Drug-induced hypersensitivity syndrome in a patient with systemic lupus erythematosus and psoriasis

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Sir,

Co-existence of systemic lupus erythematosus (SLE) and psoriasis is relatively rare. We herein describe a rare case of drug-induced hypersensitivity syndrome (DIHS) in a patient with SLE and psoriasis vulgaris, induced by an anticonvulsant.

A 39-year-old man was diagnosed with SLE at the age of 28. Seven years later, he developed psoriasis (Figs. 1a and 1b), and was treated with topical corticosteroids and vitamin D3 ointment. Moreover, he had aortic regurgitation (AR), and underwent aortic valve replacement. After the operation, he developed convulsions, and carbamazepine (Tegretol®; 200 mg/day) was administered. Four weeks later, he developed itchy eruptions on the trunk and extremities, with a fever reaching up to 40°C. On physical examination, erythema multiforme-like lesions were spread on the trunk and extremities (Fig. 1c). Tegretol® was discontinued, but one week later, erosions of the oral mucosa appeared (Fig. 1d), and the erythema on the trunk coalesced with purpura. Inguinal lymphadenopathy was observed. Laboratory data showed a normal white blood cell count (6300/μl), with a differential count of 58% neutrophils and 21% eosinophils, elevated AST (66 IU/l) and ALT (116 IU/l) levels. Serum anti-nuclear antibody was positive (1:320, homogenous), whereas anti-DNA antibody was within normal ranges and hypocomplementemia was not detected. A biopsy specimen from the abdomen revealed apoptotic keratinocytes and perivascular inflammatory cell infiltration composed of mononuclear cells and eosinophils in the edematous upper dermis (Fig. 2). Before the onset of drug eruption, the patient’s SLE was treated with oral prednisolone (12.5 mg/day); however the eruption was improved by escalating the dose of prednisolone to 25 mg/day. A lymphocyte transformation test using Tegretol® was weakly positive (S.I. 188%), and HHV-6 IgG titer (1:10) was elevated (1:640) six weeks later. HLA-A and -B genotyping revealed HLA-A*1101, A*3101, B*1501, and B*3501.

The co-existence of psoriasis and SLE is sometimes seen, and both diseases share some common pathogenesis, such as Th1/Th17 type-dominant cytokine imbalance, plasmacytoid dendritic cell activation via Toll-like

Figure 1: Clinical pictures at initial visit to our department showing malar rash and keratotic plaques on the forearm (a, b). Drug eruption showing erythema multiforme-like erythemas scattered on the trunk (c), facial erythema and mucosal erosion (d).
receptors, and interferon-α release. Our patient developed psoriasis seven years after the diagnosis of SLE. Furthermore, he had AR and received operation at the remission phase of SLE. Four weeks after administration of an anticonvulsant drug, he developed DIHS, which fulfilled the Japanese criteria, without exacerbation of either SLE or psoriasis. It is known that several autoimmune disorders, including SLE, are induced after DIHS was improved [1,2], possibly due to functional defects of regulatory T cells (Tregs) [3]. However, our case developed SLE before the occurrence of DIHS, and thus the development of SLE was unassociated with DIHS. Patients with SLE have an increased risk of developing adverse cutaneous drug eruptions [4], and cases such as toxic epidermal necrolysis (TEN) and Stevens-Johnson syndrome (SJS) rarely occurs in patients with SLE [5].

To date, only one suspected case of SLE has been reported to develop drug reaction with cosinophilia and systemic symptoms [6]. HLA-A*3101 is significantly associated with susceptibility to carbamazepine-induced adverse drug reactions in Japanese population [7]. Our case also had HLA-A*3101, therefore may have developed DIHS unassociated with SLE itself.

**Consent**

The examination of the patient was conducted according to the Declaration of Helsinki principles.

**REFERENCES**