of #42 tooth to resected border of right mandibular body, the bony cortex showed thinning and expansion and cystic radiolucent lesion was also seen on mandibular body.

Fig. 2. Axial (A) and Coronal (B) Enhanced CT scan. The mass involving right masticator space with extension to upper masticator space, and causing atrophy of upper portion of right temporal muscle, depression and erosion of lateral wall of right maxillary sinus, and right sphenoid sinus. The mass is heterogenously enhanced and the lesion similar to described above was seen in the right mandibular body.
The $^{99m}$Tc bone scan in March 1999, showed collection of radioisotope in right mandible but no evidence of metastasis to distant area (Fig. 3).

In the MRI, the mass with solid and cystic nature and lobulating contour in the right masticator space was observed, and it was extended to lower part through pterygoid muscle displacing parapharyngeal fat plane and parapharyngeal mucosal space (Fig. 4). In the enhanced coronal view, widening of foramen ovale was suspicious and the area showing similar signal intensity of the main mass was seen in the right cavernous sinus (Fig. 5). In the neck sonography, 2 cm sized round enlarged lymph node was seen in the right submandibular area (Fig. 6).

Resection operation of partial mandibulectomy and partial
maxillectomy with supraomohyoid neck dissection under general anesthesia was done. Neither cavernous sinus extension nor lymph node metastasis was identified, and biopsy specimens showed no evidence of high mitosis incidence and malignant transformation at any slice. We concluded it was a recurred ameloblastoma of plexiform type mixed with granular cell type unexpectedly extended to surrounding area.

**Discussion**

Ameloblastoma occurs mainly in the mandible but rarely in the maxilla. If once occurred in the maxilla, about a half of all maxillary ameloblastomas develop in the posterior part of the maxilla. They have the potential to break through the maxilla into the adjacent vital structures such as the infratemporal fossa, the pterygomaxillary space, the masticator space and the orbit. However, mandibular ameloblastomas rarely show extensive growth into the other vital structures unlike maxillary ones. Therefore, this case is unique because of the large soft tissue involvement although recurrence.

In this case, panoramic view was helpful in diagnosing of this huge mass by showing the pathognomonic honey-comb appearance. And CT and MR imaging showed the extent of the mass. Cause of recurrence is mainly that adequate radical therapy was not performed. When a surgery is done, 1 cm safety margin must be confirmed, so it is very useful to verify the border of mass with imaging modalities such as CT or MR imaging. Though contrast-enhanced CT scan can visualize more easily tumor tissue breaking through thin cortical boundaries than MR imaging, decreased attenuation due to fibrosis and edema may make delineation of the tumor-normal tissue interface more difficult in case that electron density of abnormal tissue approximates that of the normal surrounding tissue. MR imaging can more clearly show interfaces between mass and surrounding tissues. In the primary ameloblastoma outlined with bone tissue, CT scan may have better interpretation capabilities, but in the recurred maxillary ameloblastomas or the recurred mandibular ameloblastomas such as this case involving large soft tissue areas, MR imaging may have better interpretation capabilities because this type of tumor was developed from resected bony surface or soft tissues near resection margin.

In pre-operation MR imaging, involvement of skull base was doubted due to the widening of right foramen ovale and the high signal intensity of carvenous sinus. The widening of the foramen ovale may be caused to slight erosion of the skull base due to the mass, however, because the carvenous sinus originally have the high signal intensity in the MRI, it could not mean the extension of the mass. The mass adjacent to skull base was easily separated and no carvenous sinus extension was identified on the operation.

Recurrent ameloblastoma may not be always outlined by bony tissues and may seem to be originated from soft tissues adjacent to resected jaw bone and may involve extremely large area such as this case. Almost of all reported cases were maxillary ones, however, mandibular ones were rarely reported because ameloblastoma occurred on mandible could be easily resected with safety margin compared with ameloblastoma on maxilla presenting complex interface with adjacent structures. In this respect, MR imaging is essential to evaluate the extent of tumors. In our case, the tumor portion which was not completely resected and impacted to the adjacent soft tissues seem to originate a recurrence. This was revealed by same histopathologic result from mandible and various adja-
cent soft tissues.

This case of recurrent mandibular ameloblastoma involved adjacent soft tissues and erosion of the maxilla and skull base is uncommon. Although an ameloblastoma primarily affect mandible, exact radiographic evaluation of tumor extent to the adjacent soft tissue will assist the surgeon in the progress of rational approaches to the management of ameloblastoma.

References


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— 285 —