Disseminated Superficial Actinic Porokeratosis on the Face Treated with Imiquimod 5% Cream

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Key Words
Porokeratosis · Facial lesions · Imiquimod cream

Abstract
Porokeratoses are a group of hereditary or acquired disorders characterized by annular lesions with an atrophic center and a prominent peripheral ridge. Pathologically, porokeratosis is characterized by the presence of abnormal clones of keratinocytes that form a column of parakeratotic cells, called the cornoid lamella. Fifteen percent of patients of disseminated superficial actinic porokeratosis (DSAP) have facial lesions; other regions like the extensor surface of extremities constitute the majority of reported cases. Exclusively facial lesions are probably less frequent. Rarely, actinic porokeratosis is confined to the nose only. Of reported porokeratosis cases, 7.5% have revealed a malignancy arising within the lesion. We present a case of facial sporadic DSAP that was treated with imiquimod 5% cream in conjunction with a regular sunscreen. Follow-up of this case is important to rule out the possibility of malignant transformation of the lesions.
Case Report

A 19-year-old, single, unemployed, Caucasian female (skin type III) presented with multiple facial lesions. She had been suffering from multiple pigmented annular papules and plaques on her face for 7 years. The lesions were slowly progressive with no signs of self-healing. She had been a normal, healthy, full-term baby and had normal physical milestones of development as a child. She appeared to be immunocompetent with no history of recurrent infections or any other skin lesions. Her parents and siblings (3 brothers and 2 sisters) were all normal and healthy. She had no family history of a similar condition or other skin problems. Her medical history was unremarkable with no other obvious complaints, and she was not overexposed to sunlight as she spent most of her time indoors. Upon examination, the central part of the face, i.e. the malar area and nose including the ala nasi, showed scattered annular papules and plaques of different sizes ranging from 0.3 to 2 cm. The lesions were asymptomatic and symmetrically distributed on both sides of the face (figs. 1, 2). Routine laboratory tests were done and the results were within normal ranges. Multiple biopsies were taken from different lesions. The diagnosis was confirmed as DSAP, with a classic and very illustrative pathological picture (figs. 3–6). Treatment options were multiple, though none was considered an obvious first choice. All physical modalities were excluded to avoid the possibility of side effects like scarring and disfigurement, as the lesions were located in the central part of the face and involved the skin overlying the nasal cartilage. We first prescribed a topical retinoic acid 0.1% cream that has keratolytic and antineoplastic effects, and it was assumed that this might rectify the faulty clonal epidermopoiesis. The patient applied the cream twice daily for 3 months without a significant response. Six months later, the patient presented again and the picture was almost unchanged except for a few small new lesions in the malar area and on the nose. We decided to give her imiquimod 5% cream to be used once a day (3 times per week) for 24 weeks. She came for a follow-up every 2 weeks. Her skin started to respond after 1 month. Side effects like erythema, crustations and pruritus were experienced and were controlled by topical emollient cream; we avoided corticosteroids and calcineurin inhibitors because they can counter the effects of imiquimod. On completion of the therapy, the response was very satisfactory for both the patient and the therapist. There were some lesions with only a partial response. Central scarring and the scars across the nose improved in their color, thickness and texture, constituting an added benefit of imiquimod in this indication. The patient maintained a once-weekly application of imiquimod. There has been no relapse after follow-up for 2 years now.

Discussion

Porokeratosis is inherited as an autosomal dominant trait; however, sporadic cases are also known to occur. Chernosky and Freeman [4] first described DSAP in 1967 in a Texas population. DSAP is the most common of the clinical variants and may account for almost half of all cases in the USA [5], while in Singapore it is second to the Mibelli type in frequency...
(18 and 56%, respectively) [6]. It usually develops in the third and fourth decades of life. The incidence in females is double that of males, unlike the classic Mibelli type. DSAP is characterized by multiple lesions that are superficial, relatively smaller than the Mibelli type, slightly pigmented, annular, keratotic and with a central atrophic area. It mainly affects areas exposed to sunlight; however, paradoxically, the limbs are more affected than the face [7].

