Ichthyosiform mycosis fungoides: An atypical variant of cutaneous T-cell lymphoma: About a case and literature review

Case Report

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Abstract

Acquired ichthyosis is a known paraneoplastic sign of lymphoproliferative malignancies, with histopathologic findings that are nonspecific, revealing no insinuation of the underlying neoplasm.

Mycosis fungoides (MF) is a skin malignancy of T helper lymphocytes with a wide clinical spectrum. Among the atypical variants of MF, there is an ichthyosis-like presentation. However, to date, rare cases of ichthyosiform MF have been reported. We report here the clinical and histopathological features of patient with IMF.

Keywords

cutaneous T-cell lymphoma; mycosis fungoides; Ichthyosiform variant

Introduction

Mycosis fungoides (MF) is the most common form of cutaneous T-cell lymphoma (CTCL) and is characterized by the clonal expansion of skin-homing T lymphocytes. Various clinical and histological variants of MF have been reported.

Ichthyosiform MF (IMF) is not as well-known as other variants of MF. We report here the clinical and histopathological features of patient with IMF.

Case report

A 56-year-old man had consulted for ichthyosiform eruption (figure 1) of the lower extremities evolving for 2 years associated with maculopapules discreetly pruritic trunk of more recent appearance and polyadenopathy. Cutaneous biopsy revealed Ichthyosiform mycosis fungoides on lesions of ichthyosis and the appearance of a MF classic on the trunk lesions (figure 2). The labeling of lymphocytes was strongly positive for CD3 and CD4 and weakly positive for CD8. There was no histological lymph node involvement. The Thoracoabdominopelvic TDM was normal. The selected diagnosis was that of a stage IIA MF in its classic form on the trunk and the upper limbs and ichthyosiform type on the legs. A methotrexate was indicated in the patient, whose evolution is in progress.

Discussion

Mycosis fungoides (MF) is the most common type of cutaneous T-cell lymphoma (CTCL), and MF is a malignant lymphoma that's characterized by the expansion of a clone of the CD 4+ (or helper) memory T cells that frequently lacks other normal T-cell antigens (CD7). It usually begins as flat patches, which may or may not be histologically diagnosed as MF. Many variations of MF have been described, such as the follicular, granulomatous, hypo- or hyperpigmented and unilesional variants [1,2]. Moreover, ichthyosis-like eruption may very rarely be the first sign of MF. A few cases have recently been reported in which the ichthyosiform eruption proved to be a specific manifestation of MF [3,4]. Ichthyosiform MF differs from acquired ichthyosis...
Ichthyosiform states are rarely associated with cutaneous lymphomas. Isolated, they would often suggest first the paraneoplastic hypothesis. In the particular situation of MF, ichthyosiform eruption may also in rare cases, like that of our patient, correspond to a particular entity: MF ichthyosiform. This new, rare variant of MF is defined by a clinical aspect suggestive of ichthyosis vulgaris and a histology associating the specific appearance of mycosis fungoides and that of ichthyosis vulgaris. Ichthyosiform MF can be divided into 3 types: 1) ichthyosiform eruption as the sole manifestation of the disease, 2) ichthyosiform eruption in conjunction with additional atypical findings of MF and 3) ichthyosiform eruption in combination with the classic types of MF [6]. Marzano et al. and Hodak et al. are the previous studies have reported IMF accompanied by classical MF or follicular MF [7,8], but there have been no reports of IMF accompanied by primary cutaneous anaplastic large cell lymphoma. Our patient had the ichthyosiform eruption in combination with the hyperpigmented types of MF. Ichthyosiform MF has similar histopathologic findingsto classic MF combined with features of ichthyosis, suchas hypogranulosis and hyperkeratosis. The detection of features from both entities in the same biopsy specimen narrows the differential diagnosis down to ichthyosiform MF and paraneoplastic ichthyosis in a patient with known MF.

Conclusion

IMF can be considered as a rare type of early MF with a comparatively favorable prognosis, which is common in young patients. When examining patients with ichthyosiform lesions, careful evaluation including skin biopsy is necessary to rule out the possibility of IMF.

Références