Inflammatory, infectious, and neoplastic processes can all occur in prior areas of herpes infection (postherpetic isotopic response [PHIR]). Postzoster granulomatous dermatitis is among the most frequently encountered PHIR, but the exact pathogenesis has not been fully elucidated. Rarely, multiple diseases manifest concurrently in a PHIR. We report a case of cutaneous chronic lymphocytic leukemia (CLL) with an associated granulomatous dermatitis and medium-vessel vasculitis occurring simultaneously at the site of prior herpes zoster. Clinicians and dermatopathologists should be aware of the multiplicity of postzoster isotopic responses and should consider the possibility of multiple diseases manifesting in the same clinical lesion in this setting.

**Case Report**
A 55-year-old man with a 4-year history of CLL was admitted to the hospital due to a painful rash on the left chest, back, and left arm.

**Conditions That May Occur as a Postherpetic Isotopic Response**

- Acneform lesions
- Basal cell carcinoma
- Breast carcinoma
- Dermatophytosis
- Granuloma annulare
- Granuloma vasculitis
- Leukemia cutis
- Lichen planus
- Lichen sclerosus et atrophicus
- Lymphoma
- Metastatic carcinoma
- Pseudolymphoma
- Sarcoïdosis
- Squamous cell carcinoma
- Verrucae


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side of the face of 2 months’ duration. Erythematous to violaceous plaques with surrounding papules and nodules were present on the left side of the forehead and frontal scalp with focal ulceration. Two months prior, the patient had unilateral vesicular lesions in the same distribution (Figure 1A). He initially received a 3-week course of acyclovir for a presumed herpes zoster infection and showed prompt improvement in the vesicular lesions. After resolution of the vesicles, papules and nodules began developing in the prior vesicular areas and he was treated with another course of acyclovir with the addition of clindamycin. When the lesions continued to progress and spread down the left side of the forehead and upper eyelid (Figure 1B), he was admitted to the hospital and assessed by the consultative dermatology team. No fevers, chills, or other systemic symptoms were reported.

A punch biopsy showed a diffuse lymphocytic infiltrate filling the dermis and extending into the subcutis with nodular collections of histiocytes and some plasma cells scattered throughout (Figure 2A). A medium-vessel vasculitis was present with numerous histiocytes and lymphocytes infiltrating the muscular wall of a blood vessel in the subcutis (Figure 2B). CD3 and CD20 immunostaining showed an overwhelming majority of B cells, some with enlarged atypical nuclei and a smaller number of reactive T lymphocytes (Figure 2C). CD5 and CD43 were diffusely positive in the B cells, confirming the diagnosis of cutaneous CLL. CD23 staining was focally positive. Immunostaining for κ and λ light chains showed a marginal κ predominance. An additional biopsy for tissue culture was negative. A diagnosis of postzoster granulomatous dermatitis with vasculitis and cutaneous CLL was rendered.

Comment
Postherpetic Cutaneous Reactions—Various cutaneous reactions can occur at the site of prior herpes infection. The most frequently reported reactions are granulomatous dermatitides such as granuloma annulare, granulomatous vasculitis, granulomatous folliculitis, sarcoidosis, and nonspecific granulomatous dermatitis.1 Primary cutaneous malignancies and cutaneous metastases, including hematologic malignancies, have also been reported after herpetic infections. In a review of 127 patients with postherpetic cutaneous reactions, 47 had a granulomatous dermatitis, 32 had nonhematologic malignancies, 18 had leukemic or lymphomatous/pseudolymphomatous infiltrates, 10 had acneiform lesions, 9 had nongranulomatous dermatitides such as lichen planus and allergic contact dermatitis, and 8 had nonherpetic skin infections; single cases of reactive perforating collagenosis, nodular solar degeneration, and a keloid also were reported.1

Pathogenesis of Cutaneous Reactions—Although postherpetic cutaneous reactions can develop in healthy individuals, they occur more often in immunocompromised patients. Postherpetic isotopic response has been used to describe the development of a nonherpetic disease at the site of a previous herpes infection. The pathogenesis of postherpetic cutaneous reactions involves the activation of latent herpesvirus following reactivation of the virus. A detailed pathogenesis of postherpetic cutaneous reactions is beyond the scope of this article. Further studies are needed to elucidate the mechanisms underlying the development of these cutaneous reactions.
site of prior herpes infection. Several different theories have been proposed to explain the pathogenesis of the PHIR, including an unusual delayed-type hypersensitivity reaction to residual viral antigen or host-tissue antigen altered by the virus. This delayed-type hypersensitivity explanation is supported by the presence of helper T cells, activated T lymphocytes, macrophages, varicella major viral envelope glycoproteins, and viral DNA in postherpetic granulomatous lesions; however, cases that lack detectable virus and viral DNA in these types of lesions also have been reported.

A second hypothesis proposes that inflammatory or viral-induced alteration of the local microvasculature results in increased site-specific susceptibility to subsequent inflammatory responses and drives these isotopic reactions. Damage or alteration of local peripheral nerves leading to abnormal release of specific neuromediators involved in regulating cutaneous inflammatory responses also may play a role. Varicella-zoster virus utilizes the peripheral nervous system to establish latent infection and can cause destruction of alpha delta and C nerve fibers in the dermis. Destruction of nerve fibers may indirectly influence the local immune system by altering the release of neuromediators such as substance P (known to increase blood vessel permeability, increase fibrinolytic activity, and induce mast cell secretion), vasoactive intestinal peptide (enhances monocyte migration, increases histamine release from mast cells, and inhibits natural killer cell activity), calcitonin gene-related peptide (increases vascular permeability, endothelial cell proliferation, and the accumulation of neutrophils), and melanocyte-stimulating hormone (induces anti-inflammatory cytokines). Disruption of the nervous system resulting in an altered local immune response also has been observed in other settings (eg, amputees who develop inflammatory diseases, bacterial and fungal infections, and cutaneous neoplasms confined to stump skin). 1

Malignancies in PHIR—The granulomatous inflammation in PHIRs is a nonneoplastic inflammatory reaction with a variable lymphocytic component. Granuloma formation can be seen in both reactive inflammatory infiltrates and in cutaneous involvement of leukemias and lymphomas. Leukemia cutis has been reported in 4% to 20% of patients with CLL/small lymphocytic leukemia. In one series of 42 patients with CLL, the malignant cells were confined to the site of postherpetic scars in 14% (6/42) of patients. Sixteen percent (7/42) of patients had no prior diagnosis of CLL at the time they developed leukemia cutis, including one patient with leukemia cutis in a postzoster scar. The mechanism involved in the accumulation of neoplastic lymphocytes within postzoster scars has not been fully characterized. The idea that postzoster sites represent a site of least resistance for cutaneous infiltration of CLL due to the changes from prior inflammatory responses has been proposed.

Combined CLL and granulomatous dermatitis at prior sites of herpes zoster was first reported in 1990. In 1995, Cerroni et al reported a series of 5 patients with cutaneous CLL following herpes zoster or herpes simplex virus infection. Three of those patients also demonstrated granuloma formation. Establishing a new diagnosis of CLL from a biopsy of postzoster granulomatous dermatitis with an associated lymphoid infiltrate also has been reported. Cerroni et al postulated that cutaneous CLL in post-herpes zoster scars may occur more frequently than reported due to misdiagnoses of CLL as pseudolymphoma. Two additional cases of postherpetic cutaneous CLL and granulomatous dermatitis have been reported since 1995.

Diagnosis of Multiple PHIRs—The presence of 3 concurrent PHIRs is rare. The patient in this report had postzoster cutaneous CLL with an associated granulomatous dermatitis and medium-vessel vasculitis. One other case with these 3 findings was reported by Elgoweini et al. Overlooking important diagnoses when multiple findings are present in a biopsy can lead to diagnostic delay and incorrect treatment; we highlighted the importance of careful examination of biopsies in PHIRs to ensure diagnostic accuracy. In cases of postzoster granulomatous dermatitis, assessment of the lymphocytic component should not be overlooked. The presence of a dense lymphocytic infiltrate should raise the possibility of a lymphoproliferative disorder such as CLL, even in patients with no prior history of lymphoma. If initial immunostaining discloses a predominantly B-cell infiltrate, additional immunostains (eg, CD5, CD23, CD43) and/or genetic testing for monoclonality should be pursued.

Conclusion
Clinicians and dermatopathologists should be aware of the multiplicity of postherpetic isotopic responses and consider immunohistochemical stains to differentiate between a genuine lymphoma such as CLL and pseudolymphoma in PHIRs with a lymphoid infiltrate.

REFERENCES