Adult-onset hydroa vacciniforme-like lymphoma in a long-term resident of the United States

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INTRODUCTION

Hydroa vacciniforme-like lymphoproliferative disorders (HVLPDs) arise from chronic active Epstein-Barr virus (EBV) infection of T cells and natural killer cells. The term HVLPD was introduced in the 2016 World Health Organization classification of lymphoid neoplasms1 to reflect the spectrum of EBV-related cutaneous disorders from classic hydroa vacciniforme (HV) to severe HV and HV-like lymphoma (HVLL). Although classic HV has no geographic predilection, severe HVLPDs (severe HV and HVLL) are endemic among children and young adults in Asia and Latin America.2-4 Here we present an unusual case of severe HVLPD that arose in a Hispanic adult resident of the United States with no recent travel to endemic areas or childhood history of HV. With shifting patterns of global migration, it is important for American physicians to be familiar with this disease spectrum.

CASE

A 33-year-old man had a 2-year history of diffuse eruptions of pruritic, painful papules over his entire body. The eruptions occurred periodically in association with sunlight and heat. Over the past year, they became more frequent with greater skin involvement and associated with fevers. He was originally from Mexico but immigrated to the United States >10 years ago and has not left this country since his arrival. He has no prior history of HV or other photodermatoses. Examination revealed cervical lymphadenopathy, splenomegaly, and numerous erythematous-to-brown papules with central necrosis and crusting over his trunk, face, and extremities distributed on a background of white, stellate scars (Fig 1).

Biopsies of the papules demonstrated a wedge-shaped, superficial, and deep dermal perivascular and periaxial infiltrate of small atypical lymphocytes and eosinophils with necrosis of the epidermis and upper dermis (Fig 2, A and B). There was no vasculitis. The presence of numerous eosinophils was suggestive of a hypersensitivity reaction, such as to arthropod bites. The atypical small lymphocytes expressed CD2, CD3, CD7, CD8, and T-cell intracytoplasmic antigen 1. Rare CD56+ lymphocytes were seen. Epstein-Barr encoding region (EBER) positivity was detected in a subset of the atypical lymphocytes by EBER in situ hybridization. T-cell receptor gene rearrangement studies by PCR demonstrated clonal rearrangements of the γ and β chains. Further evaluation to classify his HVLPD included

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flow cytometry, EBV PCR, and positron emission tomography—computed tomography scan. The patient had detectable levels of EBV DNA (53,900 copies/mL) in blood. Imaging revealed hypermetabolic cutaneous nodules on his head, neck, trunk, and scrotum and lymph nodes throughout the neck, thorax, abdomen, and pelvis. Due to concern for HVLL, the patient was referred to hematology-oncology for further management.

Six weeks after his initial presentation, the patient experienced 3 days of fever (102°F), nausea, vomiting, and the eruption of new skin lesions. He was admitted to a hospital to rule out infection or hemophagocytic syndrome, which is a known complication of severe HV and HVLL. After extensive work-up, hemophagocytic syndrome was ruled out, and his symptoms were determined to be due to the progression of HVLPD. Interferon α-2b was initiated at 2 million units 3 times per week, which resulted in rapid defervescence and gradual resolution of his skin lesions.

The patient has continued to follow up with hematology-oncology on an outpatient basis. Interferon α-2b was titrated up to 7.5 million units 3 times per week with good control of his skin lesions. However, his disease burden on imaging is unchanged and his EBV DNA titers remain positive. We discussed the possibility of hematopoietic stem cell transplant to provide a potential long-term cure, but the patient has deferred this option due to lack of insurance coverage.

**DISCUSSION**

HVLPD encompasses a range of clinical entities that typically occur in children and young adults. They are linked by evidence of chronic active EBV infection and common histopathologic features, including epidermal vesicles and reticulate degeneration, with dermal perivascular lymphocytic infiltrates (Table I). Severe HV extends to sun-protected skin and is often accompanied by systemic symptoms. Up to half of cases of severe HV progress to HVLL, a systemic malignancy that follows a rapidly progressive and usually fatal course. This case describes the development of HVLPD in a man with no history of HV in childhood. His skin lesions did not correspond with a photodistribution and were accompanied by lymphadenopathy and fever. This constellation of symptoms raised concern for severe HVLPD and led to referral to hematology-oncology, where he was found to have lymphoma on the basis of findings from imaging (Table I).

In HVLPD, repeated exposure to ultraviolet light is thought to cause continual activation of EBV and draw infected T cells and natural killer cells to sun-exposed skin. Geographic variation in the incidence of severe HV and HVLL has been attributed to local differences in host human leukocyte antigen classes, EBV subtypes, and environmental factors, such as timing of initial EBV exposure.

Severe HV and HVLL are managed medically and require close follow-up with hematology-oncology to monitor for disease progression. Severe HV has a chronic waxing and waning course, making it difficult to predict if and when lymphoma will evolve. Iwatsuki et al have proposed the following indicators of severe HV progression: facial swelling with eruptions, systemic symptoms, hypersensitivity to insect bites, increased levels of EBV DNA in peripheral blood or EBER-positive cells on biopsy, and presence of cellular atypia and deep infiltrates. Although there are no randomized trials to guide treatment, immunomodulatory agents, including thalidomide, chloroquine, and interferon α, have emerged as first-line therapy for severe HV. Anthracycline-based chemotherapy, such as CHOP (cyclophosphamide, vincristine, doxorubicin, and prednisolone), can induce temporary remissions, but case series suggest higher than expected morbidity and mortality. Lastly, there is emerging...
Fig 2. Histologic examination of specimen from patient with hydroa vacciniforme-like lymphoproliferative disorder. A, Superficial and deep, wedge-shaped infiltrate with epidermal and superficial dermal necrosis. B, Infiltrate comprised of small atypical lymphocytes admixed with eosinophils. (A and B, Hematoxylin-eosin stain; original magnifications: A, ×40; B, ×200.)

Table I. Spectrum of HV-like lymphoproliferative disorders2-4,6,7*

| Category                  | Classic HV                                      | Severe HV                        | HVLL                           |
|---------------------------|-------------------------------------------------|----------------------------------|--------------------------------|
| Skin lesions              | Papulovesicular eruptions that crust and heal with depressed, stellate vacciniform scars | Necrotic papulovesicular lesions with extensive vacciniform scarring |                                |
| Distribution              | Photodistributed                                | Exposed and unexposed skin       |                                |
| Histopathology            | Reticular epidermal degeneration with spongotic vesiculation, perivascular lymphocytic infiltrates in superficial to deep dermis | Dense perivascular lymphocytic infiltrates extending to deep dermis and hypodermis |                                |
| Cellular atypia           | None                                            | +/−                              | Yes                            |
| Cell types                | CD4+ or CD8+ T cells                            | CD4+ or CD8+ T cells or CD56+ NK cells |                                |
| EBER positivity           | <25%                                            | Variable, but often >25%         |                                |
| T-cell receptor           | Polyclonal                                      | Monoclonal                       |                                |
| Systemic findings         | None                                            | Fever, lymphadenopathy, elevated liver enzymes, facial edema, hypersensitivity to mosquito bites, hemophagocytic syndrome | Fever, lymphadenopathy, weight loss, elevated liver enzymes, elevated lactate dehydrogenase, NK cell lymphocytosis, leukopenia, thrombocytopenia |
| EBV DNA in peripheral blood| Elevated                                       | Very elevated (>10,000 copies/mg) |                                |
| EBV antibody titers       | Usually normal                                   | Demonstrate chronic active EBV infection |                                |
| PET findings              | Normal                                           | Hypermetabolic foci              |                                |
| Prognosis                 | Spontaneous remission                           | Limited to cutaneous lymphoma or progression to systemic lymphoma |                                |
| Treatment                 | Sun avoidance, sun protection                   | Immunomodulatory therapy,†       | Immunomodulatory therapy,†     |
|                           |                                                  | chemotherapy‡                   | chemotherapy‡, allogenic stem cell transplant |

*Adapted from Ko et al, with addition of features further defining severe HV and HVLL.
†Interferon α-2b, thalidomide, or chloroquine.
‡Cyclophosphamide, doxorubicin, vincristine, and prednisolone with or without etoposide.
evidence that allogenic stem cell transplant, which was offered to our patient because he did not respond completely to immunomodulatory therapy, might benefit HVLPD patients who relapse or fail to respond to first-line agents.8

Given the large number of immigrants to the United States, American dermatologists will see domestic cases of HVLPD and must be familiar with its diagnosis and treatment.

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