Cheilitis Granulomatosa - An Uncommon Clinicopathological Entity: A Case Report

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Abstract

Granulomatous lesions of the Oral and Oropharyngeal submucosal tissues frequently affecting buccal and labial tissues are uncommon and present a diagnostic dilemma because of the wide variety of possible etiologic factors. The lesion affecting lips initially described by the German dermatologist Miescher as Cheilitis Granulomatosa is a rare disorder characterized by non-remissive enlargement of one or both lips.

The multiple causes and clinical features of Cheilitis Granulomatosa often create a confusing maze through which the clinician must carefully proceed in order to develop an accurate diagnosis and provide an effective treatment. Management considerations for these patients depend upon the results of the investigations, patient’s esthetic considerations and severity of the condition.

This article, besides discussing a successfully treated case of Cheilitis Granulomatosa along with gingival involvement in a 27 year old Indian woman, also highlights the importance of differentiating this condition from other lip swellings.

Keywords: Cheilitis granulomatosa; Systemic steroids; Orofacial granulomatosis

Introduction

Cheilitis Granulomatosa (CG) is a rare granulomatous disorder of unknown origin, initially described by German dermatologist Miescher in 1945 as a distinct clinicopathological entity, characterized clinically by diffuse, non tender, soft to firm swelling of one or both lips. CG has often been associated with other Orofacial Granulomatous disorders e.g. Sarcoidosis, Crohn’s disease, Atypical Tuberculosis, Anderson-Fabry disease and Allergic reaction. It is also considered as an oligo symptomatic or monosymptomatic form of Melkerson Rosenthal syndrome (MRS).

CG usually affects young adults, mostly in the 2nd decade of life with a female predilection (Ceena et al., 2006). The estimated incidence of CG is 0.08% in the population (El-Hakim and Chauvin, 2004). The degree of lip swelling can be considerable and is frequently associated with both vertical fissuring and angular cheilitis. In some cases the labial swelling is associated with hertiform eruptions (Zecha et al., 1976).

Orofacial granulomatosis is a more general term introduced by Wiesenfeld in 1985, which refers to presence of granulomatous inflammation in the oral and facial regions. The oral mucosa is thickened & edematous with the buccal and labial (70%) mucosa assuming a corrugated or lobulated appearance. Occasionally, palatal involvement has the appearance of papillary hyperplasia. A characteristic gingival hyperplasia when present has a patchy distribution & a predilection for anterior region (Sholapurkar et al., 2006).

Case Report

A 27 year old Indian female patient reported to the Department of Oral Medicine with a chief complaint of swelling in upper lip and anterior gums since last 5 months, which was painless and persistent. It appeared during the 2nd trimester of her pregnancy without any remission even after 1 month of parturition and slowly progressed to the present extent. The patient gave history of anterior gums swelling followed by upper lip swelling and the lip was everted and exposed the vestibular mucosa. There was no apparent history of trauma, allergy to any substance, insect bite, pain in the teeth, pus discharge, fever, facial paralysis or any other history of systemic ailment.

Patient consulted a physician 2 months prior to his visit for the same problem, and was prescribed symptomatic systemic medication, but got no relief. She was unclear what medication was given and did not carry any written prescriptions with her. The family history and past dental history of the patient were nonsignificant. She had no deleterious or parafunctional oral habits.

Extraoral examination revealed a diffuse swelling of upper lip causing fullness in the pre-maxillary region with smooth and intact overlying skin, having ill defined margins (Figure 1). Palpation revealed a normal temperature, nontender enlargement which was soft in consistency.

Intraoral examination revealed cracked upper lip with crustations (Figure 1). In upper anterior region, the gingiva was lobulated and enlarged, extending mesio-distally from 13 to 23 regions and superio-inferiorly from labial vestibule to partially covering the crowns with blunting and enlargement of interdental papillae extending palatally.

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in the region of 11 and 21. In the lower anterior region, gingiva was enlarged in relation to 31, 32, and 33 with gingival recession in relation to 41 with apparent inflammation. Enlarged gingiva was reddish pink in color, soft to firm, with loss of stippling & bleeding on probing (Figure 1). The dorsum of the tongue showed normal surface papillae with fissuring.

The chief complaint, history and clinical examination led to a provisional diagnosis of Chronic Idiopathic Granulomatous lesion of the upper lip and gingiva.

The differential diagnosis included Cheilitis Granulomatosa, Angioedema, Cheilitis glandularis, Neurofibroma, Exfoliative cheilitis, Plasm cell cheilitis, Sarcoidosis, Crohn’s disease, Tuberculosis, Lymphangioma, Lymphangioma, Anderson-Fabry disease and Leukemic infiltrate.

