A young woman with abrupt changes in skin and mucosa

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ABSTRACT

Cutaneous and mucosal changes due to adverse drug effects are relatively common in clinical daily practice; however, the etiology often follows unsuspected, undetermined and underreported. More severe lesions like the Stevens-Johnson syndrome should be early recognized and treated. As the entity is potentially lethal if not properly managed, and diagnosis is imminently clinical, the characteristic images might contribute to enhance the suspicion index about this syndrome.

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A 36-year-old Brazilian woman was referred to hospital because of acute onset of skin erythema and purpuric macules, in association with oral, genital, and ocular erosions and bullae, which rapidly involved approximately 10% of the body surface. Her medical antecedents were unremarkable, except for the use of anti-inflammatory pills to control a tooth ache ten days ago. Physical examination showed an acutely ill female, hypotensive, with tachycardia and tachypnea but without fever. Major changes were observed in buccal, perioral, ocular, and genital areas; moreover, skin lesions were detected on the extremities affecting palms and soles (Figure 1). Worthy of note, several skin lesions had the classical appearance of target, iris or bull’s-eye. Laboratory tests revealed mild anemia, neutrophilic leukocytosis and elevated C-reactive protein. The results of microbiological studies, including cultures and serological analysis, were negative. Histopathological study of biopsy sample showed keratinocyte apoptosis, vacuolar changes, sub epidermal blisters, and dermal perivascular lymphocytic infiltrate permeated with eosinophils. The patient was clinically managed in the Intensive Care Unit and her improvement was slow. Her treatment included nutritional support, antibiotics, corticosteroids, and topical care of lesions. The cutaneous and mucosal changes gradually improved and were all healed in about 30 days. After the hospital discharge she remained asymptomatic and without recurrences of the lesions.

What is your diagnosis?

Drug-induced Stevens-Johnson syndrome

The patient herein described had clinical, laboratory and histopathological features consistent with the Stevens-Johnson syndrome (SJS),1-5 first described in 1922 in two patients evaluated in the tenth and the twenty-second days after the onset of illness.1,4 Their eyes were exuding purulent secretion, the lips had clotted blood, and the eruption affected almost all body except for the feet and hands,1 different from the present case. The blisters and desquamation in SJS classically involve mucosae and up to 10% of the body surface.2-5 Toxic epidermal necrolysis (TEN) is a similar entity, which can be differentiated from SJS with base on the extension of over 30% of body surface area affected; whereas the overlap SJS/TEN is characterized by 10 to 30% of body surface involvement.2,4,5 Major causes of SJS are drug reactions and infections caused by Mycoplasma pneumoniae, Chlamydia pneumoniae, cytomegalovirus, human immunodeficiency virus, and disseminated candidia-
Management of patients with SJS includes prevention of secondary infection, fluid and electrolyte balance, nutrition support, glucocorticoids, antibiotics, and immunoglobulins. Systemic complications include respiratory, cardiovascular, gastrointestinal and renal involvement, and mortality rate may be up to 25%. Chantaphakul et al. reviewed 24 patients with SJS in Thailand during a 5-year period; 54.2% were males, with mean age 46.5 (20-77) years, and 33.3% with HIV infection. Oral, ocular, genital, and hepatic changes were observed in 100%, 83.3%, 41.7% and 41.7% of cases, respectively. Most of severe forms were related to adverse drug effects; 65% of cases were treated with systemic steroids during 1 to 10 days, and there was no mortality in this group of patients. Diphoorn et al. reviewed the incidence, drug exposure and mortality data of 59 cases of SJS in the Italian Lombardy region over a period of 5 years and a half. The majority of cases were associated with adverse drug reactions, mainly to allopurinol, and the mortality rate was 16.9%. Martinez-Cabriales et al. reviewed advances in clinical, pathogenesis and management of SJS and reported an annual incidence of 1.2 to 6.0 cases per million inhabitants, the growing incidence with age, and ethnic genetic predisposition. They commented the role of medications and infections by mycoplasma, cytomegalovirus and Dengue virus; and emphasized the need of histopathological confirmation, additionally to clinical features. Yamane et al. reviewed 52 cases of SJS during 2000-2013 in Japan. The mean age of patients was 55.1 years, and 31 were females; 53.85% of cases were due to adverse drug effects. The average interval between the first caustive drug intake and the onset of symptoms of SJS was 18 days, and main complications were pneumonia and sepsis. Before 2007 the mortality rate was 1.9%, and decreased from 4.5% to 0.0% between 2007 and 2013 after addition of immunoglobulins and plasmapheresis to corticotherapy. Qayoom et al. studied adverse cutaneous drug reactions in the Kashmir valley in India during one year. The incidence was 0.16%, the mean age of patients was 39.36±16.77 years with a range of 2-75 years; 50.66% were females, and the main route was oral (86.66%). Fixed drug eruptions occurred in 45.33% and SJS in only 5.33% of the patients. Antimicrobials (57.33%), non-steroidal anti-inflammatory drugs (21.33%), and antiepileptics (17.33%) were the commonest.

The present clinical images aim to enhance the suspicion index of primary care workers about severe drug-induced cutaneous reactions, and contribute to establish the earliest diagnosis.

References