WHAT IS YOUR DIAGNOSIS

DERMATOLOGY CLINICAL CASE

CASO CLÍNICO DERMATOLÓGICO

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A previously healthy nine-year-old male presented with a two-week history of disseminated asymptomatic dermatosis with numerous small erythematous fine-scaling papules on the trunk and limbs, with no apparent trigger (Figures 1 and 2). Examination was otherwise unremarkable. Histopathological exam revealed interface dermatitis with moderate-to-intense perivascular and periadnexal lymphocytic infiltrate, with epidermal exocytosis, obscuring the dermo-epidermal junction (Figure 3).

What is your diagnosis?

Figures 1 and 2 - Child presenting with small erythematous papules with adherent fine scaling, scattered across the trunk and upper limbs

Figure 3 - Lichenoid dermatitis with a lymphocytic infiltrate obscuring the dermo-epidermal junction (hematoxylin-eosin staining, 200x). Right inferior corner: Close look at the intraepidermal involvement with lymphocytic exocytosis (400x)

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DIAGNOSIS

Pityriasis lichenoides et varioliformis acuta

DISCUSSION

In the present clinical case, diagnosis of pityriasis lichenoides et varioliformis acuta was established based on clinicopathological correlation. Topical methylprednisolone aceponate was prescribed, with marked improvement. Sporadic new lesion crops developed, milder than at initial presentation and with prompt response to topical corticosteroid therapy. At six months, the patient presented post-inflammatorty hypopigmented macules.

Pityriasis lichenoides (PL) is an uncommon papulosquamous eruption with acute (pityriasis lichenoides et varioliformis acuta [PLEVA]) and chronic (pityriasis lichenoides chronic [PLC]) presentation forms.\(^1,2\)

Approximately 20% of cases are reported in children.\(^2\) Etiology is uncertain, with hypotheses of an inflammatory response to an infectious agent or other antigenic trigger, a lymphoproliferative disorder, or an immune complex–mediated reaction advocated.\(^1-3\) PLEVA most commonly presents as recurrent crops of small erythematous maculopapules on the trunk and extremities.\(^2\) Diagnosis is clinically and histopathologically established, with histopathology varying according to disease phase.\(^2,3\) Topical corticosteroids, oral erythromycin, phototherapy, topical calcineurin inhibitors, and systemic immunosuppressants are therapeutic options.\(^1-3\) Although PL usually follows a benign course, continued monitoring of these patients is recommended, even in those with spontaneous remission.\(^2\)

LESSONS FROM THIS CLINICAL CASE

PL comprises a spectrum of papulosquamous dermatoses of unclear etiology, with acute and chronic forms.

Diagnosis relies on the combination of characteristic clinical and histopathological features.

Topical corticosteroids are effective and well tolerated as first-line therapy.

ABSTRACT

Here in is reported the case of a child observed due to disseminated papulosquamous eruption, subsequently diagnosed as pityriasis lichenoides et varioliformis acuta. It is an uncommon dermatosis of unclear etiology, presenting as recurrent crops of scattered erythematous and violaceous maculopapules. Several therapeutic options have been proposed. Although clinical course is typically benign, continued monitoring is recommended.

Keywords: diagnosis; pityriasis lichenoides; treatment

RESUMO

É descrito o caso de uma criança observada devido a uma erupção papulosquamosa disseminada, subsequentemente diagnosticada como pitiríase liquenóide aguda. Trata-se de uma dermatose incomum, de etiologia ainda não totalmente esclarecida, que se manifesta por surtos recorrentes de mactulas e pápulas eritemato-violáceas dispersas. Diversas opções terapêuticas têm sido propostas. Embora de evolução geralmente benigna, é recomendada vigilância clínica regular.

Palavras-chave: diagnóstico; pitiríase liquenoide; tratamento

REFERENCES


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