Case Report

Clear cell odontogenic carcinoma in maxilla: A rare case report

Swapna Amod Patankar1, Pratik Raut2, Amod Pramod Patankar3, Rajesh Ashok Kshirsagar4
Departments of 1Oral and Maxillofacial Pathology and Oral Microbiology and 2Oral and Maxillofacial Surgery, Bharati Vidyapeeth Dental College and Hospital, 3Consultant Maxillofacial Surgeon in Pune, Pune, Maharashtra, India

Abstract
A rare and aggressive form of tumor with a female predilection and more seen with mandible is the clear cell odontogenic carcinoma (CCOC). The World Health Organization categorized it as a malignant neoplasm in 2005. This type of tumor is characterized by local recurrences as well as distant metastasis apart from being highly aggressive in nature. Due to the presence of clear cells, it is a diagnostic challenge to the pathologist. Only about 84 cases of maxillary variant have been reported in the literature. Here, we present a case report of a 31-year-old male with CCOC in the maxilla from the pathological and diagnostic point of view.

Keywords: Clear cell odontogenic tumor, clear cells, male, maxilla, rare

INTRODUCTION

A rare and aggressive form of tumor with a female predilection and more seen with mandible is the clear cell odontogenic carcinoma (CCOC). The World Health Organization categorized it as a malignant neoplasm in 2005. This type of tumor is characterized by local recurrences as well as distant metastasis apart from being highly aggressive in nature. Due to the presence of clear cells, it is a diagnostic challenge to the pathologist. About 84 cases of maxillary variant have been reported so far, and because of its rarity, the diagnosis can be missed easily.

CASE HISTORY

A 31-year-old male reported with the chief complaint of swelling on the upper left anterolateral region of the mouth. The swelling grew gradually in size over a span of 6 months. It was initially painless but later become painful. The pain was more on the outer aspect of the upper left part of the cheek. Eventually, when the patient had reported to our department, he was having unbearable pain in the nose and surrounding left eye region [Figure 1].

External examination of the swelling showed a solitary swelling measuring 3 cm × 4 cm in size that extended from the upper left part of the cheek, extending in nasal area, the zygomatic arch and lower border of the left eye. It had an irregular surface, well-demarcated border and covered by intact skin. It was firm consistency and was nontender without any signs of bleeding or pus discharge. There was no local rise of temperature over the skin covering the swelling. No abnormality was noted with the temporomandibular joint. The eyesight was not compromised, though the area around the eye was painful.

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Address for correspondence: Dr. Swapna Amod Patankar, “Shree” 35/1 Padma Darshan Housing Society Survey Number 62B, Parvati, Pune - 411 009, Maharashtra, India.
E-mail: swapna.patankar@gmail.com
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Left submental and submandibular lymph nodes along with cervical lymphadenopathy (of level 1, 2, 3 and 4) were observed in the patient. There was no relevant medical or dental history in the recent years associated with the patient.

Intraoral features of the swelling: the swelling extended from right maxillary first premolar to the first molar. The swelling was characterized by a normal mucosal covering, nonmovable/fixed, nontender, no pus or blood discharge, no ulceration of the overlying mucosa. The swelling had an irregular surface texture and well-defined borders.

Investigations
The following investigations were carried out. These include radiographs (cone-beam computer tomography), complete hemogram, incisional biopsy and abdominal ultrasonography (USG). Abdominal USG was carried out to rule out any primary or secondary tumor. USG reports were normal ruling out any metastasis to or from any other soft-tissue tumor.

Radiographic features
It showed both buccal as well as a lingual cortical plate expansion. A well-defined, corticated, unilocular radiolucent lesion with irregular borders was visible; extending from 24 to 26 regions on the orthopantamogram. General physical findings such as ultrasonography of the abdomen and chest were noncontributory. Based on the clinical and radiographic features, a provisional diagnosis of odontogenic cyst was made. The lesion was then surgically enucleated and sent for histopathological diagnosis [Figure 2].

Histopathological features
The hematoxylin and eosin staining showed that the epithelium was odontogenic in nature. They demonstrated a biphasic cellular pattern (typical of CCOC). Small islands of hyperchromatic, polygonal cells with eosinophilic cytoplasm were seen. These cells are seen surrounding the periphery of the tumor islands. Larger clear cells were separated by a clear delicate fibrous connective tissue stroma. Individual tumor cells had distinct cellular outline, round nuclei, prominent nucleoli and abundant clear cytoplasm. The cells exhibited a significant variable degree of nuclear and cytoplasmic pleomorphism. High mitotic activity was seen in the clear cells. At certain places, neural cells invaded the tissue [Figures 3 and 4].

Table 1 shows the different stains that we used and the outcome of the same.

Different stains for histopathological diagnosis and the outcomes with these histopathological features and immunohistochemical profile, the diagnosis of CCOC were established.

