Stevens Johnson Syndrome Initiated by an Adverse Reaction to Trimethoprim-Sulfamethoxazole

Adelina Buganu¹, Massud Atta¹, Matthew Solomon², Paul R. Banerjee³, ⁴, ⁵, Latha Ganti³, ⁵, ⁴

¹. Emergency Medicine, Coliseum Medical Centers, Macon, USA ². Emergency Medicine, Brown University, Providence, USA ³. Emergency Medicine, University of Central Florida, Orlando, USA ⁴. Emergency Medical Services, Polk County Fire Rescue, Bartow, USA ⁵. Emergency Medicine, Envision Physician Services, Plantation, USA

Corresponding author: Latha Ganti, latha.ganti@ucf.edu

Abstract

The authors present a case of a 54-year-old male who presented to the ED with Stevens Johnson syndrome (SJS) beginning on his upper lips, then spreading to his glans penis, airway, and buttocks. After using trimethoprim-sulfamethoxazole (TMP-SMX) to treat a pilonidal cyst diagnosed seven days prior to presentation, the patient began to have desquamating lesions on his upper and lower lips. Subsequently, he noticed desquamation on the glans penis and then between his buttocks. Before being referred to dermatology, he was treated with a high dosage of corticosteroids.

Categories: Dermatology, Emergency Medicine
Keywords: steven johnson syndrome

Introduction

Stevens Johnson syndrome (SJS) is a severe skin disorder that may arise as a reaction from certain medications. A patient suffering from SJS presents a fever, then a red or purple rash that will eventually blister. The blistering portions of the skin usually peel leaving behind a painfully eroded area. SJS can even affect the ears, mucosal surfaces of the mouth, nose, eyes, and airways as well as the genitals and urinary tract. In addition to skin manifestations, patients may develop fevers, myalgias, cough, ptyalism, and dysuria. The skin is a major protective barrier that also helps regulate body temperature with the ability to sweat. SJS is life-threatening because it can severely damage the skin [1].

Stevens Johnson syndrome occurs in 1-2 million people per year [2]. Patients suffering from HIV, Hepatitis A, herpes simplex virus/varicella zoster virus (HSV/VZV) or autoimmune diseases, such as systemic lupus erythematosus, are more likely to experience SJS. Carriers of the gene HLA-B1502 (commonly noted in individuals of southeast Asian, Chinese, and Indian descent) are at an increased risk of developing SJS [3]. Other risk factors include family history of SJS, personal history of SJS, and compromisation of the immune system. SJS is commonly caused by medications such as allopurinol, penicillin, trimethoprim-sulfamethoxazole (TMP-SMX), non-steroidal anti-inflammatory drugs (NSAIDs), and phenytoin among others. In a few reported instances, particularly in pediatric cases, viral infections have been known to cause SJS [1].

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Stevens Johnson syndrome is part of the spectrum of skin reactions. Toxic epidermal necrolysis (TEN) is a similar skin blistering disease. SJS and TEN are merely distinguished by the amount of patient body surface area affected by the skin reaction. According to the common classification of SJS and TEN, SJS affects a body surface area of 10% or less. If a patient has skin manifestations of 50% or more, it is likely to be TEN. An area in between 10% and 30% is considered an SJS-TEN combination [1]. Regardless, both diseases are considered dangerous and emergent.

**Case Presentation**

A 54-year-old male with a previous medical history of hypertension, non-insulin dependent diabetes mellitus, and hyperlipidemia presented to the ED complaining of lip swelling and a rash on his penis. The patient first noticed the swelling on his lip approximately two days prior to presentation. Later, he noticed desquamation of the glans penis. He denied any recent sexual activity and the possibility of a sexually transmitted infection (STI). Further, the patient denied any previous allergic reactions. Approximately seven days prior to presentation, the patient was diagnosed with a pilonidal cyst and was placed on TMP-SMX. He reports adherence with the medication for three days but then he stopped it. The patient restarted the medication the morning his symptoms started.

His physical examination revealed desquamating lesions on his upper and lower lip associated with swelling in his upper lip (Figure 1).

