A Case of Systemic Lupus Erythematosus with Cutaneous Leukocytoclastic Vasculitis Mimicking Bullous SLE

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Abstract
Rarely, patients with systemic lupus erythematosus (SLE) develop bullous eruptions, a disease called bullous SLE in a narrow sense that has autoantibodies against type VII collagen. We describe an unusual case in which a patient with SLE developed extensive bullae on her lower extremities. Histologically, the bullous lesions were suggestive of leukocytoclastic vasculitis with deposition of C3 within blood vessel walls. Immunoblot analyses and enzyme-linked immunosorbent assays were negative for anti-type VII collagen antibodies. We initially considered bullous SLE, but eventually made a diagnosis of secondary vasculitis in SLE. The oral prednisolone dose was increased, and the vesiculobullous lesions resolved. The clinical presentations of cutaneous vasculitis in SLE include palpable purpura, petechiae, papulonodular lesions, and livedo reticularis. Bullous lesions seem to be uncommon. Physicians need to be aware that extensive bullae can occur as a result of secondary vasculitis in SLE, even if the patient does not exhibit high disease activity.

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Introduction

It is well known that vasculitis can occur in systemic lupus erythematosus (SLE) patients, but very few studies have specifically examined vasculitis in SLE [1–3]. Thus, there is not much information about the clinical manifestations of vasculitis in SLE. Vasculitis presents in different clinical forms, depending on the size of the affected blood vessels and the organs involved [4, 5]. Cutaneous lesions, which can manifest as palpable purpura, urticaria, petechiae, papulonodular lesions, livedo reticularis, cutaneous infarctions, erythematous plaques, erythema with necrosis, panniculitis, splinter hemorrhages, and/or superficial ulceration, are the main clinical symptoms [1, 2]. Bullous lesions seem to be uncommon. Here, we describe a rare case of SLE with secondary vasculitis, in which the patient developed extensive bullous skin lesions on her lower extremities, and we considered bullous SLE and contact dermatitis as potential diagnoses. The absence of antibodies against type VII collagen ruled out bullous SLE in a narrow sense. Extensive bullous lesions caused by secondary vasculitis have rarely been reported in a patient with SLE.

Case Report

A 33-year-old female had been diagnosed with SLE 10 years ago, when she had developed a fever, a skin rash, arthralgia, anemia, and positivity for anti-dsDNA antibodies (27 U/mL). She was followed up by an internist while being treated with 10 mg/day oral prednisolone, 6 mg/day methotrexate, and 3 mg/day tacrolimus for several years. As the arthralgia recurred, treatment with 200 mg/day hydroxychloroquine was added. One month later, she developed a skin rash on her right lower extremity, and the hydroxychloroquine treatment was discontinued. However, the skin rash spread further, so she was referred to our department 5 days later. On examination, she had multiple vesiculobullous eruptions and erythematous macules on her lower extremities (Fig. 1a–c). She also developed erythematous macules on her upper arms (Fig. 1d). Her laboratory data, including her complete blood count, liver and renal function, C-reactive protein, complement component 3 (C3) and C4 levels, CH50 test results, and erythrocyte sedimentation rate, were all within the normal range. Tests for the following antibodies produced negative results: anti-nuclear, anti-ss DNA, anti-dsDNA, anti-RNP, anti-Sm, anti-SSA/Ro, anti-SSB/La, and anti-cardiolipin antibodies. Tests for PR3-ANCA and MPO-ANCA also produced negative results. Histopathologically, a biopsy from a vesicular skin lesion on her right lower extremity demonstrated edematous changes and superficial perivascular inflammatory infiltrates, consisting of neutrophils and lymphocytes, in the upper dermis, as well as fibrinoid blood vessel necrosis with nuclear debris and extravasated erythrocytes (Fig. 2a, b). The erythema on the upper arms demonstrated similar histological findings, although it did not exhibit edematous changes. Direct immunofluorescence of skin samples from both lesions showed the deposition of C3 (Fig. 3a) and fibrinogen within blood vessel walls. Indirect immunofluorescence of both normal skin and 1 M NaCl-split skin showed negative results. Immunoblot analyses of normal human epidermal extracts, recombinant bullous pemphigoid (BP)180 NC16a and BP180 C-terminal domain proteins, concentrated culture supernatant of HaCaT cells, normal human dermal extracts (Fig. 3b), and purified human laminin-332 all showed negative results. In addition, ELISA was negative for desmoglein 1, desmoglein 3, BP180, BP230, and type VII collagen. A diagnosis of leukocytoclastic vasculitis secondary to SLE was made. The prednisolone dose was increased to 25 mg/day, and the vesiculobullous lesions and erythema resolved. The prednisolone dose was then tapered and maintained at 10 mg/day for 1 year. No recurrence has since been observed.
Discussion

