TOXIC EPIDERMAL NECROLYSIS INDUCED BY LANSPORTAZOLE

Necrólise epidérmica tóxica induzida pelo lansoprazol

TAÍNÁ SCALFONI FRACAROLI
JOÃO LUZ SODRÉ
ALEXANDRE GRIPP

LUDMILLA QUEIRÓS MIRANDA
MÁRIO CHAVES

Abstract: Toxic epidermal necrolysis is a rare, severe cutaneous reaction, mostly caused by drugs. It affects the skin and mucous membranes, with involvement of more than 30% of body surface. We describe the case of a young woman, previously healthy, who developed skin detachment of more than 90% of the body surface 15 days after being administered lansoprazole for peptic disease. The treatment consisted in discontinuation of the drug involved and early administration of intravenous human immunoglobulin, which led to a satisfactory outcome of the case, substantiating the impact of early diagnosis and treatment on the morbidity and mortality of these patients.

Keywords: Epidermal necrolysis, toxic; Immunoglobulins, intravenous; Proton pump inhibitors

Resumo: A Necrólise epidérmica tóxica é uma reação cutânea rara, causada principalmente por drogas. Envolve pele e mucosas, com acometimento superior a 30% da superfície corpórea. Ilustramos um caso de uma paciente jovem, previamente hígida, que desenvolveu necrólise epidérmica tóxica, com descolamento superior a 90% da superfície corpórea, 15 dias após o início de lansoprazol para doença peptica. O tratamento consistiu na interrupção do uso da droga implicada e administração precoce de imunoglobulina humana por via venosa, que levou a um desfecho satisfatório do caso, demonstrando o impacto do diagnóstico e tratamento precoces sobre a morbimortalidade destes pacientes.

Palavras-chave: Imunoglobulinas intravenosas; Inibidores da bomba de prótons; Necrólise epidérmica tóxica

INTRODUCTION

Toxic epidermal necrolysis (TEN) is a rare disorder, having significant morbidity and over 30% mortality. It is characterized by extensive apoptosis of keratinocytes, leading to epidermal detachment and mucosal involvement. Stevens-Johnson syndrome (SJS) and TEN represent severe variants of the same process, due to etiopathogenic, clinical and histopathological similarities. These entities differ only in the percentage of body surface involved: a detachment below 10%
represents SJS, 10-30% overlapping of both and above 30% it characterizes TEN.3

The main factors involved in the etiology are drugs, mainly antibiotics, anticonvulsants, oxicam family of nonsteroidal anti-inflammatory drugs and allopurinol. The proton pump inhibitors (lansoprazole and omeprazole) are considered of low risk.4,5 Cases of TEN have been attributed to several new drugs, given that the ones with longer half-life pose higher risk.

The pathogenesis is not fully understood and involves the inability to detoxicate reactive metabolites of drugs, genetic susceptibility and immune factors related to cellular apoptosis. The main pathway of cell death in this case is the interaction of Fas receptor and Fas ligand on the surface of the keratinocyte, since the latter acquires an increased expression on the surface of keratinocytes in TEN.7-8

The clinical signs begins, on average, one week after administration of the drug, and it may range from 7 to 21 days in a first exposure. In a reexposure, the onset happens earlier, and it may happen in 2 days.5,4 It is common that unspecific symptoms such as fever, sore throat, stinging eyes and vagina, precede cutaneous manifestations by a few days. The first lesions tend to happen on the trunk, and are usually erythematous papules or purpuric macules, irregular in shape and size, which tend to coalesce. The progression of the disease, if the offending drug is not removed, occurs from 2 to 5 days or in hours; it rarely takes more than one week. The lesions become greyish red, there is intense necrosis of the whole epidermis, and flaccid blisters are formed, leaving large denuded areas. There is oral mucosa, ocular and genital involvement in more than 90% of the patients with extensive and painful erosions which lead to lip crusts, odynophagia, photophobia, dysuria and painful evacuations.1-3 Systemic manifestations happen due to acute cutaneous failure, which results in water and electrolytic disorder, hypovolemia, renal failure, thermoregulatory unbalance and higher predisposition to sepsis.1

