Primary oral and maxillofacial liposarcoma: a clinicopathological and immunohistochemical study of eleven cases

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Abstract

Introduction: The present study was aimed to characterize the clinicopathological, immunohistochemical features and treatment outcomes of primary oral and maxillofacial liposarcomas by presenting the experience over a 16-year period at a tertiary referral Chinese institution for head neck cancer.

Material and methods: This retrospective clinical study included 11 cases of pathologically confirmed primary liposarcomas treated from January 1993 to September 2009. Detailed information regarding primary site, clinical manifestations, histopathological and immunohistochemical analysis, treatments and prognosis was collected and reported.

Results: Eight female and 3 male patients aged from 8 to 76 years old. These lesions occurred in buccal (3), parotid (2), temporal (2), tongue (2), palate (1) and oropharyngeal (1) region. They were histopathologically categorized into 4 subtypes based on WHO classification scheme: atypical lipomatous tumor/well-differentiated (4), myxoid (4), mixed-type (2) and pleomorphic (1) liposarcomas. Immunohistochemical staining indicated mostly positive for Vimentin and S-100 but negative for other markers. Most patients presented as slow-growing painless masses and underwent surgery alone or combined with postoperative radiotherapy. Two patients were misdiagnosed and inappropriate treated and developed local relapse before referred to our institute. No distant metastasis and one disease-related death were recorded during the follow-up (ranging: 1-11 years, mean: 4.5 years).

Conclusions: Oral and maxillofacial liposarcoma is exceedingly rare and has atypical clinical manifestations but characteristic histopathology. Complete excision with negative margins followed by long-term follow-up is recommended as the treatment of choice for these uncommon entities.

Key words: liposarcoma, oral maxillofacial region, surgery, prognosis.
The oral and maxillofacial tumor registry of Shanghai Ninth people’s hospital affiliated to Shanghai Jiao Tong University was searched and relevant medical records were reviewed to retrieve information for patients diagnosed as primary oral and maxillofacial liposarcomas between January 1993 and September 2009. Inclusion criteria comprised an oral and maxillofacial location for primary lesion and definitive diagnosis by established pathological criteria for liposarcoma. The archived H&E and immunohistochemical slides available were further retrieved and reviewed independently by the senior oral pathologists. The research protocol was approved by the Institutional Review Committee of Shanghai Jiao Tong University Medical School. Informed consents were obtained from patients or patient relatives. The clinical and follow-up information of these patients including demographics, primary tumor site, histological subtypes, treatment modality and outcomes from medical records were collected and presented.

During surgery, these lesions were excised about 1.5 cm away from the tumor margin. After surgical removal of the lesions, the tissue specimen were immediately transferred to oral pathology unit and subjected to pathological examination by frozen sections. The three-dimensional margins were also determined by frozen sections to avoid possible residual tumor. After surgery, the surgical samples were fixed in 10% buffered formalin and sectioned in 4-μm slides for hematoxylin-eosin staining and immunohistochemical staining using standard procedures. The commercial available antibodies to Vimentin, S-100, CD34, CD99, smooth-muscle actin (SMA), CKpan, Desmin, muscle-specific actin (MSA) were used together with DAB staining system with appropriate positive and negative controls. Immunoreactivity for each protein was determined based on the proportion of stained cells and expected subcellular locations. Negative control was prepared using phosphate buffer solution (PBS) instead of primary and second antibodies.

Most patients in this series underwent complete local excision with disease-free margins. To reduce the possibility of local recurrence, postoperative radiotherapy (about 55-60 Gy) was administered to selected cases. Two recurrent patients who were misdiagnosed and inappropriate treated in local hospital were received another surgical resection in our institute followed by chemotherapy. Large lesions which were more than 5 cm in largest dimension in other two patients and they were recommended for radiotherapy after surgery. No chemotherapy before and after surgery were given due to the insensitiveness of liposarcoma to chemical agents. All patients were recommended to participate in long-term follow-up at regular intervals.