![FIGURE 1: The patient's upper and lower lips show desquamation caused by SJS.](image)

SJS, Stevens Johnson syndrome

There were no buccal or ophthalmic lesions present. He did not appear toxic. His vital signs were normal, including heart and respiratory rate. However, he did report a sensation of airway tightness. Laboratory evaluation was suggestive of underlying inflammation with an elevated C-reactive protein count (1.2 mg/dL) (Table 1).
| Sodium (136-145 mmol/L)         | 136 |
|--------------------------------|-----|
| Potassium (3.5-5.1 mmol/L)     | 4.6 |
| Chloride (99-109 mEq/L)        | 106 |
| Carbon dioxide (22.0-31.0 mEq/L)| 26.0|
| Anion gap (5-15)               | 8.6 |
| BUN (6-18 mg/dL)               | 34 H|
| Creatinine (0.55-1.30 mg/dL)   | 2.0 L|
| Est GFR (African American) (>60)| 42 L|
| Est GFR (Non-African American) (>60)| 35 L|
| Glucose (70-100 mg/dL)         | 229 H|
| Calcium (8.3-10.4 mg/dL)       | 9.5 |
| Total bilirubin (0.0-1.0 mg/dL) | 0.30|
| AST (10-37 unit/L)             | 19  |
| ALT (12-78 unit/L)             | 37  |
| Alkaline phosphatase (30-136 unit/L) | 132 |
| C-Reactive protein (0.05-0.3 mg/dL) | 1.2 H|
| Total protein (6.4-8.2 g/dL)   | 7.1 |
| Albumin (3.4-5.0 g/dL)         | 3.5 |

**TABLE 1: Patient’s laboratory values.**

BUN, blood urea nitrogen; GFR, glomerular filtration rate; AST, aspartate aminotransferase; ALT, alanine transaminase

The patient was admitted for observation due to the potential for airway compromise and treated with a high dose of corticosteroids. He had an uneventful discharge to home. Three days after discharge, he returned to the ED due to persistent symptoms and a new area of desquamation in between his buttocks. He continued with the same treatment of corticosteroids and was subsequently referred to dermatology.

**Discussion**

This patient’s case of SJS exhibited a widespread distribution of desquamation. Lesions first appeared on the patient’s upper and lower lips, though there was no presentation of buccal or ophthalmic lesions. He then experienced desquamation on the glans penis, subsequently spreading to his airway and then buttocks. As typical of SJS, the affected areas were lined with a mucous membrane. Cases of SJS have been also known to affect the urinary tract and the conjunctiva. Studies have reported that 90% of all cases of SJS affect either the mouth, genital,
or gastrointestinal tract [2]. Although not experienced by this patient, it is quite common for SJS patients to experience flu-like symptoms that precede the spread of the disease. Causes, symptoms, and treatments are summarized in Figure 2.
FIGURE 2: Causes, symptoms, and treatments for SJS

STEVENS JOHNSON SYNDROME (SJS)

The Causes, The Symptoms, and The Treatments

A rare, acute disease characterized by skin and mucosal loss

AFFECTS 1-2 MILLION PEOPLE PER YEAR

MACROSCOPIC CAUSES

- Most cases result as a reaction to particular medications (i.e., anticonvulsants, allopurinol, sulfonamides, antibiotics, NSAIDs, nevirapine)
- Few cases result from viral infections, particularly in children.

GENETIC CAUSES

- Particular variations in the human leukocyte antigen (HLA) complex cause cytotoxic T cells and natural killer cells to release granzulin, which destroys one’s skin and mucous membranes.
- Genes HLA B1502 and HLA B1508 are thought to be connected with SJS.
- Studies have shown that people of Han Chinese, South Indian, and Malaysian ancestry that carry the HLA B1502 gene are susceptible to SJS upon taking certain medications.

SYMPTOMS AND DIAGNOSIS

- Flu-like symptoms and skin pain typically precede the spread of SJS. After a few days, blisters and burning eruptions may form, typically beginning on one’s chest and face. The disease can then spread to the mucous membranes (i.e. mouth, anus, and urinary tract). It can also cause irritation in the conjunctiva.
- Skin biopsies are performed if there is uncertainty in the diagnosis.

