A Case of Familial Comedonal Darier’s Disease

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INTRODUCTION

Darier’s disease is a genetic disorder of keratinization with autosomal dominant inheritance. Its appearance is usually in the form of greasy, crusted, keratotic yellow-brown papules and plaques found particularly on seborrheic areas of the body. However, there are some clinical variants showing atypical skin lesions. Here we report an unusual case of Darier’s disease, which mainly showed prominent comedonal papules over the face. (Ann Dermatol 23(S3) S398 ~ S401, 2011)

Keywords: Comedonal type, Darier disease, Keratosis follicularis

CASE REPORT

A 31-year-old female presented with asymptomatic multiple skin colored or erythematous greasy papules on the face, which developed in her late 20s. She was otherwise well, without any relevant previous history of skin or medical disease. She said that her relatives on the maternal side, including her grandfather, mother, uncle, two aunts, and two brothers, had the same skin lesions. She had never been treated for her skin problem before. Physical examination revealed multiple skin colored or cornifying, zosteriform or linear, acute, and comedonal subtypes.

Here we report an unusual case of Darier’s disease, which mainly showed prominent comedonal papules over the face.

Fig. 1. Multiple skin colored or somewhat erythematous papules on the forehead, eyelids and areas along paranasal and nasolabial folds. The facial skin looked greasy (the arrow indicates the biopsy site).
erythematous papules on the forehead, eyelids, and areas along paranasal and nasolabial folds. Clinically, these papules looked like trichoepithelioma or syringoma, or comedones of acne vulgaris. Her facial skin was greasy (Fig. 1). Otherwise, the scalp, trunk, palms, soles, nails, oral mucosa and teeth were normal.

Skin histopathology of facial papules showed dilated follicular infundibulum, containing keratotic materials with parakeratotic cells. At the lateral aspect and base of the follicular infundibulum, suprabasal acantholysis led to formation of clefts, lacunae, and villi. Dyskeratotic cells, including corp ronds and grains, were observed, as well as brisk perifollicular inflammatory cellular infiltration (Fig. 2).

According to her clinical and histopathological features, we finally diagnosed comedonal Darier’s disease. We treated her with oral minocycline and topical tacrolimus ointment; however, her condition showed little improvement and she refused further treatment.

**DISCUSSION**

Darier’s disease, also known as keratosis follicularis, dyskeratosis follicularis or benign dyskeratosis, is a rare disorder of keratinization that primarily affects the skin. It was described independently by both Darier and White in 1889. It has a prevalence of 1:100,000 of the population and is inherited as an autosomal dominant trait. It is characterized by follicular and extrafollicular greasy hyperkeratotic papules and plaques, arising primarily in seborrheic areas. Vegetating papules, erosions, or hemorrhagic blisters may sometimes be seen. Palms, soles and the oral cavity may be affected. In Darier’s disease, there are clinical variants, including hypertrophic, vesicobullous, hypopigmented, cornifying, zosteriform or linear, acute, and comedonal subtypes, similar to our case. Major histopathologic findings include the following: 1) dyskeratosis resulting in formation of corps ronds and grains 2) suprabasal acantholysis, leading to formation of suprabasal cleft or lacunae and 3) villi, which are diagnostic with typical clinical findings.
Table 1. Reported cases of comedonal Darier's disease

| Case          | Age/Sex | Sites               | Onset (age) | F/Hx. | Features of Darier's disease | Tx.                  | Response of Tx. |
|---------------|---------|---------------------|-------------|-------|-----------------------------|----------------------|-----------------|
| Derrik et al. | 65/M    | Face, scalp         | 61          | -     | Nail, palm                  | Topical steroid      |                 |
|               |         |                     |             |       |                             | → Simple emollients  |                 |
| Derrik et al. | 55/M    | Face, trunk         | 10s         | +     | Nail, palm                  | Etretinate 50 mg/day | Improved        |
| Song et al.   | 43/M    | Face, scalp, trunk  | 16          | -     | Nail                        | Etretinate 30 mg/day | Improved        |
|               |         |                     |             |       |                             | for 6 mo.            |                 |
| Lee et al.    | 47/M    | Face, trunk         | 25          | -     |                             | Isotretinoin 30 mg/day | Improved for 4 wks |
|               |         |                     |             |       |                             |                      |                 |
| Lee et al.    | 34/M    | Face                | 24          | -     |                             | Topical adapalene   |                 |
| Aliağaoglu et al. | 42/M (Turkey) | Face, back, legs | 18          | -     |                             | + intralesional steroid |                 |
| Ours          | 31/F    | Face                | Late 20s    | +     |                             | Minocycline         |                 |
|               |         |                     |             |       |                             | + topical tacrolimus |                 |

F/Hx.: family history, Tx.: treatment, M: male, F: female.

Previously reported comedonal subtypes presented either as nodular lesions or comedonal lesions, like multiple large blackheads on the face, scalp and the upper trunk. Due to prominent follicular involvement and presence of elongated dermal villi and papillary projections, the histopathological appearance of comedonal Darier's disease differs from that of typical lesions of Darier's disease. The mechanism of comedone formation in this disease is unclear, but both follicular involvement and dilatation seem to be responsible. Diseases like acne vulgaris, trichoepithelioma, warty dyskeratosis, familial follicular dyskeratosis, and familial dyskeratotic comedones should be differentiated from comedonal Darier's disease. In particular, familial dyskeratotic comedones have very similar features with comedonal Darier's disease, but they present multiple large comedonal papules with a central keratotic plug on the forearms and thighs, whereas the face, scalp and mouth tend to be spared. In addition, corps ronds, lacunae and villi are less prominent in familial dyskeratotic comedones.

The clinical course of comedonal Darier's disease is often unpredictable and management may be challenging. Treatment with emollients, topical retinoids and topical steroids usually showed limited benefits. Recently, topical tacrolimus has also been tried. Systemic retinoids are often relatively effective, but are toxic. We recommended vitamin A derivatives, but our patient refused it, because she was planning on becoming pregnant. In some uncontrolled studies for flexural and/or hypertrophic involvement, surgical or physically destructive treatments have been used. These include excision, electrodesiccation, dermabrasion, carbon dioxide, and erbium: yttrium-aluminium-argon laser ablation.

Comedonal Darier's disease is very rare. Our patient showed scattered multiple papules resembling trichoepithelioma or syringoma on greasy facial skin and, of particular interest, she had a family history of the same skin disease. We could find only six cases of comedonal Darier's disease previously reported in the English literature (Table 1). However, only one of them had a family history. Therefore, herein, we report our interesting case of prominent comedonal Darier's disease with family history, as a very rare one.

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