Results

Demographic and clinical features

Through tumor registry search and further pathological confirmation of initial diagnosis, eleven cases of primary oral and maxillofacial liposarcomas were enrolled and briefly showed in Table I. Eight patients were females and the others were males, indicating a female predominance (8 : 3). The mean age of patients were 33.7 years. The lesions were found in buccal mucosa (3), temporal (2), parotid (2), ventral tongue (2), palate (1) and the lateral pharyngeal wall (1). Clinical manifestations of this malignancy varied largely depending on the areas involved. Most lesions typically began as slowly-growing and painless masses with dif-
ferent intervals between tumor onset and clinical presentation. The main reasons for patients to seek treatment were accelerated growth of masses and resulting facial disfigurement and the development of pain. Liposarcoma in the parotid region often resembled a benign tumor without obvious facial nerve paralysis. However, when large parotid lesions extended toward parapharyngeal space or parapharyngeal liposarcoma developed, diverse severities of airway obstruction, dysphagia and night snoring occurred (Figure 1). Computed tomography (CT) and Magnetic resonance imaging (MRI) were usually employed to determine the precise location and boundary as well as its relationship with adjacent tissues. Images of ALT/WD liposarcoma usually showed similar signal intensity equal to fat that closely resemble lipoma and characteristic thickened septa (Figure 1). Other subtypes of liposarcomas showed no remarkable difference to other soft-tissue malignancies in CT/MRI images. Among these patients, wide local excision with negative margins was the most common treatment modality. Radical resection was carried out for patients with local recurrence. Postoperative radiotherapy was also administered to four patients, mainly due to large lesions and recurrent cases. Follow-up period ranged from 1 to 11 years (mean: 4.5 years). Ten patients remained alive until last follow-up, while only 1 patient died of recurrent lesions adjacent to brain and cranial base. Two patients had recurrent lesions about 2-3 months after initial incomplete excision in local hospital before admitted to our department. No evidence of distant metastasis was recorded.

### Histopathological and immunohistochemical findings

Macroscopically, liposarcomas often appeared as soft-tissue mass containing diverse extent of hemorrhage and necrosis and varied from well-circumscribed, multilobulated and partially encapsulated to less well-delineated and more invasive and infiltrating with adjacent tissue. The colors of tumor cut surfaces ranged from yellow to grey and from greasy to mucoid and gelatinous mainly depending on different histopathologic types. Based on histopathology and the latest WHO classification for liposarcoma, eight cases were categorized as atypical lipomatous tumor/well differentiated (ALT/WD) and myxoid liposarcoma. Two cases were subtyped as mixed-type and 1 patient had pleomorphic liposarcoma.

Under microscopy, ALT/WD subtype was characterized by the presence of mature adipocytes with significant variation in cell size and scattered nuclear atypia and variable number of atypical stromal cells (Figure 2). The atypical stromal cells were spindle and often identified with hyperchromatic nuclei embedded within mature adipose tissue. Rare scattered and monovacuolated or multivacuolated lipoblasts could be found. The case with multiple recurrences was diagnosed as lipoma-like subtype of ALT/WD liposarcoma. No evidence of dedifferentiation of ALT/WD or other types of malignancy could be detected in the relapsed lesions. Myxoid liposarcoma consisted round primitive nonlipogenic mesenchymal cells and a few small signet-ring lipoblasts within prominent myxoid stroma (Figure 3). Rich and delicate branching capillary architecture was frequently observed and sometimes vessel and perineural invasions could be detected. As to pleomorphic liposarcoma, it comprised of pleomorphic lipoblasts in the background of pleomorphic sarcoma. The pleomorphic round and spindled tumor cells were mixed with multinucleated giant cells which were morphologically similar to malignant

