Pityriasis lichenoid-like mycosis fungoides

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Sir,

Mycosis fungoides is a primary cutaneous T-cell lymphoma, secondary clonal proliferation of mature skin-homing T cells, mostly CD4-positive, with a predilection for involving the epidermis. It is an indolent lymphoma that progresses over several years and represents 50% of primary cutaneous T-cell lymphomas [1]. Its clinical presentation is variable, thus leading to several clinical variants. Herein, we describe a rare variant of mycosis fungoides: pityriasis lichenoid-like mycosis fungoides.

A 45-year-old female was referred to our department with a papular rash evolving for the last year without regression. The patient had a history of breast carcinoma in complete remission for two years. A clinical examination revealed erythematous, scaly, non-itchy papules covering the entire body but sparing the face (Figs. 1 and 2). There was no scalp involvement or associated lymphadenopathy. Based on the clinical presentation, the suggested diagnosis was pityriasis lichenoid. A histological examination revealed Pautrier’s microabscesses, atypical lymphocyte infiltration along the basal layer and papillary dermis, and prominent epidermotropism (Fig. 3). There was pilotropism without mucin. Besides, hyperkeratosis with focal parakeratosis and perivascular infiltrate were noted. An immunohistochemical analysis revealed infiltrates of T cells expressing CD3, CD2, CD5, and a predominance of CD4-positive T cells in the epidermis compared to CD8-positive T cells. CD7 and CD30 were, however, negative. These findings were consistent with pityriasis lichenoid-like mycosis fungoides. The patient was classified as a IB stage and received UVB phototherapy with good progress.

Pityriasis lichenoid-like mycosis fungoides, first described by Ko et al. [2], is a rare form of mycosis fungoides, with only several cases reported in the literature [3]. It differs from the classic form by the presence of erythematous or pigmented, scaly, and non-itchy papules that resemble pityriasis lichenoid, from which it differs by the absence of spontaneous regression of the lesions. Histologically, pityriasis...
lichenoid-like mycosis fungoides is characterized by the association of features of mycosis fungoides (Pautrier’s microabscesses, lymphocytes along basal cells, intraepidermal lymphocytes larger than dermal lymphocytes, haloeed lymphocytes, stuffed lymphocytes in the dermal papilla, coarse collagen bundles in the papillary dermis) and those of pityriasis lichenoid (hyperkeratosis with focal parakeratosis, spongiosis, erythrocyte extravasation, exocytosis of lymphocytes and neutrophils, and, occasionally, the presence of apoptotic keratinocytes). Tumoral lymphocytes are more often CD8+ but may also express CD4+. In many cases, polyclonal CD3 T cells may also be present, and, in some instances, even small CD30+ lymphocytes may be detected [4,5].

In the present case, the clinical presentation, the absence of spontaneous resolution, and the histopathological findings were consistent with the diagnosis of pityriasis lichenoid-like mycosis fungoides, which remains a challenging diagnosis.

Consent

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

The authors certify that they have obtained all appropriate patient consent forms, in which the patients gave their consent for images and other clinical information to be included in the journal. The patients understand that their names and initials will not be published and due effort will be made to conceal their identity, but that anonymity cannot be guaranteed.

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