Unilateral absence of submandibular gland secondary to stones. Aplasia versus early atrophy

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Introduction
The association of bilateral aplasia of major salivary glands with other ectodermal defects has been previously described in the literature. These include lacrimeo-auriculo-dento-digital syndrome, hemifacial microsomia, Treacher Collins syndrome, and ectodermal dysplasia (1,2). Analysis of these abnormalities suggests that the aplasia of salivary glands likely results from a disturbance during fetal development of the first and second branchial arches, but the exact etiology is unknown. Unilateral submandibular gland aplasia is even rarer with less than ten cases reported in the literature (3-12). The etiopathogenesis of isolated absence of a
major salivary gland without other developmental anomalies is still unclear. Obstructive sialadenitis, attributable for example to calculi formation, leads to salivary gland inflammation and eventually to atrophy. To the authors’ knowledge, only two cases of sialolithiasis accompanied by submandibular gland aplasia has been reported (10). We describe two new cases of submandibular gland absence associated with a salivary calculus and discuss the possible pathogenesis of this entity.

**Case Reports**

**Case 1**
A 70 years old man, allergic to penicillin, presented to his dentist with two days pain and swelling at the left submandibular region. The patient clinically improved after one week treatment with clindamycin (300mg/6h PO) and diagnosis of sialadenitis was made. Two months later, he develops the same symptoms and then was referred to our Department of Oral and Maxillofacial Surgery of our Institution.

Intraoral examination revealed a large, firm, non-tender swelling in the left Wharton’s duct. A Computerised Tomography (CT) showed a 2x2.6 cm calculus located in the submandibular region (Fig. 1) and severe atrophy of the ipsilateral gland. The stone was removed under local anaesthesia and postoperative course was uneventful.

**Case 2**
A 30-year-old female patient was referred by her dentist to the Department of Oral and Maxillofacial Surgery of our Institution for evaluation of a radio-opaque imaging showed in a panoramic radiography (Fig. 2). A medical history indicated that the patient had no systemic medical problems, previous history of swelling in cervical region, dryness of the mouth or dysphagia. Intraoral examination revealed a calculus on the right Wharton’s duct, but the submandibular gland on this side could not be located. Both sublingual carunculae were normal, but no saliva was observed after manual milking of right submandibular region. An adequate amount of saliva and normal salivary flow through the papillae of the left Wharton’s duct was observed after stimulation. Clinical evaluation of parotid glands had also normal features. There were no facial or auricular malformations. CT examination

**Fig. 1A.** CT image showing a large calculus in the left Wharton’s duct. B. Neck CT demonstrating severe atrophy of the left submandibular gland.

**Fig. 2.** Panoramic radiography shows a radio-opaque imaging typical of salivary calculus.
revealed a calculus in the right submandibular duct and the absence of the right submandibular gland (Fig. 3). No other salivary gland abnormalities were found. The removal of the sialolith in a stump of Wharton’s duct and sialodochoplasty was successfully performed through a transoral approach under local anaesthesia and postoperative course was uneventful.

Discussion
We report, to the best our knowledge, the second report of two cases of submandibular gland atrophy associated with a salivary calculus. Recently, Koo et al. (10) have reported the first description of two cases with this particular association. Several studies reviewed could explain this rare condition. The ligation of the excretory duct of rat submandibular glands has frequently been used to investigate regeneration of atrophic glands. Following experimental ligation of the rat submandibular duct, Osailan et al. (13) reported an extensive glandular atrophy with neutrophil infiltrates in the early stage of obstruction (1-18 h) followed by a monocyte invasion, evident after 24 hours. After 7 days of duct ligation, the submandibular glands showed severe atrophy of most acini (14). The obstruction could have provided negative feedback to the gland with reduction in salivary flow. After removal of the obstruction, induced atrophy can be reversible. Salivary glands could recover their functionality, secreting normal amounts of saliva with a broadly normal composition (15). In the cases reported, absence of the submandibular gland may have been the result of the complete acinar atrophy secondary to an early obstruction of Wharton’s duct. The submandibular glands were replaced by fat in the CT images, as the cases reported by Koo et al. (10). This process could explain the persistence of the excretory system of the gland and the absence of other ectodermal abnormalities. Unfortunately, two cases made it difficult to determine an etiologic factor or establish an association. In our opinion, the unilateral absence of submandibular glands was secondary to a glandular atrophy caused by an obstructive sialolithiasis.

References