Case Report

Lingual location of Sweet’s syndrome: A case report

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Abstract – Context: Sweet’s syndrome is a rare acute febrile neutrophilic dermatosis and idiopathic in two-thirds of cases. The pathophysiological mechanisms of Sweet’s syndrome are poorly understood. Observation: A 35 yr old female patient has been referred from the Dermatology Department to the Oral and Surgery Department, University Hospital Center of Oran (Algeria), for a lingual nodule. For medical history, the patient had a Sweet’s syndrome diagnosed 1 yr ago and a rheumatoid arthritis treated with corticosteroids. An excisional biopsy of the lingual nodule was performed and the anatomopathological result revealed a pyogenic granuloma associated with polymorphonuclear neutrophilic vasculitis in the context of Sweet’s syndrome. Discussion and conclusion: Sweet’s syndrome is characterized by a constellation of clinical symptoms, biological and histological abnormalities and is manifested by the sudden appearance of painful skin lesions in the form of asymmetric erythematous papules, nodules or plaques. In the context of Sweet’s syndrome, faced with an oral with cutaneous lesions a correlation and/or manifestation of neutrophilic dermatosis must be suspected.

Introduction

Sweet’s syndrome (SS) or acute febrile neutrophilic dermatosis was first described by Robert Douglas Sweet in 1964 [1,2] as a rare inflammatory disease with predominantly cutaneous expression, belonging to the group of neutrophilic dermatoses [3]. Its incidence is more frequent in women and predominates in middle age. However, it can occur at any age [4]. SS is idiopathic in two thirds of cases [2,5]. However, it can also be associated with myeloid hemopathies, more rarely with solid tumors, certain drugs, autoimmune diseases, auto-inflammatory or infectious diseases or in a paraneoplastic context [2,5,6]. The pathophysiological mechanisms of SS are probably multifactorial but remain poorly elucidated [2,6]. Association with infectious diseases, autoimmune diseases, neoplasias or drugs argues in favor of a hypersensitivity reaction involving cytokines, interferon gamma and TNF for neutrophil activation and recruitment [3]. SS may occur places that have received pressure and trauma [4], these pathogenesis remains a hypothesis described in the literature [3,4]. We reported the case of a patient with simultaneous occurrence of Sweet’s syndrome associated with a rheumatoid arthritis and a lingual nodule. In previous literature, a few cases of oral lesions have been reported in patients with neutrophilic dermatosis and were mainly associated with SS [4,7]. But to our knowledge, no patient with concurrent Sweet’s syndrome, oral pyogenic granuloma associated with neutrophilic polymorphonuclear vasculitis had been reported yet in literature.

Observation

A 35 yr old female patient was referred, in January 2021, to the Department of Oral Pathology and Surgery of the University Hospital Center of Oran (CHUO), Algeria, from the Department of Dermatology of the CHUO, for the therapeutic management of a lingual nodule. The past medical history reveals a cutaneous lesions characterized by edematous and erythematous plaques on the hands, neck and face associated with joint pain. Following the investigations of the attending physicians, diagnosis of SS associated with acute rheumatoid arthritis was made in January 2020. Corticosteroids were prescribed at a dose of 0.5mg/kg which allowed an optimal regression of the cutaneous lesions. The occurrence of the lingual nodule was
synchronous with the skin manifestations that led to the
diagnosis of SS. This painless nodule was located in the anterior
third of the dorsal surface of the tongue on the midline. It was
pedunculated, soft in consistency, covered with erythematous
mucosa and progressive in evolution (Fig. 1). Through the
clinical data and the history of the disease, our diagnosis was
oriented towards a pyogenic granuloma of the tongue, however
an oral manifestation of SS cannot be ruled out. The
preoperative assessment revealed an increase in neutrophils.
Treatment consisted of an excisional biopsy with complete
removal of the lingual nodule followed by hemostasis and
realization of two separate stitches in ‘O’. The surgical specimen
was preserved in formalin for anatomopathological examina-
tion, the result was in favor of a pyogenic granuloma of the
tongue with neutrophil polymorphonuclear vasculitis as part of
sweet’s syndrome (Figs. 2 and 3). The 10-day check-up showed
optimal healing.

