Case Report
Cheilitis Glandularis: Two Case Reports of Asian-Japanese Men and Literature Review of Japanese Cases

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Cheilitis glandularis (CG) is a rare disorder characterized by swelling of the lip with hyperplasia of the labial salivary glands. CG is
most frequently encountered in the lower lip, in middle-aged to older Caucasian men; however Asian cases are very rare. In
this paper we present two cases of CG in Asian-Japanese men. One was a 23-year-old male with CG of the superficial supplicative
type. The other was a 54-year-old male with deep supplicative type. We also reviewed the Japanese cases of CG in the literature
and discussed about clinical feature of Japanese CG.

1. Introduction

Cheilitis glandularis (CG) is a rare disorder characterized by swelling of the lip with hyperplasia of the labial salivary glands. The term cheilitis glandularis was first used by Volkman in 1870 to describe a disorder that presented as a chronic, supplicative inflammation of the lower lip, characterized by swelling of the mucous glands and associated with mucopurulent discharge through dilated ductal openings [1, 2]. The histopathologic features of CG show nonspecific chronic inflammation of minor salivary gland tissue. Differential diagnosis of the lip edema may include angioedema, exfoliative cheilitis, cheilitis granulomatos, or elephantiasis nostras [3].

Cheilitis glandularis is most frequently encountered in the lower lip, in middle-aged to older Caucasian men [4, 5]; Japanese cases are very rare. In this paper we present two cases of CG in Asian-Japanese men. One was a 23-year-old male with swelling of both lips and superficial ulceration. The diagnosis was CG of the superficial supplicative type. The other was a 54-year-old male with upper lip swelling and abscess formation. This case was determined to be the deep supplicative CG type and was treated by surgical excision. We also review the Japanese cases of CG in the literature and discuss the features of Asian-Japanese CG in Japan.

2. Case Reports

Case 1. A 23-year-old Asian-Japanese man consulted the Department of Oral and Maxillofacial Surgery of Tsukuba University Hospital complaining of upper and lower lip swelling with superficial ulceration in August 2005. He noticed these masses at first in August 2004 and visited dermatology medical practitioner. He had applied cepharanthine, hochuekkito (chinese herbal remedy), triamcinolone acetonide, irsogladine maleate, and koujin powder (chinese herbal remedy) as recommended by his dermatologist, and the swelling was reduced after two months of treatment.
Figure 1: Extraoral photograph of the patient in Case 1 showing the everted lower lip.

Four months later the swelling of both lips recurred following a small superficial ulceration, and it was reduced again two months later. In June 2005 the swelling of both lips recurred, but there was no resolution of the symptoms for more than 2 months. The patient consulted his dentist, who referred him to our department for further evaluation and treatment. His previous history revealed nothing remarkable. The patient had no known drug allergies but did have a food allergy to shellfish. Clinical examination revealed a well-developed and nourished man in no apparent distress. Intraoral examination revealed good dental health. Extraoral examination revealed that the left side of his lower lip was enlarged and grossly everted, and the right side of his upper lip was slightly everted and rather firm to palpation with indiscernible borders (Figure 1). The left lower and the right upper labial mucosa had a round ulceration, approximately 8 mm in diameter (Figures 2(a) and 2(b)), and produced cloudy exudate at ductal openings. A biopsy of the tumescent region including the mucosal area in the lower lip was carried out. Histologically, the epithelium was slightly thickened, and moderate inflammatory cell infiltration with lymphocytes, plasma cells, and neutrophiles was observed at subepithelial connective tissues. The labial glands were slightly hyperplastic and associated with focal infiltration of lymphocytes and plasma cells at interlobular region (Figure 3). These findings were compatible with CG. We informed the patient of the diagnosis, but because he did not wish to have the affected regions removed surgically, followup was continued. The enlargement of both lips resolved spontaneously within a month, and there has been no relapse in two years.

Case 2. A 54-year-old Asian-Japanese man consulted the Department of Oral and Maxillofacial Surgery of Tsukuba University Hospital complaining of upper lip swelling like phlegmonous abscess. He noticed small masses in his upper lip in October 2005 but had not sought treatment. In January 2006 these masses suddenly became inflamed, and he consulted a dentist, who referred him to Tsukuba University Hospital. The patient had a history of hypertension but was not on medication for it, and he had no known allergies. Clinical examination revealed a well-developed and nourished man in no apparent distress. Extraoral examination revealed a lesion that was firm to palpation with indiscernible borders, and diffuse swelling measuring approximately $3.0 \times 2.0$ cm and extending superiorly from the right upper lip to the cheek region and posteriorly to the angle of the mouth. Intraoral examination revealed a firm, submucosal swelling in the right buccal mucosa, extending from the maxillary right first permanent molar to the lip region. There was a fistula in the center of the lesion in the mucous membranes of the inside right upper lip. There was severe periodontitis around the upper molar teeth, but probe examination indicated that the fistula was cecal and 20 mm deep, and no connection was found between the periodontal gingiva and the lip. Infectious focus is not clear, so we suspected cheilitis, and incisional biopsy of the nodular region including the mucosal area was carried out. Histologically, the labial glands were surrounded by thick fibrous connective tissues associated with lymphocyte infiltration and diagnosed as cheilitis. Fibrosis and lymphocyte infiltration also occurred at interlobular region in the labial

