A man in his 20s presented with multiple, asymptomatic, variably sized, well-defined, hypopigmented patches on his neck and trunk for the past 8 years (Figure 1). He also had multiple discrete, nontender, firm, skin-colored to slightly erythematous papulonodular lesions on the central part of face, ears, limbs, and back, varying in size from few millimeters to some reaching 1–2 cm for the past 6 years. There was a grouping of papulonodules over the nose and mouth area of the face (Figure 2). There was no loss of appendages or feeding nerve localized to the hypopigmented plaques. There was no hypoesthesia within the hypopigmented patches. Nerves were not enlarged. There was no other significant past history.

Laboratory investigations showed positive IgG serology for rk39 antibody test (rk39 antibody detection test kits by K.N.D. Research Solution, SAS Nagar, Mohali, India). Skin biopsies showed presence of basal cell vacuolisation. The dermis showed mixed inflammatory infiltrates comprising lymphocytes, histiocytes, plasma cells, and neutrophils. There was presence of well-formed epithelioid cell granulomas with scattered multinucleated giant cells. The histiocytes showed intracytoplasmic amastigote forms of leishmania (Figure 3A and B), Giemsa stain showed presence of Leishman–Donovan bodies (Figure 3C). Polymerase chain reaction from skin biopsy for leishmania was positive (positive control used was DNA from cultured promastigotes, negative control was nucleus free water). A diagnosis of post-kala-azar dermal leishmaniasis (PKDL) was established based on clinical, laboratory, and histopathological findings. The patient was treated with intravenous liposomal amphotericin B 2.5 mg/kg/day for 20 days, and there was significant reduction in all skin lesions at end of therapy (Supplemental Figure 1).

This is a case of PKDL without a history of visceral leishmaniasis (VL), which is particularly difficult to diagnose. PKDL without recognized VL is a form leishmaniasis that has garnered increasing attention in India as well as other parts of the world in recent years.1,2 Such cases are due to asymptomatic or subclinical Leishmania donovani infection.3

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1 Am. J. Trop. Med. Hyg. 00(00), 2024, pp. 1–3 doi:10.4269/ajtmh.24-0018
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However, PKDL occurs after VL only in a subset of people, and factors implicated in its cause include exposure to antimonial therapy and ultraviolet radiation (UVR); UVR acts as a potent immunosuppressant, induces production of IL-10, and alters the functioning of the antigen presenting cells. Morphologies of PKDL include hypopigmented, papulonodular, and polymorphic types, the latter of which includes multiple lesion morphologies such as papules, plaques, nodules, and hypopigmented patches. Diffuse cutaneous leishmaniasis (DCL), caused by *Leishmania amazonensis*, is a rare form of cutaneous leishmaniasis where parasites grow uncontrolled in lesions across large skin areas. Leishmania-specific immunity is higher in cases of PKDL than DCL, and rates of parasite positivity are lower in PKDL than DCL.

Lepromatous leprosy remains a close differential diagnosis of PKDL in VL-endemic regions. Subtle features on clinical examination such as involvement of the nose and mouth area, lack of sensory loss, and involvement of warm and cool areas of the body (like the spine) in PKDL helps in differentiating it from various forms of leprosy, which are typified by hypopigmented, anesthetic skin lesions. rk39 serology is often used for complementing diagnosis in PKDL; however, an important limitation of this test is that it cannot distinguish between current and past infection.

PKDL with or without antecedent VL is complex in terms of epidemiology, diagnosis, and management. The emergence of PKDL without history of VL in India indicates evolving changes in our understanding of the disease, enabled by molecular diagnosis. A high level of clinical suspicion combined with enhanced diagnostic tools is essential for developing targeted control strategies for PKDL and VL control in endemic areas.

Received January 11, 2024. Accepted for publication May 12, 2024. Published online July 23, 2024.


Acknowledgment: The American Society of Tropical Medicine and Hygiene (ASTMH) assisted with publication expenses.

Authors’ contributions: A. Sharma: conceptualization, writing—original (lead), writing—revision (lead). A Bishnoi: writing—original (equal), writing—revision (equal). D. Parsad: conceptualization, writing—original (equal), writing—revision (equal), supervision.

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