To the Editor:

Graft-versus-host disease (GVHD) is a common and serious complication seen most often with bone marrow transplantation and peripheral blood stem cell transplantation. With these therapies, functional lymphoid cells are transferred from an immunocompetent donor into a non-genetically identical recipient, or “host.” Because of the allogeneic nature of these transplants, the transplanted lymphoid cells have a high potential to recognize and treat the host’s cells as foreign, and the resultant clinical and pathologic picture is that of GVHD. The primary organ systems affected in this immune response are the skin, gastrointestinal tract, and hepatobiliary system.1,2 Cutaneous manifestations are by far the most common.3

Although notable gains have been made in elucidating the causes, risk factors, and mechanisms that result in the clinical picture of GVHD, gaps in our knowledge and understanding still exist. Our patient represents a unique case of unilateral GVHD occurring along Blaschko lines, which has important implications for both recognizing and understanding the pathogenesis of GVHD.

A 35-year-old woman was diagnosed with stage IV follicular lymphoma and received various chemotheraphy regimens over the next 4 years. Unfortunately, her disease progressed despite treatment. At 39 years of age, she underwent a nonmyeloablative allogeneic peripheral blood stem cell transplantation from a single HLA-mismatched sibling. She was placed on prednisone and cyclosporine for immunosuppression. High-dose acyclovir prophylaxis also was initiated given her history of zoster affecting the right C3 dermatome. Successful engraftment was achieved, with molecular studies showing 100% of cells following transplantation were of donor origin. Restaging at 1 and 2 years following transplantation found her to be in complete remission.

At 2 years following transplantation, she began a slow taper of immunosuppressive medications. She was successfully weaned off prednisone and continued to gradually reduce the cyclosporine dose. Toward the end of the cyclosporine taper 3 months later, she developed a pruritic eruption on the left proximal arm.

She was seen in a bone marrow transplant clinic 4 weeks after the rash developed. On examination, she had multiple, violaceous, lichenoid papules coalescing into linear band-like plaques. One plaque extended along the left upper arm and 2 others encircled the left hemithorax, respecting the midline. She was treated empirically for zoster with valacyclovir 1 g 3 times daily based on the presumed dermatomal distribution of the eruption. Despite treatment, the rash progressed, and she developed fever. Eight days later, she was admitted with concern for disseminated zoster (Figure). Viral tissue cultures and polymerase chain reaction analysis of the lesions were negative for varicella-zoster virus and herpes simplex viruses 1 and 2. Biopsies of skin lesions on the arm and trunk were both consistent with GVHD.

Given the clinical history, characteristic lesion morphology, and distinct linear distribution along with

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**Practice Points**

- Recognizing the characteristic manners in which different linear dermatoses present can aid in correctly identifying disorders that most commonly present in either a dermatomal or Blaschko-linear-type distribution.
- A blaschkoid-type distribution is the result of cutaneous mosaicism that occurs during embryological development and therefore subsequently produces a unique phenotypical presentation for various genetically influenced skin disorders, including graft-versus-host disease.
By histopathological confirmation, a diagnosis of GVHD along Blaschko lines was made. Recognizing the cause to be immunogenic rather than infectious, immunosuppressive medications were started. In addition to increasing the prednisone and cyclosporine back to therapeutic levels, she received weekly methylprednisolone. With treatment, she showed gradual but marked improvement.

Six cases of linear GVHD have occurred as an isotopic response along dermatomes previously affected by varicella-zoster virus. These cases give credence to the idea that a cutaneous viral infection may alter the skin through unknown mechanisms, predisposing it to become affected by GVHD. Notably, this phenomenon occurred despite absence of a persistent viral genome when assessed using polymerase chain reaction analysis.

An additional 3 cases of GVHD occurring in a dermatomal distribution without any prior infections in those areas have been reported. Of note, 2 of 3 patients did have episodes of zoster occur at other sites following transplantation and did not develop GVHD symptoms in any of those locations. Interestingly, controversy exists as to whether the distribution of these lesions was dermatomal or followed Blaschko lines.

Two cases of linear GVHD have been reported in which lesions were identified as occurring along Blaschko lines. The lines of Blaschko, first described in 1901, correspond to cellular migration patterns during embryological development. Postzygotic mutations causing epidermal cell mosaicism may result in skin disorders occurring in segmental areas defined by the Blaschko lines. Accordingly, the Blaschko-linear pattern in GVHD suggests cellular mosaicism as the etiology in this case. Although the host’s immune system develops immunotolerance to both cellular lineages during maturation, transplanted lymphoid cells from a nongenetically identical sibling may identify just one of the cell lines as nonself, producing a selective pattern of GVHD confined to the distribution of the genetically disparate cell line, which occurs along the lines of Blaschko in the skin. Candidate genes for mutations that would produce a mosaic following transplant GVHD include any of the 25 to 30 known minor histocompatibility antigens (or any of the several hundred yet to be found).

Although well established for monogenic dominant disorders, in 2007 it was recognized that a postzygotic mutation can cause many complex polygenic disorders, including GVHD, to manifest in a limited segmental pattern. This understanding, along with retrospective case review, has brought into question previously reported “dermatomal” or “zosteriform” presentations of GVHD, asserting that the linear patterns were misidentified and thus inappropriately attributed to a postviral response. Recognition of the Blaschko-linear distribution holds significance in both identifying lesion etiology and understanding disease pathogenesis and treatment.

Our patient illustrates a case of a Blaschko-linear GVHD. The distinctive pattern of her physical findings strongly favored epidermal cell mosaicism as the etiology of her disease. More than just a phenotypically unique case, it provided further insight into the complex etiology underlying GVHD and iterated the basic concepts of Blaschko lines and genetic alterations in development.

REFERENCES