Fifteen percent of DSAP patients have lesions on the face. Other regions such as the extensor surfaces of the extremities constitute the majority of reported cases, with cases of lesions appearing exclusively on the face being probably less frequent. Rarely, actinic porokeratosis is confined to the nose only. Of cases reported, 7.5% have revealed a malignancy arising within the porokeratosis. Unlike other forms, DSAP does not appear to have more risk of malignant change than other types [8]. It can sometimes coexist with other forms of porokeratosis (Mibelli type, linear porokeratosis, porokeratosis palmaris et plantaris disseminata and punctate porokeratosis) [9]. There is good evidence that ultraviolet light can precipitate the development of new lesions or exacerbate pre-existing lesions [5]. Our patient was minimally exposed to the sun and she spent most of her time indoors.

The exact pathogenesis of porokeratosis is not clear; however, it has been assumed that a focal clonal expansion of abnormal cells gives rise to the shape of the coronoid lamella. The coexistence of different variants in one patient or in several members of an affected family indicates different phenotypic expressions of a common genetic entity which could possibly be explained by a simultaneous expression of closely linked genes.

Genetic studies have mapped the loci responsible for DSAP to chromosomes 12q, 15q and 18q. Several mutations have been identified in the SSH1 gene on chromosome 12, which encodes a phosphatase that plays a pivotal role in actin dynamics [10, 11]. In a genome-wide scanning and linkage analysis performed on six generations of a Chinese family with DSAP, two missense mutations in SSH1 and ARPC3 were found, which are involved in the actin cytoskeleton pathway and interact with adherent junctions in the epidermal cells. This finding suggested that cytoskeleton disorganization in epidermal cells was likely associated with the pathogenesis of DSAP [12]. In a recent study on four Chinese families, in 33% of the patients, three mutations were found in the mevalonate kinase (MVK) gene which plays a role in lipid metabolism [13].

Psoriasis and phototherapy (UVA, BB-UVB and NB-UVB) have been associated with porokeratosis. DSAP has also been associated with immune suppression, HIV infection, diabetes mellitus, liver cirrhosis, acute pancreatitis, Crohn’s disease, solid malignancy, the administration of immunomodulating drugs used to treat autoimmune diseases and following the transplantation of organs (particularly in kidney transplant patients) [14, 15].

Treatment of DSAP is often difficult. Therapies such as cryotherapy, photodynamic therapy, erbium YAG and CO2 lasers or the application of topical 3% diclofenac gel or 5-fluorouracil cream are usually partially successful, but with inconsistent results [16]. Topical imiquimod 5% cream may prove to be a useful treatment option for DSAP [17]. This therapy treated the plaques effectively, there were no more keratotic elevations or furrows, and even the central scars showed an improvement in color and texture. Biologically, imiquimod is also antineoplastic and is used to treat nonmelanoma skin cancers; this is a major added advantage because DSAP may, in rare instances, be complicated by squamous cell carcinoma. Further studies are required to confirm whether imiquimod should have a place in the management of DSAP. The mechanism of action of imiquimod in the treatment of DSAP is not yet clear: it may suppress or switch off the abnormal mutant genes through its immunological effects on both adaptive and innate immunity.
In conclusion, we presented a case of facial sporadic DSAP, which was treated by application of imiquimod 5% cream in conjunction with regular sunscreens. Follow-up of this case is important to rule out the possibility of malignant transformation.

References


Fig. 1. The right side of the patient’s face before (a) and after (b) imiquimod 5% cream therapy (once a day, 3 times weekly for 16 weeks). Note the disappearance of the keratotic rim, the flattening of the lesions and the improvement in the color and thickness of the scars. Residual hyperpigmentation and hypopigmentation remained unchanged.

Fig. 2. The patient before (a) and after (b) imiquimod 5% cream therapy (once a day, 3 times weekly for 16 weeks). Notice the flattening of the lesions and the improvement of skin texture.
Fig. 3. A histopathology section. ×100. Black arrows point to invaginations in the epidermis of both specimens. Note the outward projection of a parakeratotic column called the cornoid lamella. This is a classic, characteristic picture of porokeratosis.

Fig. 4. A histopathology section. ×200. A closer view of the pathognomonic features of porokeratosis.
Fig. 5. A histopathology section. ×400. A section showing abnormal vacuolated keratinocyte clones that develop in between normal cells and interrupt the granular layer, with no granules in the abnormal cells. A column of parakeratotic cells forms, called the cornoid lamella.

Fig. 6. A histopathology section. ×400. Another lesion with similar features to figure 5.