Sweet’s Syndrome: A Classical Presentation of a Rare Disease

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Abstract
Sweet’s syndrome, also known as acute febrile neutrophilic dermatosis, is a rare disorder that typically presents with rapid appearance of tender skin lesions accompanied by fever and leukocytosis with neutrophilia. Its pathogenesis is not fully understood. The syndrome is generally classified into classical, malignancy-associated, and drug-induced categories, each of which has its specific characteristics. In this article, we present a case of classical Sweet’s syndrome in a woman who presented with an acute viral illness.

Keywords
Sweet’s syndrome, neutrophilic dermatoses, immunology

Introduction
Sweet’s syndrome, or acute febrile neutrophilic dermatosis, is a rare inflammatory condition that is characterized by appearance of abrupt painful papulonodular skin lesion in the setting of a prodrome of fever, leukocytosis with neutrophilia, and pathological findings of neutrophilic infiltration of the upper dermis in the absence of leukocytoclastic vasculitis. It is considered to be the major prototype of a subset of diseases known as neutrophilic dermatoses and is generally classified into 3 categories of classical (idiopathic), malignancy-associated, and drug-induced Sweet’s syndrome.1-5 The pathogenesis of Sweet’s syndrome remains unclear; however, the advances since its recognition have established the role of autoinflammatory processes involving both the innate and adaptive immune systems, eventually leading to their malfunction, resulting in immune-mediated hypersensitivity as well as involvement of cytokines such as interleukin-1β (IL-1β), IL-17, and tumor necrosis factor-α (TNF-α).5-10 A diagnostic approach using major and minor criteria is used globally to establish the diagnosis, and skin biopsy and finding of diffuse neutrophilic dermal aggregations in the absence of vasculitis has a pivotal role in making the diagnosis. Systemic corticosteroids remain the cornerstone of treatment strategies; however, other medications have been used as first line as well such as potassium iodide and colchicine.5-9

Case Presentation
A 41-year-old woman with past medical history of insomnia and anxiety presented with fever (as high as 103°F), sore throat, and generalized body pain for 6 days, accompanied by a painful rash involving lower extremities that later progressed to the trunk. During this period she visited the emergency department twice and was diagnosed with a flu-like illness and treated conservatively. However, her symptoms did not improve and she developed swelling of bilateral elbows, wrists, and metacarpophalangeals as well as a watery nonbloody diarrhea for 2 days before admission. On her third presentation to the emergency department she was febrile with a temperature of 39.8°C and appearing ill. She was noted to have symmetrical tender swelling of elbows, wrists, and metacarpophalangeals with decreased active and passive range of motion and dark erythematous, tender, nodular rash in bilateral thighs, abdomen, chest, and back (Figure 1). Her initial laboratory tests were significant for increased erythrocyte sedimentation rate to 85 mm/h and C-reactive protein to 131 mg/L without leukocytosis, neutrophilia, or bandemia. Blood cultures were drawn, and she was started on antibiotics and admitted to general medicine service. On evaluation by the primary team an extensive workup for infectious disease and basic rheumatologic screening was initiated. On consultation with infectious disease service, antibiotics were discontinued and skin biopsy was recommended, which was done the same day. The second day she remained febrile and...
continued to have the presenting symptoms especially the
tender skin lesions without any improvement, and laboratory
tests remained unremarkable without any leukocytosis or
growth in cultures. Therefore, rheumatology service was
consulted and she was started on pulse steroid therapy with
125 mg of intravenous methylprednisolone. On third day her
fevers resolved and the rash and other symptoms started to
rapidly improve. The next day, her infectious workup
returned negative including HIV, monospot, influenza A and
B, hepatitis C virus, hepatitis A virus, hepatitis B virus, chla-
mydia, and gonorrhea. Further the immunological workup
revealed positive anti-neutrophil antibody of 1:80 (RO/SSA
pattern); however, anti-neutrophil cytoplasmic antibody
(myeloperoxidase and proteinase 3), C3, C4, anti-Ds-DNA,
rheumatoid factor, anti-cyclic citrulinated peptide, RNP Ab,
anti-cardiolipin Ab IgM/IgG (immunoglobulin) were nega-
tive or within normal limits (Table 1). She remained afebrile
and her symptoms continued to improve and she was
switched to 40 mg of PO prednisone daily and discharged on
a prednisone taper. Later on her skin biopsy revealed dermal
aggregates of neutrophils (Figure 2), and she was diagnosed
with classical Sweet’s syndrome in the setting of a viral
infection. She was evaluated by oncology and a full workup
was unremarkable for any underlying malignancy. She also
followed up with rheumatology as an outpatient and remained
stable and symptom free.

