Pityriasis Lichenoides et Varioliformis Acuta: An Atypical Presentation

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Sir,
Pityriasis lichenoides is a group of inflammatory skin conditions that includes pityriasis lichenoides and varioliformis acuta (PLEVA) and chronic pityriasis lichenoides [1].

Pityriasis lichenoides et varioliformis acuta (PLEVA) is a rare inflammatory dermatosis of the young adults, with a sudden onset and a mostly spontaneous favorable evolution.

Although the etiology of PLEVA is unknown, it is probably an inflammatory reaction to foreign microorganisms such as infectious agents and drugs [2].

Clinically, PLEVA manifests as an acute eruption of small erythematous to brown macules evolving into papules topped by hemorrhagic crusts, by flare-up and remission.

Dermoscopy plays an important orienting role. The lesions are divided in two: early aspect of brownish amorphous zone, white scales, and dotted vessels in periphery. And lesions in late phase with structureless white areas in the center, glomerular and dotted vessels in periphery [3].

Treatment options are not codified; include phototherapy, oral antibiotics, dermocorticoids, and antihistamines [4,5]. Systemic therapies such as corticosteroids, methotrexate, cyclosporine, and retinoids may be warranted in severe manifestations of PLEVA resistant to conventional treatments.

Remission is complete, however, an evolution towards a hematoderma was reported, hence the interest of a prolonged follow-up of the patients.

Herein, we report a case of a patient who presented PLEVA. Mrs R, 51 years old, with a history of arterial hypertension under treatment and vitiligo, presented with a 2 years history of erythematosus papules with fine scaling.

Figure 1: (A) Erythematous papules with fine scaling. (B)(C): Dermoscopy: Wickham striae on erythematous background.
of a generalized rash evolving in flare ups and remission for 2 years. The patient denied any new medications, infection or vaccination. The dermatological physical exam showed multiple rounded umbilical papules, with haemorrhagic crusts, confluent for some in small patches, symmetrically located on the lower limbs, buttocks, abdomen and upper limbs (Figure 1A). The examination of the mucous membranes was normal. The lymph nodes were free, without hepatosplenomegaly, infectious signs or alteration of the general state.

Dermoscopy revealed an erythematous background, and bright whitish structures in favor of wickham’s striae (Figure 1B et 1C).

Skin biopsy showed staggered apoptotic cells with parakeratosis and major lymphocytic exocytosis scattered throughout the epidermis. The dermis was the site of a predominantly lymphocytic infiltrate fanning out towards the epidermis (Figure 2). The immunohistochemical complement excluded any sign of mycosis fungoides.

The findings were consistent with a clinical diagnosis of PLE-VA. A complete remission was noted after 3 weeks with dermocorticoïds.

The originality of our observation lies in the presence of a previously undescribed lichenoid network on dermoscopy. Further studies of dermoscopy of lichenoides pityriasis may be beneficial to increase diagnostic confidence.

Consent: The examination of the patient was conducted according to the Declaration of Helsinki principles.

Conflicts of interest: The authors do not declare any conflict of interest.

References