Perineal warty plaques: A case of verrucous porokeratosis

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Abstract
Nonvenereal genital dermatoses form an important category of disorders, and verrucous porokeratosis is a rare and less recognized entity among the same. We present the case of a young adult male with warty growths over scrotum and buttocks for a year. Characteristic cornoid lamellae with typical differentiating features were seen in the histopathology, establishing the diagnosis. This case emphasizes the rare nonvenereal cause for a condition clinically mimicking condyloma acuminata.

Key words: Perineum, porokeratosis, ptychotropica, verrucous, warty

INTRODUCTION
Porokeratosis is a group of keratinization disorder, characterized clinically by plaques with a keratotic rim that corresponds histologically to the characteristic cornoid lamellae. It has several clinical variants, each with slightly different presentation, histological features, and prognosis. Verrucous porokeratosis, which some authors also refer to as genitogluteal porokeratosis or porokeratosis ptychotropica, is a rare subtype.[1] It presents as a diagnostic challenge and an important disorder to be considered among nonvenereal genital dermatosis. This case is being reported owing to its rarity and close clinical mimic of condyloma acuminata.

CASE REPORT
A 27-year-old male, otherwise healthy, presented to the dermatology outpatient with the complaint of multiple, gradually expanding, dark-colored warty growths over scrotum and buttocks, for the past 1 year. The lesions were not associated with itch or pain. He was unmarried and disclosed a history of unprotected heterosexual encounters with multiple partners. A written informed consent was taken from the patient for clinical examination and pictures. On cutaneous examination, multiple hyperpigmented annular plaques ranging from 1 cm to 3 cm, with verrucous surface and prominent ridge-like borders over a background of hyperpigmented and velvety skin, were found over the perineal region, involving the ventral aspect of scrotum, medial aspect of both thighs and buttocks [Figure 1]. There was no previous history of genital ulcer or discharge (purulent or otherwise). His past medical history was noncontributing and systemic examination did not reveal any abnormality.

Laboratory investigations were insignificant, and viral markers were negative. A provisional diagnosis of verrucae, Bowenoid papulosis, tuberculosis

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verrucosa cutis, and porokeratosis were reached, and punch biopsy from a lesion was sent for histopathology. Histopathological examination revealed irregular psoriasiform hyperplasia with compact orthokeratotic hyperkeratosis, multiple broad cornoid lamellae overlying shallow epidermal invagination with hypogranulosis, and numerous dyskeratotic keratinocytes [Figures 2 and 3]. These features helped in establishing the diagnosis of verrucous porokeratosis. The patient was given topical imiquimod and reported mild resolution after 4 weeks of therapy but was later lost to follow-up.

**DISCUSSION**

Verrucous porokeratosis poses a diagnostic challenge, not only because it closely resembles the more common entities such as condyloma acuminata and psoriasis but also because of its rarity and lack of knowledge of the condition among treating physicians. We could find only <40 cases of verrucous porokeratosis in literature.[2] Classically, it has been described to present as symmetrical verrucous papules and plaques with satellite lesions involving the genitogluteal region.[3] The word ptychotropica itself is derived from Greek “ptyche” meaning fold and “trope” meaning turning and was first described as a separate entity in 1995 by Luckert et al. who called it “porokeratosis ptychotropica.”[4] Its pathogenesis has not been clearly deciphered yet, with repeated friction being proposed as a triggering factor, as could also have been in this case as the patient was obese and had acanthosis nigricans too; other than infections, drugs, ultraviolet radiation, and immunosuppression.[3] Furthermore, it has been found to occur more frequently in men.[5] The condition is known to evade diagnosis for years, with patients undergoing a number of therapeutic trials, and respite from seemingly none.[1] Absence of the characteristic keratotic rim makes it difficult to recognize, as does the presentation as a verrucous plaque. Although it has a predilection for genitogluteal region, lesions over thighs and lower legs have also been reported.[5]

The various subtypes of porokeratosis are differentiated mainly on clinical grounds. Verrucous porokeratosis appears as annular white-yellow structure demarcating a central scar-like area surrounded by minimal peripheral vascularization on dermoscopy, which further aids in diagnosis, though the feature is not specific to this subtype.[6] Cornoid lamellae, a histological sine qua non of porokeratosis not only helps to distinguish the entity from other clinical differentials, but also the various subtypes on the basis of its distribution, extent, and degree of prominence. Verrucous porokeratosis has a unique morphology and is
relatively refractory to treatment, therefore, requires differentiation from porokeratosis of Mibelli. This can be accomplished by the morphology of cornoid lamellae which are shown to be multiple, larger, and/or confluent, sometimes even spanning the entire surface, in verrucous porokeratosis.[7] Moreover, psoriasiform hyperplasia, hyperkeratosis, dermal telangiectasia, and papillomatosis are few other features seen that often add to the already pronounced confusion of clinical similarity with psoriasis.[8] Several cases have also been reported with cutaneous amyloid deposition secondary to frequent rubbing and scratching caused by itch.[1] A noteworthy issue is the presence of dyskeratotic cells overexpressing p53 gene underlying the cornoid lamellae, indicating the role of tumor suppressor gene in its pathogenesis.[9]