Discussion

Dr Robert Douglas Sweet first described and used the term
acute febrile neutrophilic dermatosis back in 1964 based on
his observations in 8 women. However, the name “Sweet’s
syndrome” is established as the eponym for acute febrile neu-
trophilic dermatosis and used worldwide. There is no specific
racial or ethnic predominance for Sweet’s syndrome and it is
distributed worldwide.1-5 Sweet’s syndrome is generally clas-
sified into 3 categories based on the underlying etiology and
clinical scenario, including classical (idiopathic) Sweet’s syn-
drome, malignancy-associated Sweet’s syndrome, and drug-
induced Sweet’s syndrome, all of which share the same
presenting scenario of abrupt onset of tender papulonodular
skin lesions, most commonly affecting the face, neck, and
upper extremities with asymmetrical distribution, in the set-
ting of fever and leukocytosis, with histopathological findings
de dense neutrophilic infiltration of the dermis without evi-
dence of vasculitis.1-5 The classical Sweet’s syndrome is the
most common type, which predominantly affects middle-
aged women and is usually associated with an infectious pro-
cess, usually involving the upper respiratory or gastrointestinal
tract, inflammatory bowel disease, or pregnancy.5-8 Patients
most commonly present with fever preceding the skin lesions
that may be accompanied by general malaise, arthralgia,
headache, and other symptoms such as flu-like illness, which
rapidly respond to treatment with corticosteroids. These man-
ifestations are very similar to those of familial Mediterranean
fever; hence, there has been suggestion of possible common
underlying pathophysiology.6,10 Symptoms may recur in up to
one third of the patients, with or without treatment.5-9
Malignancy-associated Sweet’s syndrome was initially con-
sidered a subset of classical Sweet’s syndrome. The symp-
toms of Sweet’s syndrome can present concurrently, precede,
or follow after the presentation of the associated malignancy.
Skin lesion in cases associated with malignancy can be bul-
lous or become ulcerated and resemble those of pyoderma
gangrenosum.5 The most common malignancies associated
with Sweet’s syndrome are hematological malignancies, most
commonly acute myelogenous leukemia. Solid tumors with
carcinomas of the genitourinary tract, breast, and gastrointes-
tinal tract have been reported as well.5-9 Drug-induced form
of Sweet’s syndrome is most commonly observed with gran-
ulocyte-colony stimulating factor, all-trans retinoic acid,
trimethoprim-sulfamethoxazole, and azathioprine.5,8 The diag-
nostic criteria described by Walker and Cohen in 1996 relies
on a temporal relation between administration of a specific
medication and development of the specific symptoms as
well as relapse of the symptoms with re-administration and
resolution with discontinuation.5,8

Sweet’s syndrome is considered a subset of other neutro-
philic dermatoses such as pyoderma gangrenosum and
Behcet’s disease, all of which share the common pathophysi-
ologic characteristics of autoinflammatory processes leading
to neutrophilic infiltrations. The true pathogenesis of Sweet’s
syndrome is unknown as of date and it is believed to be mul-
tifactorial and nonuniform between subtypes of the dis-
ease.5,12 Hypersensitivity is believed to be the underlying
inciting mechanism driving the pathogenesis and the immune
cascades leading to the disease manifestations; however,
scarce evidence has been available to reveal the role of immune complexes and immunoglobulins, making the hypersensitivity hypothesis less strong. Photoinduction and Koebner phenomenon have also been suggested as possible inciting etiologies. Recent advances has led to the understanding that beside the innate immune system, the adaptive immune system also plays a significant role, evidenced by the elevated levels of IL-1α, IL-1β, IL-2, and interferon-γ, which are Type 1 helper T cells (Th1)-related cytokines. Further immunohistochemical examinations of the skin biopsies has revealed decreased Type 2 helper T cells (Th2), which indicates hyperexpression of Th1 cells and increased levels of TNF-α and interferon-γ, which lead to activation and recruitment of neutrophils. The role of proinflammatory T helper 17 (Th17) and secretion of IL-17 in neutrophil activation and recruitment as well as basement membrane remodeling have also been identified in the pathogenesis of Sweet’s syndrome. Moreover, recently there has been identification of certain genes and their roles in pathogenesis of neutrophilic dermatoses including Sweet’s syndrome. Human leukocyte antigen B54 has been associated with Sweet’s syndrome, as well as heterozygous mutations in MEFV gene that is observed in familial Mediterranean fever. These mutations seem to be activating the inflammasome and the innate immune system, leading to IL-1 production and neutrophilic cutaneous inflammation. Nevertheless, a unique pathway to the pathogenesis of Sweet’s syndrome remains to be elucidated. Diagnosis of Sweet’s syndrome was proposed by Su and Liu in 1986 and further modified by von den Driesch in 1994 (Tables 2 and 3). It is based on clinical suspicion and mandates as skin biopsy, and exclusion of other potential differentials including infectious, inflammatory, and neoplastic processes. Management of Sweet’s syndrome lacks a universally accepted guideline; however, corticosteroids remain the cornerstone of first-line treatment, and an excellent response to steroids comprises one of the minor diagnostic criteria. Topical and intralosional steroids can be used for milder forms of the disease as well. Systemic steroids are usually used in doses of 0.5 to 1 mg/kg/day in both oral and intravenous forms, although higher doses up to 2 mg/kg/day have

