Tinea Versicolor Mimicking Pityriasis Rubra Pilaris caused by *Malassezia globosa*

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**Abstract**

Tinea versicolor (TV) is a common fungal skin disease caused by the *Malassezia* species. This disease usually presents as hypopigmented- or hyperpigmented coalescing scaly macules, papules, patches or plaques on the trunk and upper arms. Herein, we report a rare clinical manifestation of TV in a 29-year-old man presenting with marked follicular, erythematous, and hyperkeratotic papules on the trunk with erythematous scaly macules and patches on the upper extremities with intermittently spared skin. We initially suspected pityriasis rubra pilaris, however, skin biopsy results and mycological examination revealed TV. Polymerase chain reaction-based sequence analysis revealed *Malassezia globosa*. The patient was successfully treated with oral itraconazole and topical terbinafine.

**Key Words**: *Malassezia*, *Malassezia globosa*, Tinea versicolor, Pityriasis rubra pilaris

**INTRODUCTION**

Tinea versicolor (TV) is a common cutaneous fungal disease caused by the *Malassezia* species, mostly *M. globosa*. Although these organisms usually inhabit the human skin as normal follicular flora in the yeast phase, they produce skin lesions when they grow to the hyphal phase. Typically, TV presents in one of the three forms: papulosquamous, folliculitis, or inverse. The papulosquamous form is the most common manifestation and is characterized by hypopigmented or hyperpigmented coalescing scaly macules, papules, patches or plaques on the trunk and upper arms. However, other unusual clinical presentations such as the atrophic form and mimics of pityriasis rotunda, mycosis fungoides, and pityriasis rubra pilaris have also been reported. In this case report, the authors describe a patient who showed hyperkeratotic follicular papules and scaly coalescent erythematous macules over the trunk and upper extremities with intermittently spared skin. Although we first suspected PRP, skin biopsy results confirmed a diagnosis of TV. To the best of our knowledge, this is the third reported case of TV.
mimicking PRP.

**REPORT OF A CASE**

A 29-year-old man presented with a 10-year history of skin eruption. He denied any histories of medication and other cutaneous or systemic diseases, although topical steroid was applied on the skin lesion without improvement. On presentation, marked follicular, erythematous, hyperkeratotic papules were noted on the trunk (Fig. 1A, 1B). Erythematous scaly macules and patches with...

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**Fig. 1.** (A, B) Marked follicular, erythematous, and hyperkeratotic papules noted on chest and abdomen on first presentation. Erythematous scaly macules and patches with intermittent spared skin noted on (C-F) upper extremities.
intermittent spared skin were also noted on upper extremities (Fig. 1C-F). He reported that the skin lesions progressed from his neck in a cephalo-caudal distribution to involve his chest, abdomen, back, proximal arms, and proximal legs. Owing to the appearance, PRP was primarily raised as a possible diagnosis and skin punch biopsy was performed. Histologic examination revealed thick basket-weave stratum corneum with septate hyphae and spores (Fig. 3A, 3B), consistent with TV. On the next visit, direct microscopic potassium hydroxide (KOH) examination revealed short, thick, fungal hyphae and large numbers of spores. Fungal culture of skin scrape showed growth of *Malassezia* species.

**Fig. 2.** Markedly improved skin lesions with remnant mild pigmentation at follow-up visit.
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on a modified Leeming and Notman agar medium plate. PCR-based sequence analysis of the internal transcribed spacer region of the fungal rRNA gene was performed and showed 99% homology with *M. globosa* in the GenBank database (accession no. AY743630.1). The patient was treated with itraconazole 200 mg/d for 1 week and topical terbinafine for 3 weeks. At the follow-up visit, KOH revealed no fungal organisms and skin lesions improved with remnant mild pigmentation (Fig. 2A-F).

