GENERALIZED LICHEN NITIDUS ASSOCIATED WITH DOWN'S SYNDROME - CASE REPORT

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Abstract: Lichen nitidus is a disease of unknown etiology, characterized by flesh-colored, shiny papules of 1-2 mm and generally asymptomatic or with mild pruritus. The most common sites of occurrence are genitalia, upper limbs, trunk and abdomen. The generalized form is rare. This is the fourth reported case of lichen nitidus associated with Down Syndrome.

Keywords: Down syndrome; Lichen nitidus; Lichenoid eruptions; Lichens

INTRODUCTION

Lichen nitidus is a disease of unknown etiology, first described in 1907 by Pinkus. It is characterized by flesh-colored, shiny papules of 1-2 mm. The localized form affects preferentially the genitalia, upper limbs, trunk and abdomen. There is no sex or race predilection, but it is more common among children and young adults. It is generally asymptomatic but discrete itching might occur. The prognosis is favorable with spontaneous resolution in the majority of cases. Koebner phenomenon can be present. There are clinical variations including confluent, vesicular, hemorrhagic, palmoplantar, spinous follicular, perforating, linear and generalized forms. The generalized forms are rare and the association with Down’s syndrome has been reported in the literature, but it is not clearly understood.

CASE REPORT

A 4-year-old, female, black Brazilian patient presented asymptomatic lesions on trunk that appeared three years ago and subsequently became generalized. The patient was born with Down’s syndrome and...
hypothyroidism. On physical examination 1-2 mm hypochromic, shiny papules were distributed on face, trunk, upper and lower limbs and genitalia (Figures 1 and 2). Palmoplantar areas were spared (Figure 3). The histopathological study showed lymphohistiocytic granulomatous infiltrate focally enlarging papillary dermis, limited by lateral epidermal ridges resembling a “claw” or “baseball glove and ball” image (Figures 4 and 5). The clinical-pathological correlation was compatible with lichen nitidus. The patient was treated with moisturizing cream and had discrete improvement after a two-month follow-up.

DISCUSSION

To our knowledge this is the fourth case reported in the literature associating generalized lichen nitidus and Down’s syndrome. The first case was described by Patrizi et al. in 1991 and the subsequent reports were done by Laxmisha & Thappa and Henry & Metry. All reported cases were on children and only one presented pruritus. Koebner phenomenon was present in one case. The proposed treatments, mostly for isolated cases included topical steroids, ammonium lactate lotion, topical immunomodulators and oral cetirizin. In this case we prescribed moisturizing cream because there were no symptoms. Some authors suggest that lichen nitidus is a lichen planus variant. The co-existence of two entities has already been described and in some cases lichen nitidus can progress into lichen planus. However, these entities have distinctive clinical and histological characteristics. Smoller & Flynn demonstrated that the infiltrate was different on lichen nitidus and lichen planus biopsy samples, using immunohistochemical techniques.

The histopathology of lichen nitidus is characteristic, showing lymphohistiocytic granulomatous infiltrate limited to papillary dermis. The epidermal ridges are thickened into dermis involving the infiltrate that resembles the “baseball glove and ball” image.

Cutaneous alterations described in patients with Down’s syndrome include: atopic dermatitis, milia-like calcinosis cutis, elastosis perforans serpiginosum, onychomycosis, tinea corporis, anetoderma, folliculitis, psoriasis, cheilitis, xeroderma, alopecia areata, palmoplantar hyperkeratosis, syringoma, keratosis pilaris, vitiligo, seborrheic dermatitis and livedo. Generalized lichen nitidus has already been described in association with Crohn’s disease, amenorrhea and after treatment with alpha interferon. These reports are sporadic and no relationship between lichen nitidus and systemic disease has been established. The proposed treatments in the literature have
inconstant results, but good outcomes were described with PUVA, UVB narrow band, astemizole, topical and systemic steroids, oral antihistamines and cyclosporin.

Despite these treatments, we considered an expectant observation of asymptomatic cases as a good option, since spontaneous resolution has been reported in the literature. We present this case because of the rare association between Down’s syndrome and generalized lichen nitidus. Further studies are necessary to explain if this association is just a coincidence.

**REFERENCES**


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