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Lymphoepithelial Cysts of Oral Mucosa: Two Cases in Different Regions

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Abstract

Lymphoepithelial cyst of the oral cavity is a rare cystic lesion that presents as an asymptomatic, well-circumscribed, yellowish small submucosal nodule covered by normal overlying mucosa, usually located in the floor of the mouth or in the ventral or posterolateral surface of the tongue. Histopathological examination reveals a cyst lined by a stratified squamous epithelium surrounded by lymphoid tissue. In this paper, we report two cases of oral lymphoepithelial cyst, one on the anterior floor of the mouth and another on the posterior lateral tongue. In both cases, the patients were women aged approximately 50 years and the lesions were treated by local surgical excision, with no evidence of recurrence after 2 years of follow-up.

Key words: Lymphoepithelial cyst — Ectopic lymphoid tissue — Oral lymphoepithelial cyst — Oral cysts — Benign oral lesion

Introduction

Oral lymphoepithelial cyst (OLC) is a relatively rare lesion that develops within lymphoid tissue of unknown etiopathogenesis. These cystic lesions, usually located in the lateral aspect of the neck, were first described by Bhaskar and Bernier as “branchial cyst” in 1959.

Clinically, OLCs present as movable, painless submucosal nodules with a yellow or yellow-white discoloration. Occasional cysts are transparent. Usually the OLC is located on the floor of the mouth or on the ventral or posterolateral surface of the tongue. Approximately half of all intraoral cases are located on the floor of the mouth.

Histopathologically, it presents as a central cystic lesion lined with stratified squamous epithelium with desquamated keratin in the lumen. Surrounding the cyst lining there are variable amounts of lymphoid tissue and a
fibrous connective tissue capsule. Two cases with OLC are reported in this paper.

Case Reports

1. Case 1
A 57-year-old woman presented with a nodule on the floor of her mouth. Her personal and family medical histories were unremarkable. Intraoral examination revealed a submucosal nodule 2 mm in diameter. The nodule was non-elastic, movable, and soft, appearing as a yellow discoloration on the left side of the floor of the mouth, adjacent to the lingual frenum (Fig. 1). The mucosal surface was intact and the patient complained of no pain or discomfort. The clinical diagnosis was lipoma or OLC. The lesion was totally excised under local anesthesia. Histopathological examination revealed a cystic lesion lined with thin parakeratinized stratified squamous epithelium; lymphoid tissue exhibiting germinal centers was observed on the cyst wall (Fig. 2). The diagnosis was lymphoepithelial cyst. No complications were encountered in the postoperative period of two years.

2. Case 2
A 55-year-old woman visited a dental clinic for a routine examination. On physical examination, an elastic, hard, submucosal nodule, yellow in color and 15 mm in diameter was observed on the posterior edge of the right side of the tongue (Fig. 3). The overlying mucosa was normal and the patient complained of no discomfort. The clinical diagnosis was lipoma or OLC. Under local anesthesia, the nodular lesion was excised together with surrounding normal tissues and sent for histopathological examination. Microscopically, the cystic cavity was lined with a thin flattened layer of parakeratinized stratified squamous epithelium, and was surrounded by a well-circumscribed mass of lymphoid tissue, with desquamated cells and lymphocytes present in the lumen (Fig. 4). The

![Fig. 1](image1.png)
Fig. 1 Submucosal nodule, 2 mm in diameter, with yellow discoloration, on left side of floor of mouth adjacent to lingual frenum

![Fig. 2](image2.png)
Fig. 2 Cystic lesion lined with thin stratified squamous epithelium; lymphoid tissue exhibiting germinal centers observed on cyst wall
pathological diagnosis was OLC. Three years after surgery, the postoperative course was uneventful.

**Discussion and Conclusion**

Bhaskar and Bernier first recommended the term lymphoepithelial in 1959 in a report of 468 cases of branchial cysts of the neck\(^{14,25}\). In 1962, Gold and Lewittown described an OLC as a case report, which was the first one in the world\(^{14,25}\). A lymphoepithelial cyst may occur in virtually any organ. After the neck and oral cavity, the most frequent sites reported in the literature are the pancreas and thyroid gland. Pancreatic lesions are also believed to be benign epithelial inclusions embedded within the pancreas during embryogenesis\(^{30}\).

OLCs are rare cystic lesions lined with stratified squamous epithelium surrounded by lymphoid tissue and have been reported as branchial cysts, branchial cleft cysts, branchiogenic cysts, or pseudocysts\(^{3}\).