Vasculitis is an uncommon, but serious, manifestation of SLE [5]. The prevalence of vasculitis in SLE is reported to range from 11% [2] to 36% [1] (visceral and/or cutaneous), while the prevalence of cutaneous vasculitis was reported to range from 10% [2] to 17.3% [3]. Episodes of vasculitis are known to often occur during lupus flare-ups accompanied by symptoms, such as fever, fatigue, and weight loss [4]. In addition, a previous study demonstrated that compared with SLE patients without vasculitis, patients with vasculitis had a higher mean European Consensus Lupus Activity Measurement (ECLAM) score (5.86 vs. 3.87; \( p < 0.001 \)) [2]. However, it is worth noting that episodes of vasculitis do not always occur during periods of high disease activity [4], as was seen in our case.

In 2 large cohort studies of SLE patients with vasculitis, the most common type of vasculitis was small vessel vasculitis (as was seen in our case), which is a type of leukocytoclastic vasculitis that can be limited to the skin [1,2]. The clinical presentation of vasculitis in SLE is heterogeneous: the most common skin manifestation is erythematous punctate lesions on the fingertips and palms (36%), followed by palpable purpura (25%), ischemic lesions and/or ulcers (14%), erythematous papules/macules (14%), urticarial lesions (11%), and nodular lesions (5%). Bullous lesions, however, seem to be extremely uncommon. To the best of our knowledge, only 1 case of SLE in which the patient developed severe widespread bullae with histological evidence of leukocytoclastic vasculitis has been reported [6]. However, in the latter case, the cutaneous bullae arose after acute steroid withdrawal, which is different from our case [6].

Up to 29% of patients develop a combination of the skin manifestations of vasculitis [2]. Our patient also had 2 different types of lesions: erythematous macules on her upper arms and vesiculobullous lesions on her lower legs.

The differential diagnoses in our case included bullous SLE, a blistering disorder that occurs in patients with active SLE, which is characterized clinically by widespread vesiculobullous skin lesions, histopathologically by dermal-epidermal separation in the basement
membrane zone, and immunologically by immunoglobulin deposition in the basement
membrane zone in a linear, granular, or mixed linear and granular pattern (shown by direct
immunofluorescence) [7]. In bullous SLE, the upper dermis is characterized by edema and the
accumulation of neutrophils in dermal papillae, which are similar to the findings seen in
dermatitis herpetiformis. In addition, patients with bullous SLE in a narrow sense have auto-
antibodies to type VII collagen. Another type of bullous lesion that can arise in SLE is epidermal
elevation due to vacuolar degeneration of the basal layer and dermal edema [8]. We were able
to rule out bullous SLE and contact dermatitis based on the histopathological, immunofluo-
rescence, immunoblotting, and ELISA findings.

As the clinical presentations of the cutaneous lesions of SLE vary significantly, it is
sometimes difficult to determine whether cutaneous lesions are related to SLE and/or
high SLE disease activity. Physicians need to be aware that extensive bullae can occur as
a result of secondary vasculitis in SLE, even if the patient does not exhibit high disease
activity. However, since previous studies of vasculitis in SLE were largely based on the
clinical manifestations of the condition [3] with only small numbers of vasculitis cases
being confirmed by histopathology [1,2], clinical symptoms like those seen in our case might
be underreported.
Statement of Ethics

This research complies with the guidelines for human studies and was conducted ethically in accordance with the World Medical Association Declaration of Helsinki. Written informed consent was obtained from the patient for publication of the details of her medical case and any accompanying images.

Conflict of Interest Statement

The authors have no conflicts of interest to declare.

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Author Contributions

F.M. conceived of and designed the study, analyzed and interpreted the data, and wrote the manuscript. K.O. was responsible for the data analysis and interpretation. T.H. contributed to the data acquisition, analysis, and interpretation and wrote the manuscript. H.A. was responsible for the data interpretation and approved the manuscript.

Data Availability Statement

All data generated or analyzed during this study are included in this article. Further enquiries can be directed to the corresponding author.

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