Type2 Segmental Manifestation Of Disseminated Superficial Actinic Porokeratosis In An 8 Year Old Girl

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Abstract

Porokeratosis is a form of genodermatosis of different clinical types. The combination of different types of porokeratosis in one individual is rare. Here we report a rare case in which two different types of porokeratosis linear and disseminated type coexisted. An 8 year old girl developed a centrifugal lesion on her right chest at the age of 2 years, scattered lesions on her both cheeks at the age of 3 after ultraviolet exposure and linear porokeratosis on her left side of body at the age of 5. No other family member and siblings had similar skin eruptions. Histopathology revealed cornoid lamella, dyskeratosis in the epidermis and scanty dermal perivascular lymphocytic infiltration. We diagnosed her facial lesions as disseminated superficial porokeratosis and linear porokeratosis on left side of body. The combination of two varieties may be a result of the loss of heterozygosity. We consider that this case may represent segmental manifestation of disseminated superficial actinic porokeratosis.

Discussion

Porokeratosis is a clonal disease characterized by keratotic abnormalities with autosomal dominant inheritance. Each type of porokeratosis has distinct clinical features and distribution. Nevertheless, all
these types are associated with cornoid lamellae as a common histological feature [1].

Here, we described a case of porokeratosis in which two different clinical types of porokeratosis existed together in a girl. We diagnosed her facial eruption as DSAP because the small lesions were distributed symmetrically on both cheeks, did not follow the lines of Blashko and appeared after UV exposure at the age of 3. On the other hand we are certain about the LP diagnosis concerning the eruptions on the left and right side of her body surface because of the linear distribution of the eruptions along the lines of Blashko. Happle proposed the loss of heterozygosity caused by somatic recombination as an explanation for this association [6]. In the case of an autosomal dominant trait, at the early stage of embryogenesis, somatic crossing-over, nondisjunction or deletion may occur involving this gene locus and this may cause the loss of heterozygosity. In our patient the combination of LP and DSAP may be a result of the loss of heterozygosity. It is possible that our patient is susceptible to ultraviolet light for some reason and DSAP appeared first after strong ultraviolet exposure at the age of 3.

Various treatments have been applied for porokeratosis, such as keratolytic agents, topical or intralesional corticosteroids, topical tretinoins, topical 5-flurouracil, systemic retinoids, cryotherapy, electrodessication, CO2 laser or derma abrasion. However the results were unsatisfactory. We tried tretinoin and cryotherapy with moderate effect. However long term follow up is necessary to determine the efficacy of these treatments.

References

Illustrations

Illustration 1

Figure 1: centrifugal lesions on right chest and scattered lesions on both cheeks.

Illustration 2

Figure 2: numerous small keratotic lesions on her left hand in a linear fashion
Illustration 3

Figure 3: numerous small keratotic lesions on her left leg in a linear fashion

Illustration 4

Figure 4: cornoid lamella and dyskeratosis in the epidermis.
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