Darier White disease

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Abstract
Darier’s disease was described independently by Darier and White in 1989 AD. Reported prevalence varies from 1/100,000 in Denmark to 1 in 30-35,000 in northern England and Scotland.\(^1\) Darier’s disease has been reported from other parts of the world, however, there has been no reported cases in Nepal.

It is an inherited autosomal dominat disease characterized by dark crusty lesions over the seborrheic areas of the skin. We are reporting a case of Darier’s disease in a Nepalese patient who developed skin lesions over his face and body at the age of 12 years. We took skin punch biopsy from the lesions, which showed findings suggestive of Darier’s disease. Our case was treated with isotretinoin and topical keratolytics. He responded with the given treatment within two weeks of therapy.

Introduction
Darier’s disease is an autosomal dominant condition characterized by a persistent eruption of hyperkeratotic papules, histological examination of which shows suprabasal acantholysis with a distinctive overlying dyskeratosis. Sporadic cases are common. It has worldwide distribution.\(^1\)

Case Report
A 27 years old gentleman presented with yellow-brown greasy crusted lesions over his face, chest, and back and buttocks. He noticed the lesions for the first time at the age of 12 years. Initially the lesions were few and was present over the forehead only.\(^2\) later it progressed over to the face, retroauricular areas and chest. The lesions were asymptomatic but was often complicated by secondary infections.

He visited various health facilities for this problem, where secondary infection was treated, but the lesions did not disappeared.

On examination firm rough papules, which was yellow-brown to skin coloured, it was present on the forehead, scalp, temporal regions and margin of the scalp, nasolabial folds, ears, chest, back and buttocks (Figure 1). There were large warty lesions on behind the pinna as well. Nail changes revealed red and white longitudinal bands of varying width with pathognomonic notch at the free margin of the nail (Figure 2). However, there were no lesions on the mucosal layers, palm and soles. Laboratory evaluation of renal and liver function tests, hematology, lipid profile and random blood sugar were within the normal limits. A skin punch biopsy was taken from a lesion over the chest. Histopathological examination revealed areas of hyperkeratosis and suprabasal clefts with irregular upward proliferation of papillae, which
were lined by single layer of basal cells (Figure 3). The histopathology confirmed the diagnosis of Darier’s disease. The Patient was treated with isotretinoin 20mg OD, omeprazol 20mg OD, 5% salicylic acid& 3% coal tar shampoo, sunscreen and azithromycin for 3 days (for secondary infection). Within two weeks of therapy, the lesions were flattened and crusted lesions were reduced. Renal Function Tests, Liver Function Tests and lipid profile was done regularly every month.

Discussion

Though Darier’s disease an inherent autosomal dominant disease, in 47% cases, there is no clear family history. These patients may have sporadic disease or family member with milder form of disease.

The first skin lesions are present during teenage and are pruritic. They are exacerbated by heat, sweat, humidity, sunlight, UVB exposure, lithium, oral corticosteroids, and mechanical trauma.

Darier’s disease is caused by the disruption of desmosomal junction, which allows entry of adhesion proteins into the acantholytic cells. The aetiology of darier disease is though to be due to mutations in the ATP2A2 gene at chromosome 12q24.1, which encodes the sarco/endoplasmic reticulum calcium ATPase type 2 (SERCA2). SERCA2 is a member of a family of ion pumps, which maintain high calcium concentration in the endoplasmic reticulum.

Many patients with mild disease require no treatment other than emollients, simple hygiene and advice to avoid sunburn. Topical tretinoin and isotretinoin, adapalene and tazarotene have been reported as effective but irritation is limiting factor and effect can be seen only in 3 months. Antiseptics may help infected plaques, which may respond to topical steroid/antibiotic combinations. For those with more severe disease, oral retinoids are usually effective; both acitretin and isotretinoin are used. Severe inflammatory exacerbations of Darier’s disease occur in some patients, and may respond to cyclosporin; The use of biological agents has not been reported. Dermabrasion or laser treatment may prove useful, usually in limited areas. Benefit from photodynamic therapy, and with topical 5-fluorouracil in two patients has been reported.

Amongst newer modalities, laser ablation of recalcitrant plaques has been reported in seven keratosis follicularis (Darier disease) patients. Of these, three patients were treated with carbon dioxide lasers with Er:YAG lasers and two with pulsed-dye lasers. In all of these cases, treatment was successful, with only one patient developing disease recurrence in her axilla in seven months after treatment. Another report describes resolution of disease using 1550-nm erbium-doped fractional fiber laser.

Carbon dioxide laser ablation with adjunctive dermabrasion, curettage, and shave excision in various combinations has also been reported to cause disease remission for 8 months to 2 years.

Photodynamic therapy with 5-aminolevulinic acid was used to treat keratosis follicularis (Darier disease)
lesions in 6 patients, with 4 patients showing sustained improvement or clearance for a follow-up period of 6 months to 3 years.\textsuperscript{11, 12}

Surgical excision of hypertrophic intertriginous keratosis follicularis has been described in one case report.\textsuperscript{13}

**Conclusion**

In our case we started Isotretinoin, he showed marked improvement, so we continued the treatment with regular follow up.

Darier’s disease is genetic disease with mild to severe presentation. Depending upon the severity of the presentation we can treat this condition with medication and other modalities like dermaabrasion or lasers.

**References**