Figure 2: (a) and (b) show the upper and lower lip of the patient, respectively. There was painless swelling, and induration on both lips with abscess. The surface mucosa showed a color change and produced cloudy fluid at the sites of ductal openings.
gland (Figure 5(a)). In a week the inflammation was resolved by drainage and antibiotic medication, but a small nodule with a mucus-discharging punctum remained. A cystic lesion (9 × 4 mm) was detected in the upper lip by magnetic resonance imaging (MRI). The signal intensity of the mass was relatively low compared with the surrounding soft tissue on T1-weighted magnetic resonance (MR), but remarkably high on T2-weighted and short T1 inversion recovery MR, which suggested fluid collection (Figures 4(a), 4(b), and 4(c)). Five months later, the patient had a persistent nodule that was exudative in his upper lip, and he was scheduled for surgical excision of the nodule. We designed a spindle-shaped incisional line on the nodule that included the opening of the duct and excised the nodule along with the surrounding scar tissue under a general anesthetic. The tissue specimen submitted for histopathologic examination measured 1.5 × 1.0 × 1.0 cm. Histologically, extensive fibrosis associating with lymphocyte infiltration was observed at the central region of the specimen. A few atrophic labial glands and dilated duct were scattered among the fibrosis (Figure 5(b)). These findings were compatible with CG. The patient is currently being seen on a regular recall basis and is doing well after 1 year.

3. Discussion
Cheilitis glandularis (CG) is a rare disorder characterized by swelling of the lip with hyperplasia of the labial salivary
Cheilitis glandularis is most frequently encountered in the lower lip, in middle-aged to older Caucasian men [4]; however, our cases were in Asian-Japanese men. In the first case, both lips were affected, and superficial suppurative type CG was diagnosed. In the second, the upper lip was affected and deep suppurative type CG was diagnosed. Asian-Japanese cases of CG are very rare and are mainly reported in Japanese domestic journals. To our knowledge, 17 cases, including ours, have been reported in the domestic literature between 1952 and 2009 [7, 10, 11]. Our findings from this review of Japanese cases are summarized in Table 1. The mean age is 58 years (range 1 to 76) and the sex ratio is 6:11 (f:m). In this literature search, 11 of the 17 Japanese cases affected the upper lip or included both lips. There are only two reported cases of the simple type. Among the reported cases of GC in the international literature, few involve the upper lip [12]. However, our literature search showed that the upper lip of Asian-Japanese patients was often affected. The reason for this difference in morbidity is not clear, but it may reflect racial or etiological differences. The etiology of CG is still unknown, but it has been suggested to be an autosomal dominant hereditary disease for which smoking, poor oral hygiene, chronic exposure to sunlight and wind, and a compromised immune system are predisposing factors [2, 8, 9, 13, 14]. Swerlick and Cooper reviewed forty-eight cases of CG described in the literature and suggested that CG patients can be categorized...
into three groups [15]. The largest group develops CG as a result of marked, chronic sun and wind exposure. The second group has a history suggesting a coexistent atopic diathesis. The third group has factitious cheilitis. In their review, CG represents the usual clinical presentation of cheilitis that develops after chronic exposure to one or more irritants. Our presented cases did not have obvious causes of CG; however, in the first case, diathesis is suspected because the patient had a food allergy. In the second case, the patient had periodontitis of a molar tooth near the CG, which could act as a chronic irritant to facilitate CG development.

CG is an unusual disease, characterized by a painless, enlarged everted lip with mucoid-draining pores, that may be encountered in clinical practice. In some cases, CG may degenerate into squamous cell carcinoma [5, 16, 17]. However, like our first case, it is sometimes self-healing. Thus, there may be many unreported cases of CG in which the patient does not seek treatment because the CG resolves spontaneously. Further research of CG, including thorough investigation of its etiology, is necessary to improve our understanding of this disease.

4. Conclusion

We reported two cases of CG affecting the upper and both lips in Asian-Japanese men. One was superficial suppurative type and the other was deep suppurative type.

References


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