The pathogenesis of OLCs is uncertain. The two theories that have been proposed are the entrapment and obstruction theories. Several authors have suggested that ectopic foci of glandular epithelium become entrapped within nodal tissue and may proliferate ultimately to form a cyst\(^{5,27}\). However, other authors have suggested that OLCs are the result of obstruction of crypts of otherwise normal oral tonsils\(^{17}\).

Clinically, an OLC presents as a small asymptomatic, well-circumscribed, yellowish, elevated, mobile, submucosal tumor of varying duration\(^{3}\). Most of the lesions are located in the floor of the mouth (60%) or on the lateral and ventral surfaces of the tongue (40%)\(^{5,25}\). In our cases, the sites and diameters of the lesions concurred with the literature. Giunta and Cataldo\(^{13}\) and Buchner and Hansen\(^{7}\) also observed that these cysts may vary in diameter from 2 to 15 mm. OLCs occurring on the parotid gland are often associated with immunocompromised patients and lymphoproliferative disorders\(^{20}\). Table 1
shows cases reported in the literature, since the first report by Gold and Lewittown (1962)\(^{14}\), excluding cases related to immunocompromised patients and salivary glands. The OLC is covered with intact mucosa, clinically interpreted most often as mucocele or lipoma. Although most of the cases reported are from male patients\(^{27}\), our two cases were from women, in their fifth decade of life. The actual prevalence of OLCs is not known, because many of these cystic lesions are not diagnosed. The cysts are often misdiagnosed as mucous cyst, other cystic lesions or even as a lipoma, and many times they are

<table>
<thead>
<tr>
<th>Publication</th>
<th>Cases</th>
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<th>Age</th>
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<tr>
<td>Gold &amp; Lewittown, 1962(^{14})</td>
<td>1</td>
<td>Floor of mouth</td>
<td>M</td>
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<td>Floor of mouth</td>
<td>M</td>
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<td>Vickers &amp; Von Der Muhl, 1966(^{27})</td>
<td>1</td>
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<td>M</td>
<td>30</td>
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<td>Bhaskar &amp; Colonel, 1966(^{5})</td>
<td>24</td>
<td>Floor of mouth (n = 15); Tongue (n = 8); Palate (n = 1)</td>
<td>M = 17/F = 7</td>
<td>40 (15–65)</td>
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<td>Young &amp; Claman, 1967(^{29})</td>
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<td>Floor of mouth</td>
<td>M</td>
<td>42</td>
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<td>13</td>
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<td>M = 12/F = 1</td>
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<td>Acevedo &amp; Nelson, 1971(^{3})</td>
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<td>Giunta &amp; Cataldo, 1973(^{3\text{a}})</td>
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<td>Buchner &amp; Hansen, 1980(^{7})</td>
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<td>Floor of mouth (n = 19); Tongue (n = 14); Palate (n = 4); Retromolar area (n = 1)</td>
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<td>Tongue</td>
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<td>1</td>
<td>Tongue</td>
<td>F</td>
<td>27</td>
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<tr>
<td>Khelemsky &amp; Mandel, 2010(^{8\text{c}})</td>
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<td>Floor of mouth</td>
<td>M</td>
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<td>Floor of mouth (n = 2); Tongue (n = 4); Palate (n = 1); Oropharynx (n = 2)</td>
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<td>38 (16–60)</td>
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<tr>
<td>Yang et al., 2011(^{27})</td>
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<td>Floor of mouth (n = 46); Tongue (n = 60); Palate (n = 6); Buccal mucosa (n = 3); Others (n = 5)</td>
<td>M = 37/F = 83</td>
<td>44 (2–75)</td>
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</table>
not examined microscopically, and incidence may vary among different institutions.

Differential diagnosis is important\(^{10}\) for definitive therapy. Several lesions appear as submucosal nodules in the floor of the mouth. Lipomas are usually located in the buccal mucosa, and rarely affect the floor of the mouth. It can also have a yellowish appearance but is usually larger than an LC. Some salivary lesions must also be considered as LCs in differential diagnosis. Mucoceles are the most frequent benign salivary gland in the oral cavity and can have the same clinical aspects as an LC. Fluid drainage and size fluctuation facilitates differential diagnosis between the two lesions. Salivary stones or sialolithiasis can also present as a submucosal mass in the floor of the mouth, and can induce pain depending on duct obstruction and salivary gland swelling.

An LC is usually encapsulated and small in size, facilitating surgical excision in an ambulatorial environment. There are no LC recurrences reported in the literature, and recovery after surgical removal should be fast and without recurrence.

References


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