| Laboratory Tests | Results | Laboratory Tests | Results |
|-----------------|---------|-----------------|---------|
| ESR             | 85 mm/h | HIV Ab/Ag       | Nonreactive |
| CRP             | 131 mg/L| HIV DNA         | Undetectable |
| ANA             | Positive 1:80 (RO/SSA pattern) | Monospot | Negative |
| ANCA            | Negative | Rapid flu A/B | Nonreactive |
| C3              | 176 mg/dL | HCV | Nonreactive |
| C4              | 35.8 mg/dL | HAV IgM | Nonreactive |
| Anti ds-DNA     | Negative | HBC IgM | Nonreactive |
| RF              | <10 IU/mL | HBs Ag | Nonreactive |
| Anti-CCP        | 12 U | Chlamyda trachomatis | Negative |
| RNP Ab          | 0 AU/mL | Neissera gonorrhoeae | Negative |
| ACA IgG         | <9 GPL | β-2-macroglobulin | 2.6 HI |
| ACA IgM         | 11 MPL | β-2-glycoprotein I Ab (IgM, IgG, IgA) | WNL (2, 1, 4) |

Abbreviations: ESR, erythrocyte sedimentation rate; HIV, human immunodeficiency virus; Ab, antibody; Ag, antigen; CRP, C-reactive protein; ANA, anti-neutrophil antibody; ANCA, anti-neutrophil cytoplasmic antibody; C3, complement 3; HCV, Hepatitis C virus; C4, complement 4; HAV, hepatitis A virus; Ig, immunoglobulin; ds-DNA, double-stranded DNA; HBC, hepatitis B core; RF, rheumatoid factor; HBs, hepatitis B surface; CCP, cyclic citrullinated peptide; ACA, anti-cardiolipin Ab.
been used for more severe forms as well.5-8 Steroids are usually tapered in 3 to 5 days and when a desired clinical response is observed. Commonly used first-line treatments consist of potassium iodide (900 mg/day) and colchicine (1.5 mg/day). Second-line treatments are also available such as clofazimine (100-200 mg/day), cyclosporine (2-4 mg/kg/day), dapsone (100-200 mg/day), and indomethacin (50-15 mg/day). Other less used medications such as antibiotics like doxycycline and metronidazole in cases of secondary infections, as well as other antimetabolite agents like cyclophosphamide, methotrexate, and anti-TNF agents such as etanercept and infliximab, have been reported in case reports and small series. Novel approaches to treatment have also been reported such as immunoglobulin and Anakinra (IL-1 receptor antagonist) in refractory cases.5-9 Efficacy of these treatment strategies and the variability in their mechanism of action, and advances in our understanding of neutrophilic dermatoses’ pathophysiology, especially TNF-α, IL-1β, and IL-17, indicates that both innate and adaptive immune systems play a pivotal role in pathogenesis of Sweet’s syndrome. Future research potential lies in further investigation of underlying immunologic signaling pathways, role of genetics in pathogenesis, associations and prognostic value in other autoimmune and myeloproliferative diseases.

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### Ethics Approval

Patient was provided written consent and informed about the purposes of this case report. This case report was performed in accordance with the MedStar Health ethical standards of the institutional and/or national research committee.

### Informed Consent

A signed written informed consent was obtained from the patient.

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### Table 2. Diagnostic Criteria for Classical and Malignancy-Induced Sweet Syndrome

| Major criteria: |
|---|
| 1. Abrupt onset of painful erythematous plaques or nodules. |
| 2. Histopathology evidence of a dense neutrophilic infiltrate without evidence of leukocytoclastic vasculitis. |

| Minor criteria: |
|---|
| 1. Preceded by URT or GI infections or vaccinations; accompanied by a hematologic or visceral malignancies; inflammatory disorders; or pregnancy. |
| 2. Pyrexia >38°C. |
| 3. Leukocytosis >8000, neutrophilia >70%, ESR >20 mm/h, positive CRP (3 out of 4). |
| 4. Excellent response to systemic corticosteroids or potassium iodide. |

Abbreviations: URT, upper respiratory tract; GI, gastrointestinal; ESR, erythrocyte sedimentation rate; CRP, C-reactive protein.

Two major criteria and 2 minor criteria are required to establish the diagnosis.

### Table 3. Diagnostic Criteria for Drug-Induced Sweet’s Syndrome

| Major criteria: |
|---|
| 1. Abrupt onset of painful erythematous plaques or nodules. |
| 2. Histopathologic findings of dense neutrophilic infiltrate without evidence of leukocytoclastic vasculitis. |

| Minor criteria: |
|---|
| 1. Pyrexia >38°C. |
| 2. Temporal relationship between drug ingestion and clinical presentation, or temporally related recurrence after intake. |
| 3. Temporally related resolution of lesions after discontinuation or treatment with systemic corticosteroids. |

All criteria are required to establish the diagnosis.
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