Familial case of Darier’s disease: A rare genodermatosis

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Abstract:
Darier’s disease, also known as keratosis follicularis is a rare autosomal dominant inherited genodermatosis characterized by abnormal keratinization. A 39-year old non-diabetic, normotensive man presented to dermatology OPD with discretely distributed dark brown greasy, warty, malodorous, hyperkeratotic papules over his whole body for last 15 years. He has two young daughters and both has started initial stage of similar lesions for last 6 months. Characteristic v-shaped nicking at the tip of the nails with longitudinal red & white ridges were seen. Nails at the distal ends were broken. Biopsy of skin for histopathological examination showed more specific and confirmatory ‘corps ronds and grains’. This article reports a case of familial darier’s disease. Good improvement was noticed after acitretin therapy.

Key words: Darier’s disease, Autosomal dominant, Genodermatosis, Acitretin

Introduction
Darier’s disease is a genetic, inherited skin disorder characterized by brown keratotic papules that tend to coalesce into patches in a seborrheic distribution. Early lesions are small, firm papules, later covered with a greasy, gray-brown crust. Over years, the papules grow and may fuse to form malodorous, papillomatous, vegetating growths. Although this is a genetically transmitted disease, according to a larger series about 47% had no clear family history presumably because of incomplete penetrance and variable expressivity. The disease is caused by mutations in the ATP 2A2 gene, which encodes the sarco/endoplasmic reticulum Ca²⁺ ATPase isoform. The prevalence of this disorder in population is 1:100000. The sex incidence is equal, although the males appear to be more severely affected than females. Here we are reporting a case of familial Darier’s disease affecting father and daughters.

Case report
A 39 year old man presented with eruptions on the body for 15 years, with malodor from same areas for 5-6 years and has been under topical and herbal medications intermittently but with no improvement. Initially, he developed pruritic dark brown, discrete papules with a greasy texture distributed over the face, neck and chest. Gradually the lesions spreaded all over the body forming warty, vegetating growths and infected during summer. His nails showed v-shaped triangular nicking of the free edges with longitudinal alternating white and red nail lines. There were no associated systemic complaints. He has two young daughters (6 and 4 years respectively) and both has started initial stage of similar lesions for last 6 months. His hair and mucous membrane examinations were normal.

Fig 1: showing multiple dark brown, greasy keratotic papular lesions on face, trunk and upper limb, some are coalescing into plaques on face.
Considering signs and symptoms, we considered differential diagnosis of Darier’s disease, Hailey-Hailey disease, Bazex syndrome, ichthyosiform dermatosis, Langerhans cell histiocytosis and severe atopic dermatitis. General examination and other systemic examination revealed no abnormality. His CBC, LFT, blood sugar, renal, thyroid all were within normal limit. Histological examination showed suprabasal clefts in the epithelium with acantholytic and dyskeratotic cells represented by “corps ronds and grains”.

Biopsy report is confirmatory of Darier’s disease. So, our final diagnosis is familial Darier’s disease. Patient was treated with systemic retinoid (Tab acitretin 25 mg/daily), antihistamine, topical moisturizer and steroid cream. After 1 month of treatment patient came for follow up. Good improvement was noticed.

**Discussion**

Darier’s disease is a rare keratinization disorder with skin involvement and relatively subtle oral mucosal lesions. In Darier’s disease abnormal dissolution of desmosomal plaque proteins is seen in electron microscopy. Calcium ion (Ca++) dependent cell-cell adhesion molecules (epithelial cadherins) are greatly reduced on the acantholytic cells of patients of Darier’s disease contributing to acantholysis.6

Darier’s disease is usually manifested during childhood or adolescence. But the full picture is seen in the 4-5th decade of life and our case also belongs to this category. Numerous erythematous pruritic small, firm papules appear first on seborrheic areas, later on become grayish brown, ulcerates and gets crusted. Foul odor may also be present as a result of secondary infection. Palmer and planter keratosis may be
present with nail changes 7. The frequency of oral lesions range from 15% to 50% and is present on the palate showing a cobblestone appearance 8,9. The present case showed skin lesions and nail abnormality without mucosal involvement. Precipitating factors were heat and humidity, mechanical trauma like friction, sunlight and secondary bacterial infections. In the literature, associated anomalies have been described including mental retardation and psychosis 10. In our patients such anomalies were absent.

Darier’s disease is more associated with cosmetic and esthetic than functional implications since this is a benign dermatosis. But depending on severity of the disease, the emotional status may be damaged by esthetic reasons. The systemic treatment of the Darier’s disease is symptomatic. The lesions relapse because of hereditary etiopathogenesis, especially in patients with severe and generalized form of disease.

Conclusion

Most patients with severe Darier’s disease should receive genetic counseling, including information of inherited condition and risk of transmission to offspring. Biopsy is necessary to arrive at definitive diagnosis. Dental and psychiatric opinion should follow in severe cases to prevent possible complications. So multidisciplinary approach is needed for better management.

References: