Keratotic spines in a patient with pruritic and dyskeratotic dermatosis: A new clinical finding

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INTRODUCTION

The Polyomaviridae family includes 13 small, nonenveloped, double-stranded, circular human DNA viruses. Polyomavirus infection is usually latent and asymptomatic. Clinical disease, although rare, primarily occurs in patients who are immunosuppressed iatrogenically. In such patients, viral reactivation can cause disease involving the kidneys, central nervous system, or skin. Cutaneous manifestations associated with the Polyomaviridae include Merkel cell carcinoma, trichodysplasia spinulosa (TS), and, more recently, pruritic and dyskeratotic dermatosis (PDD). The viruses responsible for disease are Merkel cell polyomavirus, TS-associated polyomavirus, human polyomavirus 6 (HPyV6), and human polyomavirus 7 (HPyV7). We report a patient with an unusual presentation of PDD that included a lichenified dermatosis associated with alopecia and follicularly-based keratotic papules.

CASE REPORT

A 44-year-old man who had undergone kidney and pancreas transplantation 7 years prior presented with a 2-year history of a diffuse, pruritic, burning rash associated with dyspigmentation and lichenification as well as tender follicular-based papules involving the dorsal hands and elbows and pits involving the wrists and palms (Fig 1). A biopsy from the upper extremity revealed mild acanthosis, spongiosis, and patchy dyskeratotic dermatosis extending somewhat linearly into a hyperkeratotic stratum corneum (Fig 2). The paraffin-embedded tissue was tested for HPyV6, HPyV7, and TS-associated polyomavirus.

Abbreviations used:

HPyV6: human polyomavirus 6
HPyV7: human polyomavirus 7
PDD: pruritic and dyskeratotic dermatosis
TS: trichodysplasia spinulosa

was performed for untreated hypertension and diabetes mellitus. Prior to the presentation, his eruption had been diagnosed as psoriasis by another physician, and he had been treated with intramuscular triamcinolone acetonide, triamcinolone 0.1% cream, and narrowband ultraviolet B phototherapy without improvement. At the time of presentation, his medications included doxazosin 8 mg/day, gabapentin 300 mg twice daily, metformin 500 mg/day, metoprolol tartrate 100 mg/day, pantoprazole 20 mg/day, tacrolimus 1 mg/day, and everolimus 2.5 mg twice daily.

Physical examination revealed diffuse, hyperpigmented, scaly, lichenified patches, patchy alopecia, and follicular-based keratotic papules and pits involving the elbows, wrists, and palms (Fig 1). A biopsy from the upper extremity revealed mild acanthosis, spongiosis, and patchy intraepidermal dyskeratosis extending somewhat linearly into a hyperkeratotic stratum corneum (Fig 2). The paraffin-embedded tissue was tested for HPyV6, HPyV7, and TS-associated polyomavirus.
HPyV7 was detected in both specimens. The patient was treated with acitretin, and after 3 months of therapy he has had complete clearance of his pruritic eruption. His follicular lesions and palmar pits remain but are less tender.

**DISCUSSION**

PDD is a rare disease in immunocompromised individuals caused by the reactivation of HPyV6 or HPyV7 in the epidermis and sparse involvement of the papillary dermis. Since its initial documentation in 2015, there have been 4 additional reports in the medical literature, all occurring in immunocompromised individuals. However, none of the previously reported patients have demonstrated prominent follicular-based keratotic papules and alopecia.

PDD clinically manifests with pruritic, brown to gray, lichenified plaques involving the trunk and extremities. Ho et al first described the histologic finding of “peacock plumage,” which refers to the irregular columns of parakeratosis noted on histopathologic evaluation. In addition, histopathology reveals variable papillomatosis, intraepidermal dyskeratotic keratinocytes, and a sparse perivascular lymphocytic dermal infiltrate. The cases of PDD reported in the literature are listed in Table I. We report a patient with PDD with an unusual clinical presentation, including keratotic follicular-based papules and alopecia, which have not been previously reported in HPyV6 or HPyV7 cutaneous infection.

TS is characterized by the presence of spine-like follicular papules that involve the face, ears,
Table I. Summary of all literature-documented cases of PDD due to HPyV6 or HPyV7

