Aplasia of Left Submandibular Gland- A Case Report

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The congenital absence of major salivary glands is a very rare disorder. The cause of the absence has not been determined. Isolated absence of unilateral submandibular gland is an unusual entity with less than 14 cases reported in the literature. The etiopathogenesis of isolated absence of major salivary gland without other developmental anomalies is still unclear. In this case we report a 42 year old male who came with a complaint of suprasternal neck swelling and on ultrasonography and magnetic resonance imaging incidentally left submandibular aplasia was detected. The literature on this rare occurrence is reviewed and significance of the patient is discussed.

INTRODUCTION-
Congenital major salivary gland is a very uncommon disorder (1, 2, 3, 4). The precise incidence of major salivary gland agenesis is difficult to establish due to asymptomatic nature of many cases (3).

The first case of salivary gland agenesis was described in 1885; and about 30-40 cases have been reported in the literature (1, 5, 6).

Most of the cases of congenital absence of major salivary glands; may be associated with multiple other developmental anomalies of the face (1, 4). These anomalies may be like atresia of lacrimal puncta, mandibulo-facial dysostosis or congenital malformations of the temporo-mandibular component (1, 2, 4, 7, 8, 9).

Developmental anomalies of major salivary gland include total absence of major salivary gland (aplasia) or reduced glandular tissue associated with salivary hypofunction (hypoplasia). Any of the major glands may be absent unilaterally or bilaterally (9).

Analysis of these abnormalities suggest that the aplasia of salivary glands likely results from the disturbance during fetal development of first and second branchial arches but the exact etiology is not known (10).

Unilateral submandibular gland aplasia is even rarer with less than 10-14 cases have been described (1, 10, 11-20). Most of them were incidental findings because of lack of symptoms.

In this study we report a case of an incidentally detected unilateral aplasia of left submandibular salivary gland demonstrated by ultrasonography and magnetic resonance imaging.

CASE REPORT-
A 42 year male patient was referred for ultrasonography for the complaint of suprasternal swelling in the neck. On examination no such swelling was palpated and the patient didn’t complain about any symptoms apart from anxiety. The patient was otherwise healthy.

Inspection showed an asymmetry in the submandibular region. Extra oral palpation was unremarkable. Oral mucosa had no alterations. The rest of oropharyngeal and cervical examination was normal. There was no enlargement of submandibular gland region.

Neck ultrasonography was performed for initial radiological evaluation

Fig no-1: Ultrasonography of right and left submandibular region showing the absence of left submandibular gland.

Left sublingual gland was normal; no areas of necrosis and calcification were noted. But left submandibular gland could not be identified. No anomalies were observed in remaining major salivary glands. Cervical adenopathy was absent. These findings were suggestive of absence of unilateral i.e. left submandibular gland with normal left sublingual gland.

Our study was completed with magnetic resonance imaging which revealed the absence of left submandibular gland and normal left sublingual gland. Right sublingual, right submandibular and both parotids were normal. Patient was asymptomatic hence required no treatment.
Isolated unilateral major salivary gland aplasia is a rare entity with only few cases reported in the literature to date. The aplasia is likely due to arrest in organogenesis but the exact etiology is unknown. Agenesis of major salivary glands may be associated with ectodermal defects of first and second branchial arches, lacrimal punctum aplasia or hypoplasia and agenesis of lacrimal glands and may be the result of some disturbing influence in early fetal development.

Submandibular gland aplasia may be associated with severe caries in mandibular permanent incisor teeth as well as other signs of salivary hypofunction, such as dry mouth, disturbed oral sensation and oral infections. In our case the patient did not suffer from any problems related to salivary hypofunction and was asymptomatic as the aplasia was confined to one gland only and the secretions of other glands compensated for it. Additionally all other cases of unilateral submandibular gland aplasia reported in the literature had no accompanying developmental disorders. Also in our case we did not encounter any associated anomaly.

In previous case reports hypertrophy of sublingual salivary gland in association with absence of ipsilateral submandibular salivary gland was observed. However in our case no significant hypertrophy of left sublingual gland was seen.

Salivary gland aplasia can be diagnosed with variety of imaging techniques which include computed tomography, magnetic resonance imaging, ultrasonography, sialography or nuclear medicine. Among these methods ultrasonography is cheap, non-invasive and widely available; and thus very important.

In our patient we were able to confirm the diagnosis of left submandibular gland aplasia by using ultrasonography and magnetic resonance imaging.

CONCLUSION-
In conclusion unilateral submandibular gland aplasia is an extremely rare disorder and its incidence is unknown. Careful evaluation of patients who complain of neck mass may lead to detection of new cases of this entity. Ultrasonography and magnetic resonance imaging techniques should be performed before any intervention.

In our opinion unilateral agenesis of submandibular gland always will not be associated with hypertrophy of ipsilateral sublingual gland as well as any facial anomaly which was revealed in our case.

REFERENCES