There is a severity score for TEN (SCORTEN), which may be useful to assess the prognosis of these patients, if performed within the first 48 hours of onset (Table 1).7 Factors such as lymphadenopathy, increased transaminasis and neutropenia also imply worse prognosis.1,3 Management in the acute stage involves prompt identification and withdrawal of the culprit drug, support therapy in intensive care unit or burn intensive care and eventual specific drug therapy. Early ophthalmologic evaluation is important to avoid late complications, such as synechia and amaurosis.5,9

Systemic corticosteroids were the main therapy over decades. Currently, their use is controversial, some experts suggest there is increased risk of sepsis and death, while others show that short course of high dose, at the onset, may be beneficial.9,10 The administration of high doses of intravenous immunoglobulin (IVIG) seems to be a promising alternative, as it showed reduction in mortality in some studies. Its probable mechanism of action consists in blocking the Fas receptor-Fas ligand binding, thus inhibiting the apoptosis of keratinocytes. The dose has not been established yet, but the literature shows a higher benefit with the early administration of high doses (2-3 g/kg divided in 3-4 days).1,9,10 Other drugs may be used, such as cyclosporine, cyclophosphamide and TNF-alpha antagonists.4,7

**CASE REPORT**

A previously healthy, 23-year-old woman, began to have ocular and vaginal itching and later symmetrical erythematous macules appeared on the limbs, mainly hands and feet. Within 7 days there was significant spread of lesions, which evolved into blisters. She sought the emergency care unit, where prednisone 2 mg/kg was administered. After 2 days, she was transferred to our service. She reported use of lansoprazole 15 days before the appearance of the lesions.

On examination, the patient was pale 2+/4+, febrile and tachycardic. Epidermal detachment involved more than 90% of the body surface, and there were some areas of erosion (Figures 1 and 2). Mild involvement of the oral mucosa and lips with crust formation. She was admitted to the intensive care unit, lansoprazole was stopped and intravenous immunoglobulin, 2 g/kg was administered for 3 days, in addition to skin debridement and daily dressings. The patient developed significant reepithelialization in 15 days (Figures 3 and 4). After two months, she presents only residual hyperchromic macules.

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**TABLE 1: Score of severity of TEN - SCORTEN**

| Prognostic factors | Points |
|--------------------|--------|
| Age > 40 years     | 1      |
| Tachycardia > 120 bpm | 1     |
| Neoplasia          | 1      |
| Initial detachment > 10% | 1    |
| Serum urea > 10 mmol/L | 1    |
| Serum bicarbonate < 20mmol/L | 1 | 1
| Blood glucose > 14 mmol/L | 1 |

| SCORTEN | Mortality (%) |
|---------|---------------|
| 0-1     | 3             |
| 2       | 12            |
| 3       | 35            |
| 4       | 58            |
| ≥ 5     | 90            |

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DISCUSSION

The proton pump inhibitors are drugs which are little related to the development of TEN, and only 5 cases have been reported in the literature to date. Nonetheless, as they are often used, they may not be considered as cause of pharmacodermia. Our patient showed symptoms 15 days after the administration of lansoprazole and had nonspecific complaints 1 day before the onset of skin lesions, as described in the literature. However, the first lesions were on the extremities and there was palmoplantar involvement, which is not frequent. Although the patient has presented more than 90% of epidermal detachment, there were no severe systemic complications, and the process ended after withdrawal of the culprit drug and immunoglobulin administration.

We believe that SCORTEN is useful, but in this
case it was not used because the patient was admitted nine days after onset of symptoms. Early ophthalmologic evaluation and daily eye care were performed, as ocular synchia is the main long-term sequela. Due to the involvement of the skin and mucous membranes (oral, ocular and genital), an interdisciplinary assessment is important.

The diagnosis is clinical and must not be delayed due to high mortality. It is hard to establish a therapy of choice for TEN, due to the rarity of the disorder and the difficulty of performing controlled clinical trials, but intravenous immunoglobulin is a promising therapeutic option, as there is evidence it may stop the necrosis process.

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MAILING ADDRESS:
Tainá Scalfoni Fracaroli
Avenida 28 de Setembro, 77
Vila Isabel
20551030 Rio de Janeiro, RJ
E-mail: tsfracaroli@yahoo.com.br

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