Discussion

SS is characterized by a constellation of clinical symptoms,
biological and histological abnormalities that may be
associated with various diseases [4,6]. The typical manifes-
tations of this syndrome are high fever and the sudden
appearance of painful skin lesions in the form of papules,
nodules or asymmetrical erythematous plaques located
preferentially on the upper extremities, of the face and neck
[6]. Involvement of the oral mucosa is rare and less painful
than on the skin [7]. Oral manifestations are superficial
aphtoid lesions (oral mucosa and tongue), bullae and
hemorrhagic vesicles (labial and gingival mucosa), gingival
hyperplasia, ulcerative periodontitis, necrotic nodules (labial
mucosa), papules (macerated: palate and tongue), pustules
(individual and grouped: palate and pharynx), swelling
(tongue), ulcers (palate) [8]. Biological abnormalities include
an accelerated sedimentation rate (ESR) higher than 20 mm in
the first hour, elevation of C-reactive protein (CRP), and
neutrophilic hyperleukocytosis [4]. Histology reveals a diffuse
infiltrate of mature neutrophils typically located in the
superficial dermis [3]. The presence of a vasculitis is no
longer a criterion for eliminating the diagnosis, fibrinoid
necrosis of the vessel wall and leukocytoclastic can be observed
in forms typical of SS. [2]. A skin and/or mucosal biopsy for
routine histopathological evaluation is essential to confirm a
clinically suspected diagnosis of SS [4,7]. The positive
diagnosis is based on clinical, histological and biological
arguments. Two major and 2 minor criteria are required for
diagnosis according to the classification of Su and Liu, modified
by Von den Driesch (Tab. I) [9–11]. The treatment of SS is based
on anti-inflammatory drugs that allow a rapid regression of
symptoms and mucocutaneous lesions [3]. Other treatments
described as effective can be administered depending on the
clinical situation, such as colchicine, dapsone, ciclosporin,
anti-interleukin-1 and anti-tumor necrosis factor [6]. SS can be
associated with other diseases such as solid tumors or
pleomorphic adenoma. In this case surgical intervention is
needed, no recurrence of SS lesions is reported after a complete
excision of tumor [12]. The prognosis of SS varies depending on
the underlying cause because this condition can be associated
with other diseases, including malignancy. Generally, with
timely diagnosis and appropriate treatment, the lesions of SS
resolve without scarring [13]. In our case, according to the
clinical and histopathological data, the diagnosis of pyogenic
granuloma with a neutrophilic polymorphonuclear vasculitis in
the context of SS was retained.

Pyogenic granuloma is defined as a common, acquired,
benign vascular tumor according to International Society for
the Study of Vascular Anomalies (ISSVA) classification 2018
[14–16]. Pyogenic granuloma is a reactive tumorlike lesion that
arises in tissues such as the skin and mucous membranes
(mostly oral cavity) [16]. Clinically, pyogenic granuloma
presents as sessile or pedunculated exophytic mass with a
smooth or lobulated surface which has a tendency to bleed
easily [17]. Various factors (chronic local irritation, trauma,
and hormonal changes) are implicated in the etiopathogenesis
of this entity, but the exact cause is unknown [15].
Histologically, these lesions show an excessive proliferation

Fig. 1. Clinical aspect of the lingual nodule.

Fig. 2. Histological details showing the squamous epithelium and
inflammatory infiltration in the chorion (Magnification ×10).
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of vascular type of connective tissue to a nonspecific infection [17]. The differential diagnosis of oral pyogenic granuloma includes other vascular tumors (hemangioma, lymphangioma) peripheral giant cell granuloma, other malignancies that can mimic pyogenic granuloma (malignant lymphomas, basal cell carcinoma, Kaposi’s sarcoma, angiosarcoma, non-Hodgkin’s lymphoma) [14,15].