**Vitality test** in relation to 11-23 and 31-42 regions was performed and the respective teeth were found to be vital. **Blood investigations** (Hb%, BT, CT, ESR, TLC, DLC), RBS, HbsAg and HIV were in the normal range. **Intraoral periapical radiographs** in relation to 11-21 and 31-42 region revealed mild angular bone loss with intact lamina dura with normal periodontal ligament (PDL) space width (Figure 2) and **Orthopantomograph** revealed generalized mild angular bone loss with no gross bony pathology. **In vitro Hypersensitivity test** was negative. An **Incisional biopsy** of the lip and gingiva was performed under local anaesthesia.

Biopsy specimen revealed stratified squamous parakeratinized epithelium overlying an edematous connective tissue stroma with delicate to dense collagen fibres and few ill-defined areas of noncaseating granulomatous infiltrate consisting of lymphocytes, foamy histiocytes, epitheloid cells and Langerhan’s type of multinucleated giant cells (Figure 3), which were suggestive of chronic granulomatous lesion of the lip.

The histopathological findings of the gingival tissue revealed the aforementioned features along with superficial connective tissue showing edematous areas and few areas of thick collagen fibres which was suggestive of a granulomatous lesion.

The chief complaint, history, clinical examination and subsequent investigations led us in arriving at a final diagnosis of Cheilitis granulomatosa of the upper lip.

Treatment given to the patient consisted of Systemic Prednisolone 10 mg tid for 7 days. Patient reported after 1 week with significant reduction in signs & symptoms of lip swelling. Same treatment was continued for another 14 days with follow up every 1 week. After 3rd week of commencement of treatment, the dose of prednisolone was tapered down gradually and finally withdrawn after 3 months (Figure 4).

Gingiva did not show any remarkable response to systemic glucocorticoid therapy, hence surgical treatment of gingivoplasty was advised (Figure 4).

**Discussion**

The clinical presentation of CG is highly variable. The etiology & pathogenesis of CG is unknown. Hornstein suggested that the disorder is polyetiologic and may be caused by an alteration in autonomic nervous system function localized to facial skin, resulting in increased vascular permeability and edema. Chronic infectious foci, food sensitivities and obstruction of lymphatic vessels have all been suggested as contributing factors (Ceena et al., 2006).
Management of CG is dependent on accurate diagnosis of the condition and identification of any precipitating factors. Patients without dental infections who present with clinical features suggestive of CG should be questioned regarding the presence of systemic signs and symptoms of Crohn’s disease, sarcoidosis, or a history of angioedema. In the presence of positive findings, the patient should be referred for appropriate medical evaluation.

In our case the patient had an extensive lip swelling; hence it was initially treated with systemic glucocorticoids i.e prednisolone for three months with regular follow up and patient got complete regression in signs and symptoms after 3 months.

CG remains an enigmatic disorder with multiple causes. The dental practitioner is likely to encounter patients with this disorder and a proper knowledge of this idiopathic condition is imperative and places a major role in its successful diagnosis and treatment.

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**Table 1: Differential diagnostic possibility for patients with lip swelling (Sholapurkar et al., 2006; Grave et al., 2009; Rogers and Bekic, 1997).**

| Diagnostic considerations | Features not consistent with Cheilitis granulomatosa |
|---------------------------|-----------------------------------------------------|
| Angioedema                | Swelling resolves in 24-48 hrs and recurs again on antigenic stimulation. |
| Cheilitis glandularis     | Inflamed orifices of secretory ducts with red macules on mucosa. |
| Neurofibroma              | Slowly progressive enlargement; pathognomonic and histology. |
| Exfoliative cheilitis     | It is precipitated by lip biting, trauma, along with the presence of epithelial tags. |
| Plasma cell cheilitis     | The classic clinical appearance is a flat-to-slightly raised, eroded plaque or patch, usually on the lower lip of elderly patient. |
| Sarcoidosis               | Characteristic skin lesions; mediastinal involvement; lungs and liver involvement. |
| Crohn’s disease           | Gastrointestinal signs and symptoms; oral ulcerations and fissures. |
| Tuberculosis              | Past history of tuberculosis, and other clinical features of TB. |
| Hemangioma                | Lesions congenital; vascular proliferation. |
| Lymphangioma              | Congenital lesion and characteristic appearance. |
| Anderson-Faty disease     | Characteristic angiookeratomas of skin; history of pain affecting extremities. |
| Leukemic infiltrate       | Histologic features of atypical infiltrate; abnormal peripheral WBCs. |