Traumatic ulcerative granuloma with stromal eosinophilia: A clinical study

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Abstract

Background: Oral ulcers are a common symptom in clinical practice. Among different types of ulcers in oral cavity, traumatic ulcerative granuloma with stromal eosinophilia (TUGSE) appears to be quite neglected by the clinicians due to the limited knowledge and awareness. Traumatic ulcerative granuloma with stromal eosinophilia (TUGSE) is a reactive benign and self-limiting lesion of the oral mucosa. It may mimic squamous cell carcinoma as well as other malignant lesion. The cause is unknown but a traumatic background has been suggested. Oral TUGSE shows regression after a conservative surgical treatment.

Objectives: To analyse the clinical features of 8 TUGSE lesions.

Materials and Methods: 8 cases of TUGSE were retrieved from the files of the Department of Oral medicine and radiology. Their clinical data and histopathological features were examined, collected, and analyzed.

Results: The reported cases included six males and two female patients. Age ranged from 9 to 81 years with mean age of 48 years. The most common site was the tongue followed by buccal mucosa.

Conclusions: Traumatic ulcerative granuloma with stromal eosinophilia (TUGSE) appears to be quite neglected by the clinicians due to the limited knowledge and awareness. It is important to recognise this mucosal entity as it has a close resemblance with malignancy of the oral cavity. Prognosis of TUGSE is normally favourable and aggressive surgical treatment is usually not required.

Keywords: clinical feature; oral cavity; traumatic ulcerative granuloma with stromal eosinophilia, chronic traumatic ulcer

Introduction

Traumatic ulcerative granuloma with stromal eosinophilia (TUGSE) is a rare, benign ulcerative lesion of the oral mucosa. Although the tongue is the most commonly location involved, oral TUGSE lesions can also occur on other oral mucosal sites including buccal mucosa, retromolar area, floor of the mouth, vestibular mucosa, gingiva, and palatal mucosa. Clinically, an oral TUGSE lesion manifests as a slow-healing ulcer with an elevated or rolled border, mimicking a squamous cell carcinoma. The duration of healing ranges from 1 week to 1 year. Delayed healing of TUGSE lesions has been reported to be associated with the lack of secretion of transforming growth factor (TGF)-a and TGF-b by eosinophils infiltrating the lesions. The etiology of TUGSE is still not clear, but traumatic irritation is considered to be the most likely cause. Histologically, TUGSE is characterized as a lesion with surface ulceration and underlying granulomatous tissues showing an inflammatory infiltrate rich in eosinophils. The same histological manifestation in infants is called Riga Fede disease, in which the lesion usually occurs on the ventral surface of tongue and is caused by irritation of erupting mandibular incisors. Oral TUGSE is generally considered to be a reactive lesion. However, CD30+ cells and a monoclonal rearrangement of the T-cell receptor chain gene similar to those found in primary cutaneous CD30+ lymphoproliferative disorders have been reported in part of TUGSE lesions. Therefore, some authors even suspected that TUGSE may be the oral counterpart of primary cutaneous CD30+ lymphoproliferative disorder.
Materials and Methods: 8 cases of TUGSE were retrieved from the files of the Department of Oral medicine and radiology. Demographic data, including the sex and age of patients as well as the location, clinical diagnosis, symptoms and signs, treatment, and recurrence of the lesions were obtained by reviewing the dental and medical charts.

Results
The reported cases included six males and two female patients (Fig. 1).

Fig 1: Gender distribution

Age ranged from 9 to 81 years with mean age of 48 years, with the highest incidence being in the fifth and sixth decades. After reviewing the medical and dental charts, we found that pain was the general symptom and oral ulceration was the universal sign for all oral TUGSE lesions. The most common site was the tongue followed by buccal mucosa. Oral TUGSE lesions were correctly diagnosed in only 4 cases. Marginal induration was palpable in 4 cases (32.4%), and thus the initial clinical diagnosis was “rule out cancer” for these 4 cases. The remaining 4 cases were diagnosed as traumatic ulcer or nonspecific chronic ulcer. Moreover, none of these 8 patients had concomitant cutaneous lesions or developed lymphomas. The lesions were resolved after the biopsy. All patients were followed up for four months, no recurrence had been found.