| Case no. | Age [years] | Gender | Primary sites       | Pathologic subtype | Treatment  | Treatment outcome |
|----------|-------------|--------|---------------------|--------------------|------------|-------------------|
| 1        | 21          | Female | Right parotid region | Myxoid             | SE + PR    | NED               |
| 2        | 26          | Female | Right pharyngeal region | ALT/WD             | SE         | NED               |
| 3        | 25          | Female | Left buccal region   | ALT/WD             | SE + PR    | R, D              |
| 4        | 11          | Female | Left buccal region   | Pleomorphic        | SE         | NED               |
| 5        | 8           | Female | Left ventral tongue  | Myxoid             | SE         | NED               |
| 6        | 46          | Male   | Left palate          | Mixed-type         | SE         | NED               |
| 7        | 43          | Female | Left temporal region | Myxoid             | SE         | NED               |
| 8        | 28          | Female | Left buccal region   | Mixed-type         | SE + PR    | NED               |
| 9        | 48          | Female | Right temporal region | Myxoid             | SE + PR    | NED               |
| 10       | 39          | Male   | Right parotid region | ALT/WD             | SE         | NED               |
| 11       | 76          | Male   | Left ventral tongue  | ALT/WD             | SE         | NED               |

ALT/WD – atypical lipomatous tumor/well differentiated liposarcoma, SE – surgical excision, SE + PR – surgical excision + postoperative radiotherapy, NED – no evidence of disease, R – recurrence, D – death

Table I. Clinicopathological characteristics and treatment outcome of 11 oral and maxillofacial liposarcomas
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fibrous histocytoma (Figure 4). High mitotic activity and intramuscular invasion of tumor could be observed. Mixed-type liposarcoma showed the diverse morphological features of myxoid liposarcoma and ALT/WD or pleomorphic liposarcoma in the lesions with clear boundary. Among the mixed-type lesions, different parts had their own characteristic cell morphology and histopathological features (Figure 5).

Immunohistochemical staining results available for 7 cases showed that vimentin and S-100 were positive most cases. However, CD99, α-SMA, MSA, CKp and Des commonly appeared negative (Table II). Despite limited values for direct diagnosis for liposarcoma based on immunohistochemical staining, these results were valuable for differential diagnosis among diverse types of malignancies.

Discussion

Liposarcoma, first described by Virchow in 1857, is considered as one of the most common soft tissue sarcomas in adults. However, it rarely involves the oral and maxillofacial region [9, 12]. Its rarity and lack of characteristic symptoms and signs reduced the opportunity to early detect and management [13]. The lack of adequate information regarding the clinicopathological features and prognostic factors for this rare tumor further makes early diagnosis and treatment planning difficult. Our data reported here may improve our understanding of clinicopathological features of oral and maxillofacial liposarcoma to avoid misdiagnosis and mistreatment in the clinic.

Previous reports showed a male/female ratio of about 2 : 1 for liposarcoma in the head and neck region.
region, indicating a male preponderance for this tumor [2, 11]. However, an obvious female preponderance (female/male: 8 : 3) was found in our series. Similarly, Zheng et al. reported female preponderance with the female/male ratio of 4 : 1 in their series of Chinese origin [5]. The discrepancy of gender distribution may be due to limited number of cases analyzed and/or genetic in origin. Moreover, the average age (33.7 years) in the Chinese cohort is also lower than that in previous studies, as previous reports indicated the age of presentation ranging from 30 and 60 years with a peak in the sixth decades [14]. Notably, two female pediatric patients with 8 and 11 years old were included, being myxoid and pleomorphic subtype arising from buccal mucosa and ventral tongue, respectively. Pediatric liposarcoma is extremely unusual and significantly different from adult cases. Alaggio et al. reported that conventional myxoid liposarcoma was the most common subtype with excellent prognosis in patients younger than 22 year of age [15]. These two pediatric patients underwent wide local excision with negative margins and had no local recurrence and metastasis during the follow-up. This finding reminds the clinicians that oral and maxillofacial liposarcoma can even occur in children and adolescent.