DISCUSSION
Hansen in 1985 coined the term “clear cell odontogenic tumor.” 1992 WHO classifications did not include this. However, later on the subsequent reporting and documentation led to its inclusion. Hence, the term tumor
### Table 1: Different Stains used and their outcome

| Stain                                      | Finding                                                                 | Figures |
|--------------------------------------------|-------------------------------------------------------------------------|---------|
| Per Iodic Acid Schiff Stain (PAS)          | Tumour cells are showing abundant diastase degradable PAS-positive granules, but they are negative for mucin |         |
| Mucicarmine Stain                         | Negative for mucin                                                     |         |
| Congo red stain                            | Negative for amyloid deposit                                            |         |
| CK5/6                                      | Tumor cells showed diffuse positivity                                   |         |
| CK-7                                       | Tumor cells showed and strong positivity                                |         |
| CK- 19                                     | Tumor cells showed weak positivity                                     |         |
| p63                                        | Diffuse and strong nuclear positivity of the nuclear cells             |         |
| Calretinin                                 | Negative in all tumour cells                                           |         |
| S-100 protein                              | Negative in all tumour cells                                           |         |
| SMA                                        | Negative in tumour cells. Staining was noted in smooth muscle cells and blood vessels in the stroma |         |

### Table 2: The different considerations for the clear cell variety of tumors

| Is it odontogenic? | Is it salivary gland origin? | Is it metastatic? |
|--------------------|-----------------------------|-------------------|
| Clear cell ameloblastoma: Odontogenic follicles exhibit VickersGorlin criteria Clear cell CEOT: polyhedral epithelial cells with prominent intercellular bridges and presence of amyloid and calcifications | Mucoepidermoid carcinoma: triphasic pattern (epidermoid, mucous and intermediate cells) with microcyst and macrocyst formation | Uncommon to jaws Metastasis from kidney, prostate, liver and thyroid considered Renal clear cell carcinoma exhibits rich dilated prominent sinusoidal vascular network |
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was replaced with carcinoma (CCOC).\textsuperscript{[1]} CCOC has a female predilection with the male/female ratio of 1:1.8, and the majority of cases have been diagnosed in patients older than 40 years (81.0\%). The mean age is 54.2 years, with 58.2 years for women and 41.8 years for men. Mandible is involved in 77.0\% and maxilla in only 23\% of cases.\textsuperscript{[3,4,5]} Of the reported cases, 6.8\% of cases have shown a radiopaque finding. Hence, lesion may be either radiolucent or radiopaque or mixed in nature.\textsuperscript{[6,7]} CCOC may exhibit biphasic, monophasic and ameloblastomatous patterns. Monophasic patterns have all clear cells throughout the lesion. Most of the tumors report a biphasic pattern, having clear nest cells and islands of polygonal cells. These polygonal cells surround the tumor periphery. In case, it resembles the pattern like that of ameloblastoma; there are clear cells within the network of follicular cells.\textsuperscript{[8]} The extent to which pleomorphism or hyperchromatism can occur varies from case to case and is not always a consistent parameter.\textsuperscript{[4]} The differential diagnosis of clear cell carcinoma of the jaws has a long list such as odontogenic tumors, tumors of the salivary glands and metastatic renal carcinoma.\textsuperscript{[9]}

Table 2 shows the different considerations for the clear cell variety of tumors.\textsuperscript{[10]}

Based on the present case findings, we were able to rule out the salivary gland as S-100 protein test was negative and renal carcinomas since intratumoral hemorrhage was clinically not present. This could have been a case of ameloblastoma, but since the site and presentation were not conventional,\textsuperscript{[11]} we ruled that out too. However, palisading was seen in the present case, which actually made us consider ameloblastoma as a differential diagnosis. The absence of amyloid as well as Liesegang’s ring calcification ruled out CCCEOT also.\textsuperscript{[12]} Mucoepidermoid carcinomas are distinguished by a triphasic architecture consisting of mucin-positive mucous cells, squamoid cells, along with intermediate cells.\textsuperscript{[13]} CK19 showed weak positivity, hence cannot be always definitive diagnosis.\textsuperscript{[1-2]} Histopathological and immunological overlaps result in difficulty for differentiating CCOC from the clear cell carcinoma of the salivary gland in the maxilla. This type of tumor is reported to have a recurrence rate of 38.25\%. Hence, long-term follow-up is necessary.

Three main mechanisms explain the clear cell change in cells.\textsuperscript{[12]}

1. Sparsity/loss of cell organelles
   - Rough endoplasmic reticulum is reduced, its cisternae show vacuolization
   - Secretory granules are decreased
   - The predominance of immature granules with limited optical density
   - Mitochondria are swollen, show the vacuolar transformation.

2. Accumulation of intracytoplasmic substances (glycogen, mucin, lipids, tonofilaments and zymogen granules)

3. Artifacts due to fixation and histologic techniques.

**Treatment**

Surgical control of CCOC was carried out with an en bloc resection of bone and soft-tissue involvement keeping 1 cm safe margins, to decrease the risk of recurrence. Adjuvant radiation therapy was given to this patient as there were extensive soft tissue and perineural invasion, positive nodes and/or extracapsular spread. Because of the potential for locoregional recurrence and/or late metastatic spread, the patient is kept under observation.\textsuperscript{[14]}

**Recurrence**

A recurrence of the same lesion at the same site was
observed after 1 year, which was treated with the radical surgical approach followed by radiation.

CONCLUSION

CCOC is a rare tumor. It has variations in its histological appearance. Hence, considering a holistic view of the tumor is essential. Differential diagnosis is important to prepare the treatment plan for a long-term follow-up of the patients. More robust reporting is needed to understand the biological nature of such tumors.

Declaration of patient consent
The authors certify that they have obtained all appropriate patient consent forms. In the form, the patient has given his consent for images and other clinical information to be reported in the journal. The patient understands that his name and initials will not be published and due efforts will be made to conceal identity.

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Conflicts of interest
There are no conflicts of interest.

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