**DISCUSSION**

TV is a common skin fungal infection caused by the *Malassezia* species. These organisms are usually present in 75~98% of healthy individuals as yeasts and are considered normal follicular flora. They produce skin lesion, which is TV, when they grow to hyphae. The prevalence of *Malassezia* is reportedly up to 50% in hot and humid regions compared to 1.1% in colder climates. *M. globosa*, the major pathogen of TV, is a dimorphic, lipophilic organism that requires lipid for growth. This characteristic is apparent in that skin lesions are common among adolescents whose sebaceous glands are more active and in oily areas of skin, such as the chest, abdomen, back, pubis, and intertriginous areas. Hyperhidrosis, oral contraceptives, systemic corticosteroid use, Cushing's disease, immunosuppression, and malnourishment are also predisposing factors for TV.

Typically, a diagnosis of TV is based on a typical presentation of hypopigmented- or hyperpigmented, scaly, oval to round macules scattered over the upper trunk and arms, and can be confirmed through a microscopic KOH examination. A combination of short, thick fungal hyphae and variably sized spores on KOH examination is commonly called "spaghetti and meatballs". *M. globosa* reveals whitish, dull, and friable colonies on a modified Leeming and Notman agar medium plate. Histologic examinations show clusters of oval budding yeast cells with short, septate, hyphae inhabiting the stratum corneum. TV should be differentiated from seborrheic dermatitis, pityriasis rosea, PRP.
confluent and reticulated papillomatosis, pityriasis alba, Hansen's disease, syphilis, and vitiligo\textsuperscript{8,9}. Systemic regimens used for treating TV include itraconazole, ketoconazole, and fluconazole\textsuperscript{8}. Although topical terbinafine has been proven effective in TV treatment, it is ineffective via the oral route\textsuperscript{8}. In this case, oral itraconazole and topical terbinafine are considered to have contributed to clinical improvement.

PRP is a rare, chronic papulosquamous disorder characterized by reddish-orange scaly patches, hyperkeratotic follicular papules, and palmoplantar keratoderma\textsuperscript{1}. Skin lesions may progress to erythroderma with sharply demarcated islands of normal skin ("nappes claires"). The disease shows a bimodal age distribution with peak incidences in the first and fifth decades of life. It is known that there are no predilections for PRP regarding race and gender. PRP is classified into six types based on clinical characteristics such as age, pattern of lesion, prognosis, and medical or family history\textsuperscript{1}. The classic adult type accounts for over 50% of all cases, shows a cephalo-caudal progression of skin lesions, and often resolves within three years. For the 10% of patients with atypical adult type, however, PRP shows a predilection for skin lesions on the legs, and runs a chronic course with skin lesions persisting for over 20 years. Histologically, erythematous lesions show acanthosis with broad and short rete ridges, and alternating orthokeratosis and parakeratosis in both vertical and horizontal directions\textsuperscript{10}. Follicular papules show dilated infundibula filled with an orthokeratotic plug with perifollicular shoulders of parakeratosis at times. In our case, however, none of these histologic findings was apparent, indicating no concomitant PRP.

As far as we know, there have only been two cases of TV imitating PRP. Darling et al.\textsuperscript{6} described a 54-year-old patient with a 20-year history of pruritic erythematous patches and keratotic follicular papules involving the trunk and upper arms, the patient was diagnosed with TV with only KOH findings. Berry and Khachemoune\textsuperscript{2} confirmed TV with skin biopsy results in a 65-year-old patient presenting with similar skin lesions distributed along the trunk and proximal extremities. Although diagnoses were made, the exact pathogen could not be elucidated in both these cases. In our case, however, we confirmed TV with skin biopsy results and identified \textit{M. globosa} as the causative organism. We believe that the long disease duration with improper use of corticosteroids may have led to this finding.

This case report represents an unusual presentation of TV mimicking PRP, which we consider unique and educational. Clinical presentations including scaly erythematous patches with islands of normal skin and hyperkeratotic follicular papules first led us to an initial diagnosis of PRP, however, skin biopsy confirmed TV. We suggest that dermatologists should always perform basic tests such as KOH when confronting patients with scaly erythematous patches before performing invasive examinations. Also, it is possible that TV with long duration and inappropriate treatment may demonstrate non-typical features mimicking many other diseases.

\textbf{Conflict of interest}

In relation to this article, I declare that there is no conflict of interest.

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