Dowling-Degos disease: a rare genodermatosis

Abstract

Dowling-Degos disease (DDD) is a rare, autosomal dominant genodermatosis, characterized by multiple small, reticular pigmented macules distributed in the flexural areas. In the present report, there is a case of the disease that affects 41-year-old female patients, with multiple clinical complications of the disease.

Keywords: genes, female, hyperpigmentation, facial scars, Dowling-Degos disease

Introduction

Dowling-Degos disease (DDD) is a rare genetic disease of the skin (reticulated pigmented anomaly), clinically characterized by flexural brown pigmented reticulate macules, comedo-like papules on the back, neck and pitted perioral or facial scars. The diagnosis is established based on clinical and histopathological correlation. In this paper we report a patient of 41-year-old females, with multiple clinical complications of DDD.

In 1978, Wilson Jones and Grice first described Dowling Disease (DDD). It is rare disease that affects women between the second and fourth decade of life. It presents a late onset with benign evolution; nevertheless, it is extremely unaesthetic. It is further characterized by its clinical picture and histopathology.

Clinically, therefore, it presents with acquired hyper pigmentation of reticulated pattern predominantly in flexures that do not modify or suffer influence of the solar exposition. They are also verified lesions type cribriform scars and perioral acne, with no previous history of acne. In addition, DDD patients with comedo-like lesions on the face, dorsum and other areas mentioned above are observed in patients with DDD.

Histopathology of the skin showed presence of horny cysts and absence of melanocytes hyperplasia. In addition, there was acanthosis characterized by fine and irregular elongation of the inter papillary cones with hyper pigmentation at the extremities. The present case report is about a patient with typical DDD and aims to show the importance of histopathological evaluation, and especially the clinical examination for conclusive diagnosis.

A 43-year-old female, from Campinas- Brazil, black, complained of acne form lesions in the face 30 years ago, and lesions in the armpits and intergluteal region of appearance 3 months ago. At the dermatological examination there were multiple atrophic scars and open comedo on the face, cervical and intermammary regions. There were also erythematous nodules with exit of purulent secretion with open toes in areas of fibrosis and retractions in inframammary regions, armpits and inguinal region. Brownish hyper pigmentation of the reticular pattern located in the infra axillary and intergluteal region was also observed.

Figure 1 Open joints in anterior cervical

The histopathological finding was similar in the two biopsied samples (inframammary and left axilla) and presented extensions of the epithelial cones and hyper pigmentation of the basal. Sometimes the epidermis exhibited foci of atrophy and rectification. Confirming the diagnosis through clinical examination and histopathology. Patient, man, 61 years old, complained of asymptomatic lesions growing asymptomatic dark lesions growing in number and size that started...
20 years ago. He also affirms acne form lesions with spontaneous improvement. Dermatological examination revealed brownish, clustered and confluent macules varying from 2-4mm in folds areas (inguinal region and armpits), intergluteal cleft, posterior thigh root, pelvis and dorsum (Figure 3). It was also observed, atrophic scars in the bilateral and posterior cervical and dorsal. Histopathological study in two points showed in both samples acanthotic thickening of inters papillary ridges with hyper pigmentation of the same, with compatible Dowling Degos.

Hyper pigmentation is symmetrical, asymptomatic and progressively increasing over time, appearing initially in the armpits and in the groin with late extension for the intergluteal folds, cervical and inframammary region and internal face of the arms and thighs.

Comedo-like lesions affecting the dorsal region, armpits and intermammary region, keratoacanthoma, epidermoid cysts, seborrheic keratosis and hidradenitis suppurativa represent the additional characteristics of the presentation of this disease in some patients, which corroborates the existence of a defect in the pilosebaceous epithelial proliferation in these patients.

Regarding histopathology, there was no melanocyte hyperplasia, presence of corneal cysts, with adenoid type seborrheic keratosis as the main histological differential diagnosis. In this, however, there is no infundibulum involvement and papillomatosis. In addition, acanthosis is characterized by fine and irregular elongation of the interpapillary cones with hyper pigmentation at the extremities, which also extends to the follicular infundibulum and there is sporadic follicle blockage. 

It is still necessary to advance in the treatment, all of which still have unsatisfactory results.

Conclusion

These case reports, therefore, have the purpose of exposing that despite the rarity of DDD, this is a disease of easy and quick diagnosis, since it is established simply based on clinical and histopathological correlation. There is no need for another complementary examination for the diagnosis aid.

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None.

Conflicts of interest

The authors declared that there are no conflicts of interest.

References

1. Zimmermann CC, Sforza D, Macedo PM, et al. Dowling-Degos disease: classic clinical and histopathological presentation. An Bras Dermatol. 2011;86(5):959–982.
2. Gontijo B, Pereira LB. O espectro da Doença da Kitamura–Doença de Dowling–Degos: Revisão da literatura e apresentação de dois casos. An Bras Dermatol. 1993;68(2):89–92.
3. Kim YC, Davis MD, Schanbacher CF, et al. Dowling–Degos disease (reticulate pigmented anomaly of the flexures): a clinical and histopathologic study of 6 cases. J Am Acad Dermatol. 1999;40(3):462–467.
4. Betz RC, Planko L, Eigelshoven S, et al. Loss of function mutations in the keratin 5 gene lead to Dowling–Degos disease. Am J Hum Genet. 2006;78(3):510–519.
5. Azulay Abulafia L, Porto JA, Souza MAJ, et al. Dowling Disease. An Bras Dermatol. 1992;67:2758.
6. Nirmal B, Dongre AM, Khopkar US. Dermatoscopic Features of Hyper and Hypopigmented Lesions of Dowling Degos Disease. Indian J Dermatol. 2016;61(1):125.