STANDARD TREATMENTS

- SJS can be most directly treated if the medication or infection causing the disease is identified.
- Pain killers, anesthetic mouth washes, topical corticosteroids, and eyedrops are often used to treat the symptoms of SJS.
- Patients may require fluid replacement or treatment from an ICU or burn unit.

ADDITIONAL DISEASE INFORMATION

SJS vs. TENS

- Toxic Epidermal Necrolysis (TENS) is a much more severe version of SJS.
- SJS covers less than 10% of the body. SJS-TENS overlap covers between 10-30% of the body. TENS covers over 30% of the body.
- While SJS can often be treated with few long-term consequences, TENS can be fatal.
- TENS is less common than SJS with 0.4-1.2 million cases per year.

PREVENTION

- Getting a DNA test and avoiding particular medications if tested positive for the associated genes
- HIV/AIDS, chemotherapy, and lupus patients are at higher risk
- Consider family medical history

POTENTIAL COMPLICATIONS

- Pneumonia
- Cellulitis
- Dry eyes
- Myocarditis
- Vaginal atresia
- Uveitis
- Scarring of the penis
- Corneal ulceration
- Uveitis

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Stevens Johnson syndrome is widely thought to be associated with particular variations in the human leukocyte antigen (HLA) complex causing keratinocyte apoptosis upon exposure to particular medications [1]. In this process, cytotoxic T lymphocytes and natural killer cells release cytotoxic molecules, mainly granulysin, causing epidermal necrosis [4]. Among the most common medications responsible for this reaction is TMP-SMX, which has accounted for about 20% of cases in several studies [5]. In this case, the patient used bactrim for several days to treat his pilonidal cyst. TMP-SMX is a combination of both trimethoprim and sulfamethoxazole, which makes it likely that this patient’s case of SJS was triggered by this medication. Reports indicate that in order to initiate recovery from the disease, the patient must first stop taking the causative medication.

At the ED, the patient was treated with a high dosage of corticosteroids for airway management and continued symptoms and desquamation before it was confirmed that he had SJS. The use of corticosteroids to treat SJS has been controversial. On one hand, prolonged use of corticosteroids is known to increase the chance of secondary infection. However, a retrospective study at Vajira Hospital in Bangkok, Thailand concluded that a limited short-course of corticosteroids can reduce the mortality rate of SJS without inflicting a secondary infection [6]. Current research is also being done on other potential treatments for SJS. Studies have indicated that cyclosporine is an effective immunomodulator due to it specifically targeting granulysin, halting the dissemination of the disease. Early reports also suggested the efficacy of tacrolimus and cyclophosphamide; however, further studies must be conducted to support these results [7].

Conclusions

Stevens Johnson syndrome can widely affect the skin and mucosal regions of the body without preceding symptoms. Physicians must be aware that a given medication used to treat one condition may have the potential to cause SJS. As the disease is commonly genetic, physicians must understand their patient’s family medical history with regard to SJS when prescribing medication. Several treatments are currently being studied for more severe cases of SJS—TENS, though the most essential and basic management is to identify and stop using the causative medication.

Additional Information

Disclosures

Human subjects: Consent was obtained by all participants in this study. HCA Centralized Algorithms for Research Rules on IRB Exemptions (CARRIE)/ IRB manager issued approval 2020-588. HCA Centralized Algorithms for Research Rules on IRB Exemptions (CARRIE)/ IRB manager issued approval 2020-588. Based on the information provided and attested as true, the research plan described does not require IRB oversight. This is because you are either a) not engaging in research with human subjects as defined by federal regulations; b) engaging in research with human subjects deemed excluded from IRB oversight per 45CFR46.102(l) OR c) engaging in research with sufficient human subject protections in the design to meet one or more IRB exemption criteria set forth in 45CFR46.104. Conflicts of interest: In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that no financial support was received from any organization for the submitted work. Financial relationships: All authors have declared that they have no
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