Discussion
Although the true etiology for oral TUGSE lesion is still unknown, the most possible etiology is traumatic irritation. Sharp tooth margins, ill-fitting dentures, and incisional biopsy for oral leukoplakia have been found as possible causes.1,4 Only one case developed TUGSE lesions after wide surgical excision of their primary oral squamous cell carcinomas, and were suspected as recurrent cancers clinically. However, most reported TUGSE lesions are deficient of obvious clinical causes to confirm the traumatic origin of the lesion. The dysfunction of the immune system may be taken into consideration. A previous study showed a lack of TGF-α and TGF-β production in eosinophils infiltrating in oral TUGSE lesions 6 and this may be why oral TUGSE lesions have a prolonged healing process and are often suspected as malignant lesions 6. In addition to oral squamous cell carcinomas, clinical differential diagnoses of TUGSE lesions may include deep fungal infections, necrotizing bacterial infections, syphilis, and granulomatous disorders due to the slow-healing process and the presence of elevated or rolled border of TUGSE lesions 1-4,8,11 However, oral TUGSE is a self-limiting lesion, and aggressive surgical treatment is usually not required. Moreover, healing of the lesion is usually uneventful after incisional or conservative excisional biopsy of the lesion 1-3. A soft diet and application of corticosteroid or nonsteroidal anti-inflammatory drug ointments are suggested for patients with oral TUGSE to reduce the uncomfortable feelings caused by the lesion. The most common location for oral TUGSE lesions is the tongue 1-3, which was also seen in this study. In addition, the second common location for TUGSE lesions is the buccal mucosa. The high frequency of occurrence of TUGSE lesions on the tongue and buccal mucosa is because the tongue and buccal mucosa are more susceptible to injury by teeth during occlusion or chewing than other oral mucosal sites. The finding that oral TUGSE lesions occurring on the tongue and buccal mucosa also suggests the possible traumatic etiology of oral TUGSE lesions. There was a male predilection for this study, with the male to female ratio of 3:1, which was not comparable to the male to female ratio of 1.38:1 reported by Fonseca et al.,2 which could be due to small sample size in out study. In the series by Elzay et al. of 41 patients with TUGSE (the sex is unknown in one patient), there is an equal number of male and female patients (n =20) 3. However, in the series by Salisbury et al. of 37 patients with TUGSE, a female predilection with the male to female ratio of 1:1.85 is found 1. Fonseca et al. 2 demonstrated no predilection of eosinophils for the muscle fibers surrounding the TUGSE lesion. CD30 is a transmembrane protein of the tumor necrosis factor (TNF) family. CD30 positivity is found in various lesions ranging from reactive conditions to T- or B-cell lymphomas.6,8,12 However, strong and homogeneous CD30 expression in most neoplastic cells is restricted to classic Hodgkin lymphoma, anaplastic large cell lymphomas, and primary cutaneous CD30+ T-cell lymphoproliferative disorders.9 CD30+ cells are also observed in a part of reported TUGSE lesions. Scattered or clustered infiltrates of CD30+ cells were found in 5 of 12 (42%), 11 of 19 (58%), and 26 of 37 TUGSE lesions (70%) reported by Hirshberg et al., 4 Fonseca et al.,2 and Salisbury et al.1 respectively. These CD30+ cells in TUGSE lesions include large mononuclear cells and small reactive T cells.2 Using immunohistochemistry for CD30 protein and polymerase chain reaction (PCR) analysis for the TCR γ chain gene, some atypical TUGSE lesions were found to have CD30+ cells and a monoclonal rearrangement of the TCR γ chain gene.1,3,6 However, negative expression of anaplastic lymphoma kinase protein, a marker associated with large cell CD30+ lymphoma, supports the reactive nature of TUGSE lesion 4. Alobid et al. 6 even found the same pattern of monoclonal rearrangement of the TCR γ gene in specimens of both oral and cutaneous eosinophilic ulcers of the same patient. They thought that TUGSE lesions seem to be the oral counterpart of primary cutaneous CD30+ lymphoproliferative disorders. The large mononuclear cells in TUGSE lesions exhibit irregular nuclear contour, fine chromatin, and abundant cytoplasm.4 They are suspected of heterogeneous origins. As mentioned before, large mononuclear cells have been found to be positive for CD30,2 B-cell marker CD20,27 T-cell marker CD3, 8,10 macrophage marker CD68,11 or Langerhans cell marker CD1a.12 Moreover, Abdel-Naser et al. 10 also discovered the expression of Epstein-Barr virus (EBV) encoded latent membrane protein 1 in CD3-positive
large mononuclear cells. However, this finding is contradictory to the fact that EBV-infected cells are usually of the B-cell lineage. Recurrence of TUGSE lesion has been reported in only two cases with CD30+ cells and a monoclonal rearrangement of the TCR γ chain gene\(^{[18]}\).

**Conclusion**

Traumatic ulcerative granuloma with stromal eosinophilia (TUGSE) appears to be quite neglected by the clinicians due to the limited knowledge and awareness. It is important to recognise this mucosal entity as it has a close resemblance with malignancy of the oral cavity. Prognosis of TUGSE is normally favourable and aggressive surgical treatment is usually not required.

**References**