| Case report | Age (Y) | Sex | Immune status | Clinical presentation | Treatments attempted | Outcome |
|-------------|---------|-----|---------------|-----------------------|---------------------|---------|
| Ho et al4   | 1. 73/M |     | Single lung transplant recipient on alemtuzumab, prednisone, azathioprine, and tacrolimus | 1. Multiple pruritic, brownish-pink, thin, scaly papules and plaques on the trunk and neck sparing the acral areas and face | 1. Topical corticosteroids and antihistamines | 1. Worsening progression of symptoms |
|             | 2. 72/M |     | Double lung transplant recipient on alemtuzumab, tacrolimus, mycophenolic acid, and prednisone | 2. Pruritic, brown, well-defined thin papules and plaques on the trunk, axillae, neck, buttocks, and legs | 2. Topical corticosteroids and diphenhydramine | 2. Diphenhydramine led to minimal improvement, and topical corticosteroids worsened rash |
| Nguyen et al2 | 1. 52/M |     | Kidney and pancreas transplant recipient on tacrolimus and rapamycin | 1. Pruritic rash present for over 2 years | 1. Unknown | 1. Patient deceased |
|             | 2. 36 /F |     | HIV/AIDS | 2. Severe pruritic rash worsening over 12 months | 2. Unknown | 2. Initial worsening for 2 years and then several years of resolution with no immunosuppression |
|             | 3. 54/M |     | None at time of diagnosis but was hospitalized for sepsis, parapharyngeal abscess, and pneumonia | 3. Pruritic rash worsening over 2 years followed by resolution over several years | 3. Unknown | 3. Patient deceased |
| Smith et al5 | 59/M |     | Bilateral lobar (living) transplant recipient on prednisone and tacrolimus | Diffuse, gray, lichenified plaques with confetti-like islands of sparing on the trunk and proximal extremities and generalized pruritus | • Oral steroid • Topical steroid • Antihistamine • Topical (1% and 3%) cidofovir | • Oral steroid: minimal improvement of rash and pruritus • Topical steroid: no relief • Antihistamine: no relief • Topical (1% and 3%) cidofovir: clinical improvement of the rash on the flanks, near clearance on the extremities, and resolution of generalized pruritus |
| Canavan et al3 | 48/M |     | Cardiac transplant recipient on sirolimus and tacrolimus | Severely pruritic, pink to gray macules, thin papules, and plaques on the trunk and extremities, sparing acral areas and the face | Cyproheptadine, doxepin, emollients, topical corticosteroids, pameoxine cream, intravenous cidofovir, and oral acitretin 25 mg | No improvement with topical treatments or antihistamines. Temporary improvement with intravenous cidofovir with subsequent relapse within 4-6 weeks. Complete clinical resolution with oral acitretin |
| Current report | 44/M |     | Kidney and pancreas transplant recipient on tacrolimus and everolimus | Hyperpigmented scaly patches all over body and prominent follicular spines on the elbows and palms | Oral acitretin 25 mg daily | Complete clearing of patches with improvement in keratotic papules at 1-month follow-up |

HPyV6, Human polyomavirus 6; HPyV7, human polyomavirus 7; PDD, pruritic and dyskeratotic dermatosis.
extremities, and trunk, also showing lichenification of the skin and alopecia of the eyebrows. The histopathologic features include enlarged, dystrophic hair follicles and accumulation of eosinophilic, perinuclear globules within inner root sheath cells, and the intrafollicular accumulation of keratotic/parakeratotic material. The unusual keratotic papules present in our patient suggest that TS and PDD can have similar clinical presentations characterized by keratotic papules and spines, though the clinical distribution of lesions and histopathologic features should allow the distinction of the 2 conditions. Based on our report of alopecia and keratotic papules similar to TS, physicians might consider testing for all 3 polyomaviruses known to cause cutaneous disease to establish a definitive diagnosis in cases showing overlapping clinical features, especially if the histopathologic findings fail to distinguish the conditions. The presence of high sequence homologies in the coding region for all 3 polyomaviruses likely explains the clinical overlap in dermatoses resulting from HPyV6, HPyV7, and TS-associated polyomavirus infection. The pathogenic mechanism by which HPyV7 causes PDD is unknown, though it is postulated that human polyomavirus causes disease via upregulation of viral oncogenes that include tumor antigens. The small tumor antigen of HPyV7 is noted to be elevated in virus-positive eruptive pruritic dermatoses. In a recent study, Wu et al found that HPyV7 small tumor antigen was found to dysregulate protein phosphatase 2A through physical interaction, resulting in increased activation of MEK/ERK/c-JUN and 4E-BP1, which may lead to dysregulation of keratinocyte growth and hyperproliferation. Further studies are needed to elucidate the pathogenic mechanism.

PDD is a rare cutaneous disease with a limited documentation of effective treatment. The reduction of immunosuppression and initiation of antiviral treatment are effective options to control viral replication and clinical progression if clinically feasible. Anecdotal reports describe treatment failure with antihistamines and corticosteroids. Successful treatment has been documented with topical cidofovir, a cytosine analog that inhibits human polymerase activity required for polyomavirus replication, and/or acitretin, a vitamin A analog that increases keratinocyte turnover. Cidofovir and acitretin have also shown efficacy in the treatment of TS, though acitretin was coadministered with antiviral therapy (oral valganciclovir). In summary, we present an additional report of a recently described HPyV7-associated dermatosis in a patient who underwent solid organ transplantation that was characterized by lichenified plaques, with novel additional findings of follicular-based papules and pits and alopecia. PDD should be included along with TS in the differential diagnosis of a diffuse pruritic rash with keratotic papules and pits and/or alopecia in an immunosuppressed patient. While keratotic papules and spines, alopecia, and response to cidofovir treatment reflect the similar viral class of origin of PDD and TS, the histologic features of the 2 conditions appear to be disparate. Based on the few cases of PDD reported in the literature, a diffuse eruption of lichenified patches or plaques would favor infection with HPyV6 or HPyV7, as the histologic findings fail to show follicular epithelial dysmorphism characteristic of TS. Despite the lack of a specific viral cytopathic effect in lesions of PDD, the presence of columnar parakeratosis and/or spotty intraepidermal dyskeratosis should alert the pathologist to the diagnosis when there is a history of immunosuppression.

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