In our case, given the clinical criteria of the lingual nodule, we evoked as a differential diagnosis peripheral giant cell granuloma, a benign tumorlike and benign conjunctival tumors (vascular) of the oral mucosa in an isolated context not related to SS. Nevertheless, new discoveries have shed light on the role of inflammatory signaling, disease induction, and relationship with malignancy [18], that’s why we keep in mind anatomo-pathological surprises. Although the appearance of the nodule coincides with the cutaneous manifestations of SS, however, only the histological criteria of SS allow it to be isolated or to be associated with this syndrome. The anatomopathological result of our patient revealing a neutrophilic polymorphonuclear vasculitis associated to pyogenic granuloma classified it as part of SS.

Like our case, SS has been reported to occur in the setting of a variety of autoimmune conditions among which rheumatoid arthritis [18,19]. There are several case reports indicating an association between SS and rheumatoid arthritis. Proinflammatory cytokines are considered to play a vital role in the pathogenesis of both rheumatoid arthritis and SS [19]. It is possible that co-existence of autoimmune conditions and SS is merely coincidental, and that there is no etiologic link between autoimmune disease and SS. However, as may be the case in SS occurring in the setting of infections, a dysregulated immune system in autoimmune states lends itself well to the development of SS and maybe reported to trigger of this syndrome [18]. Moreover, it is possible that the concomitantly occurrence of the nodule in the setting of SS be merely a coincidence. However, in addition, of the fact that SS and pyogenic granuloma share the same trauma pathogenic hypothesis (trauma), mechanistically, it is possible and plausible that SS sets up a pro-inflammatory milieu, that is propitious to the development of this pyogenic granuloma associated with neutrophilic polymorphonuclear vasculitis.

**Table I.** Diagnostic criteria of Sweet’s syndrome [9].

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<th>Classification of diagnostic criteria</th>
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| Major criteria | Sudden appearance of painful erythematous plaques or nodules.  
Typical histology (mostly neutrophilic dermal infiltrate without vasculitis). |
| Minor criteria | Nasopharyngeal prodromes or gastrointestinal infection or vaccination or association with an inflammatory disease, cancer (hematological malignancy, solid tumors), or pregnancy.  
Fever >38 °C and general malaise.  
At the beginning of the eruption: ESR >20 mm/h; increased CPR; leukocytosis >8000 elements/mm³ including >70% polyneutrophils (3 of these 4 data are necessary).  
– Rapid response to treatment with steroids or potassium iodide. |
In the literature and in the context of oral SS, Cohen indicates in his article published in 2003, nodular necrotic lesions especially in the labial mucosa as well as aphthoid and ulcerative lesions especially in the oral mucosa and the tongue. Popular lesions and swelling have been described especially in the tongue area [4]. Kato and al., in 2008, found oral erosions [20], while Fricain and al., in 2015, observed recurrent oral ulcers associated with lingual pustulosis [7]. Bangaru and al., in 2022, reported one case associated with pleomorphic adenoma [12]. In our case, clinically, we found a lingual nodule who’s the histopathological reports associated a pyogenic granuloma with a neutrophilic polymorphonuclear vasculitis correlated to SS (Tab. II). Like Bangaru and al., in 2022, no recurrence of SS lesions of our case is reported after drug therapy based on corticosteroid therapy and a complete excision of tongue tumor.

**Conclusion**

SS has been reported in conjunction with a host of autoimmune and pro inflammatory states; it is unclear, however, whether these associations are due to a shared etiology or are just coincidental occurrences [17]. Although rare, the inaugural symptoms of SS can be found in extracutaneous sites, including the oral mucosa. The diagnosis of SS is based on the presence of clinical, biological, and histological abnormalities showing sterile neutrophilic inflammation in the involved organ and sometimes vasculitis that are no longer a criterion for eliminating the diagnosis can be revealed by anatomopathological exam.

In the context of diagnosed SS, any oral lesion should be correlated with this syndrome until proven otherwise. Faced with an oral lesion with cutaneous lesions, correlation and/or manifestation of neutrophilic dermatosis must be suspected.

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The authors declare that they have no conflict of interest.

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### Authors contributions
All authors contributed equally in the article.

### References