Case Report
Porokeratotic Eccrine Ostial and Dermal Duct Nevus

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PEODDN is a rare benign cutaneous disorder that clinically resembles comedo nevus but favors the palms and soles, where pilosebaceous follicles are absent. Widespread involvement along Blaschko’s lines can also occur. It is a disorder of keratinization involving the intraepidermal eccrine duct (acrosyringium) and is characterized by eccrine hamartoma and cornoid lamellation in pathology. The patient is a 29-year-old man with an 8-year history of pruritic skin lesions on his right lateral ankle. In the pathologic examination, multiple small epidermal invagination with overlying parakeratotic cornoid lamellation, loss of granular layer, and few dyskeratotic cells at the base of epidermal invagination are revealed. After clinic-pathologic correlation, the diagnosis of porokeratotic eccrine ostial and dermal duct nevus (PEODDN) was made. Late-onset and rare clinical presentation as pruritic lesion are the characteristic features that make this patient an extraordinary presentation of PEODDN.

1. Introduction

The term Porokeratotic Eccrine Ostial and Dermal Duct Nevus (PEODDN) was first described in 1980 by Abell and Read [1]. However, it was first described by Marsden et al. in 1979 [2]. PEODDN is a very rare skin condition which is classified as porokeratotic dermatoses. It is characterized by cornoid lamella which is a column of parakeratotic cells and is associated with dyskeratosis in the spinous layer as well as reduction in the number of granular zone cells. It is in close association with subjacent acrosyringia [3]. Here, we report a case with this rare condition.

2. Case Report

A 29-year-old Iranian gentleman was referred to the Dermatology Department of RAZI Skin Hospital with severe pruritic skin lesions on his right lateral ankle that had presented since 8 years ago. Physical examination revealed multiple keratotic papules of similar size in a linear distribution forming verrucous plaques over the lower part of his right lateral ankle (Figure 1).

One of the papules was removed by punch biopsy and examined under the microscope. Pathologic evaluation revealed multiple small epidermal invagination with overlying parakeratotic cornoid lamellation and underlying, slightly tortuous, eccrine duct nearby the epidermis. Loss of granular layer and few dyskeratotic cells were evident at the base of epidermal invagination (Figure 2).

3. Discussion

POEDDN is firstly reported by Marsden et al. as a comedo nevus of the palm in 1979 [2]. According to our best knowledge, there are about 41 case reports and 6 literature reviews about that all around the world. Many of them present at birth or at young ages although some may occur in adults or even in the elderly [3]. It is usually asymptomatic, although it may be accompanied by a mild pruritus, hyperhidrosis, or anhidrosis. Association with other conditions is rare and includes neurological problems, scoliosis, palmoplantar keratoderma, onychodysplasia, alopecia, and hyperthyroidism [4–6].

Etiologically, it has been proposed that the invagination of the epidermis may result from an abnormal clone of epidermal cells which leads to the formation of cornoid lamella [7]. Another hypothesis suggests that the invagination is a dilated acrosyringeal and dermal duct which is keratin-plugged [8]. It is also supposed that Porokeratotic Eccrine Nevus may be caused by somatic connexin 26 mutations [9].
Histopathology is the mainstay of diagnosis; cornoid lamella with the involvement of acrosyringia is pathognomonic for PEODDN. It is usually associated with the dilation of eccrine duct. Differential diagnoses include porokeratosis plantaris discreta, inflammatory linear verrucous epidermal nevus, nevus comedonicus, linear epidermal nevus, linear psoriasis, spiny keratoderma, linear porokeratosis, congenital unilateral punctate porokeratosis, and porokeratosis of Mibelli [10, 11].

In conclusion, although PEODDN is often an early-onset and asymptomatic or mildly pruritic lesion, we describe a case of late-onset PEODDN with severe pruritus on his right lateral ankle. Because of the rarity of this condition, each diagnosed case of PEODDN should be reported to enhance our knowledge regarding this condition. We would like to emphasize that late-onset pruritic lesions with blaschkoid distribution could be one of the clinical presentations of PEODDN.

**Conflict of Interests**

The authors declare that there is no conflict of interests.

**References**