As the sites of predilection for oral liposarcomas, the buccal mucosa is the most common site in this series which is consistent with previous studies [16]. Of note, two parotid liposarcomas were found and both were pathologically classified as ALT/WD. No such lesions were found in minor salivary glands. One patient was initially misdiagnosed as pleomorphic adenoma and inappropriately treated in local hospitals, then developed recurrent lesion 2 month later. To our knowledge, although ten cases of liposarcomas in the parotid region were reported in the English literature so far, only one case with definitive diagnosis of parotid ALT/WD was previously documented by Fanburg-Smith et al. with unspecified treatment and no recurrence during 2 years follow-up [9]. Therefore, our two cases might add some important information to better understanding of parotid ALT/WD and remind surgeons to be alert about the diagnosis and treatment of such rare tumor in the parotid region.

The liposarcomas within oral and maxillofacial region often appear as painless, slow-growing swelling or soft-tissue masses. On the first examination, these disease entities tend to be perceived as benign tumors. Three patients in this series had initial misdiagnosis and incomplete excision and 2 patients developed local recurrence before

Figure 2. Histopathological and immunohistochemical examination of ALT/WD liposarcoma. Mature adipocytes with diverse size, multi-vacuolated lipoblasts and variable number of atypical stromal cells as showed by H + E staining (A). Immunohistochemical staining for vimentin (B), S-100 (C), CD99 (D). Magnification 200×
Figure 3. Histopathological and immunohistochemical examination of myxoid liposarcoma. Multivacuolated and signet-ring primitive cells within the prominent myxoid stroma as showed by H&E staining (A). Magnification 100×. Immunohistochemical staining for, S-100 (B), vimentin (C) CD34 (D). Magnification 400×

Figure 4. Histopathological examination of pleomorphic liposarcoma. The tumor consists of pleomorphic sarcoma (left area) and pleomorphic lipoblasts (right area) as showed by H&E staining. Magnification 100× (A). The pleomorphic sarcoma is similar to malignant fibrous histocytoma and contains round and pleomorphic tumor cells with mitosis. Magnification 400× (B). The lipoblasts shown nuclear atypia. Magnification 400× (C)

Figure 5. Histopathological examination of mixed-type (ALT/WD & myxoid) liposarcoma. The lesion consists of ALT/WD (upper area) and myxoid liposarcoma (lower area) with apparent boundary as showed by H&E staining. Magnification 100× (A). The ALT/WD (B) and myxoid part (C) each shows characteristic histology. Magnification 400×
referred to our department. Therefore, differential diagnosis of such masses is essential and should include various benign and malignant lesions such as lipomas, lymphadenopathies, branchial cyst and Hodgkin lymphoma, et al. The presence of fat in CT and MRI images may suggest a lipomatous tumor such as lipoma and liposarcoma. Several characteristic findings including more than 75% fat, thick septa may contribute a lot to differential diagnosis between lipoma and ALT/WD [17, 18]. However, it’s fairly difficult to definitely diagnose such masses between liposarcomas and other soft tissue neoplasm based on radiography.

Histological examination of liposarcoma is pivotal for diagnosis establishment. Tumor cells with characteristic morphology contribute to diagnosis and subtype classifications. The most common histologic subtypes in our series, in agreement with previous findings, are ALT/WD and myxoid liposarcoma. The lipogenic differentiation marker S-100, indicative of the presence of lipoblasts, was stained positive in ALT/WD and myxoid liposarcoma, but negative in other subtypes. Although no specific molecular markers exist for its diagnosis, however, such markers by immunohistochemistry may be beneficial for differential diagnosis and subtype classification. The cytogenetic feature of ALT/WD and dedifferentiated liposarcoma has been revealed in the form of supernumerary ring or giant marker chromosomes with 12q13-15 amplification [19]. Several tumor-associated genes including MDM2, HMGA2 and CDK4 were identified and proven beneficial for differential diagnosis by immunohistochemical detection or fluorescent in situ hybridization [20]. Moreover, fine needle aspiration biopsy was suggested as a valuable and promising technique for preoperative diagnosis especially in the deep region [21]. Therefore, further studies were warranted to determine the roles of such tumor biomarkers in early detection, pathological categorization and prognostic prediction for liposarcoma [22].