There is a lack of literature regarding difference in presentation of verrucous porokeratosis in immunocompetent and immunocompromised individuals, though porokeratosis in general has more extensive presentation in immunocompromised patients.[9] Nature of the disease as revealed by clinical and histopathologic findings is essentially the same in both the group of patients.[10]

Treatment options are largely limited for verrucous porokeratosis with the condition being mostly refractory to conventional therapy and none so far has been reported to achieve complete clearance. 5-fluorouracil, imiquimod, steroids, calcipotriol, tacrolimus, Vitamin A, and cychotherapy have been used but with variable results.[11] Photodynamic therapy was found to be useful as an alternative treatment, but its utility remains limited to providing symptomatic relief in itch and partial clearance of lesion, and it did not achieve curative response.[12] Smaller lesions can be approached for surgical intervention as has been shown in a few case reports, but it is difficult to comment upon the rate of recurrence on the basis of only these few.[13] Further studies are required to ascertain the role and prognosis of surgery in verrucous porokeratosis.

Close follow-up is necessary, for there is a yet uncertain risk of malignancy given the known risk of 7%–11% with porokeratosis, mainly squamous cell carcinoma, and also Bowen disease and basal cell carcinoma.[8] Although no such transformation has yet been reported with verrucous porokeratosis, this figure can run as high as up to 24% in porokeratosis in patients with risk factors (linear variant, long-standing large lesion, late onset, old age, and history of prior radiation exposure).[14] Verrucous porokeratosis is known to run a long course before being adequately diagnosed, and so need to be monitored to look for any possible malignant degeneration.

To conclude, verrucous porokeratosis is an important differential to be considered among nonvenereal dermatosis and needs histological confirmation for diagnosis. Refractoriness to conventional therapy and a possible potential of malignant transformation emphasize the need to keep the condition under monitoring. Better understanding of pathogenesis and advent of further therapeutic modalities may improve clinical outcome of these patients.

**Declaration of patient consent**
The authors certify that they have obtained all appropriate patient consent forms. In the form the patient(s) has/have given his/her/their consent for his/her/their images and other clinical information to be reported in the journal. The patients understand that their names and initials will not be published and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

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**Conflicts of interest**
There are no conflicts of interest.

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Lepromatous leprosy as a presenting feature of HIV

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Abstract
Various bacterial, mycobacterial and fungal opportunistic infections occur frequently in immunocompromised individuals, however, leprosy in retroviral disease is a relatively rare association. Hereby, we report a case of lepromatous leprosy that presented with clinical features mimicking other opportunistic infections and subsequently led to the diagnosis of HIV. The myriad challenges associated with the diagnosis and management of HIV–leprosy coinfection are also discussed. Thus, although uncommon, atypical cutaneous lesions in HIV-seropositive patients warrant investigation for leprosy.

Key words: HIV, lepromatous leprosy, opportunistic infections

INTRODUCTION
Leprosy patients with HIV clinically present similar to patients with leprosy in the general population. Sometimes, on starting antiretroviral treatment (ART), subclinical leprosy manifests as clinical leprosy or the preexisting leprosy worsens immune reconstitution inflammatory syndrome (IRIS).[1] HIV-positive patients with low immunity may present with various opportunistic infections such as cutaneous cryptococcosis, deep fungal infections, extensive molluscum contagiosum, and extensive viral warts.[2] Our case of lepromatous leprosy is reported for its rare opportunistic infection-like manifestations. Thus, leprosy can be considered the presenting feature in this patient that led to the detection of his HIV-positive serostatus.

CASE REPORT
A 50-year-old married male presented with asymptomatic reddish raised lesions over the face, trunk and extremities for 1 month, with a history of anorexia, progressive weight loss and mild abdominal pain.

On examination, multiple erythematous to skin-coloured papules and nodules were present on the face, trunk and extremities with a looser distribution on the distal extremities. The lesions were non-tender and non-pruritic. The patient was otherwise asymptomatic.

Laboratory investigations showed a CD4 count of 350 cells/mm³. Mantoux testing was positive. Tuberculosis polymerase chain reaction (PCR) was negative. HIV serology was positive.

Histopathology of skin biopsy showed a lepromatous lesion with large, non-acid-fast bacilli (AFB) positive macrophages. The patient was diagnosed with lepromatous leprosy.

The patient was started on antiretroviral therapy (ART) and multidrug anti-leprosy therapy (MDT) with dapsona, rifampicin, and clofazimine. The patient is still on follow-up and is asymptomatic.

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