Complete surgical excision with free margins is the primary treatment option for liposarcomas [2, 9, 13]. The values of adjuvant radiotherapy and chemotherapy for liposarcoma still remain controversial. Radiotherapy alone is occasionally considered as an alternative for selected cases especially for impossible total excision or recurrent tumors. Careful follow-up after complete excision has been recommended for monitoring recurrence and distant metastasis. Local recurrence, usually resulting from incomplete excision becomes a common challenge for successful management of liposarcoma. Although these tumors sometimes appear well encapsulated, wide excision should be emphasized to avoid any microscopic residual disease. In our cases, excisions were performed at least 1.5 cm from the palpable margins of lesions to guarantee complete removal when possible. However, the proximity of vital neurovascular structures and ensuing severe morbidities caused by wide excision sometimes limited the extent of excision. Therefore, adjuvant postoperative radiotherapy is sometimes needed for selected cases especially when complete excision was unattainable. It has been suggested that radiotherapy delayed tumor growth and prevented local recurrence [23]. Colledge et al. reported that patients treated with surgery alone had a 5-year survival rate of 83% as compared with 63% for those treated with combined surgery and radiotherapy [2]. Four patients reported here underwent surgery combined postoperative radiotherapy and had satisfactory results without evidence of recurrence. Based on our experience and previous reports, adjuvant radiotherapy should be considered as a valuable treatment alternative for local disease control after incomplete excision or suspicious disease left. In addition, no study has offered conclusive evidence supporting the treatment utility of chemotherapy for liposarcoma.

Although previous studies suggested that tumor size, histological subtype and treatment correlated with biological behaviors and prognosis of liposarcomas, we didn’t confidently draw conclusions about the relationship between prognosis and these clinicopathological variables largely due to limited number of cases. But, in general, oral and maxillo-

| Case | Vlm | S-100 | CD34 | CD99 | α-SMA | CKp | EMA | Des |
|------|-----|-------|------|------|-------|-----|-----|-----|
| 1    | +   | +     | –    | –    | –     | –   | –   | –   |
| 4    | +   | –     | –    | –    | –     | –   | –   | –   |
| 6    | +   | ±     | –    | –    | –     | –   | –   | –   |
| 8    | +   | ±     | +    | –    | –     | –   | –   | –   |
| 9    | +   | –     | –    | –    | –     | –   | –   | –   |
| 10   | +   | +     | –    | +    | –     | –   | –   | –   |
| 11   | +   | +     | ±    | –    | –     | –   | –   | ±   |

Vlm – vimentin, α-SMA – α-smooth muscle actin, MSA – muscle-specific actin, CKp – pan-cytokeratin, EMA – epithelial membrane antigen, Des – desmin, positive: +, negative: –, partial positive: ±
Facial liposarcomas had good prognosis as supported by our data and others findings, presumably owing to easy detection and timely therapeutic intervention.

In conclusion, we retrospectively reviewed 11 cases of primary liposarcomas within oral and maxillofacial region and described the clinicopathological features and treatment outcomes in detail. Our study indicated that these rare tumors developed from various anatomic sites, most frequently in the buccal mucosa and usually identified as ALT/WD and myxoid subtypes by histopathology. The treatment outcomes of these patients receiving surgery or together with radiotherapy were favorable with 90.9% disease-free survival during the follow-up, rare recurrence and no metastasis. It’s critically essential for clinicians to keep the possible diagnosis of liposarcoma in mind when diagnosing masses in the oral and maxillofacial region. More retrospective and perspective studies are still needed to establish the optimized diagnosis and treatment regimes for oral and maxillofacial liposarcoma.

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