Bullous pemphigoid successfully treated with omalizumab

Sir,
Bullous pemphigoid is an acquired, autoimmune, bullous disease.[1-4] The pathogenic autoantibodies are usually of the IgG subtype, but some patients with bullous pemphigoid have IgE autoantibodies against type XVII collagen, which have been shown to be pathogenic.[1,3] Ten cases of bullous pemphigoid treated with omalizumab, an anti-IgE monoclonal antibody, have been reported in the last few years.[1-5] Herein, we present a patient with bullous pemphigoid, who had a very high IgE level and who was treated successfully with omalizumab.

A 70-year-old man, weighing 70 kg, presented with multiple, tense bullae and eroded areas on the trunk and extremities, particularly on the knees, hands and feet [Figure 1] and several oral erosions. The Nikolsky sign was negative. The skin lesions were itchy and had gradually progressed over the previous 3 months. In addition, he had psychosis and was positive for anti-hepatitis C virus antibody. Investigations revealed a markedly raised IgE level (2500 IU/mL, N: 0–87 IU/mL) and eosinophilia (2.9%, N: 0.03–0.4%). Hepatitis C viral RNA load was 400,000 IU/mL. Histopathological examination showed sub-epidermal separation and accumulation of eosinophils and neutrophils in the bulla cavity [Figure 2a]. On direct immunofluorescence with fluorescein isothiocyanate labeled anti-C3 and anti-IgG antibodies, there was linear staining of the dermoepidermal zone [Figure 2b]. Bullous pemphigoid was diagnosed based on the clinical, histopathological and direct immunofluorescence findings. His lesions did not respond to 6 weeks of treatment with topical clobetasol propionate 0.05% and oral tetracycline (4 × 500 mg/day). Hence, these were stopped and oral corticosteroid therapy (1 mg prednisolone/kg/day) was started. Three weeks later, dapsone 100 mg/day was added as new lesions continued to appear. However, dapsone therapy had to be discontinued because of an increase in transaminase levels. His psychosis also deteriorated, most likely because of the systemic steroids. Though disease activity persisted, we were hesitant to step up his immunosuppressive medication because that would require prior treatment of his hepatitis C infection with interferon which was very likely to exacerbate his psychosis. Hence, additional immunosuppressive agents were not considered. Based on the available reports, omalizumab therapy was started at a dose of 300 mg subcutaneously every 4 weeks as an alternative treatment. New lesions stopped appearing within 1 week of treatment initiation and eroded areas gradually re-epithelialized [Figure 3]. Several days after the omalizumab injections, thrombocytopenia was detected (80,000/uL, N: 150,000–372,000/uL), but this did not require cessation of the drug and the platelet count gradually improved. When omalizumab became unavailable for 2 months, after the seventh dose, his bullous pemphigoid lesions exacerbated, but the lesions disappeared with re-administration of omalizumab. The platelet count fell minimally after each omalizumab injection. Although the platelet count gradually increased between successive injections, it never reached the normal range. The hepatitis C viral RNA load was stable. Till date, 11 injections of omalizumab 300 mg/injection have been given to the patient. The immunoglobulin E level was found to be decreased to 851 IU/mL just before the 12th dose. His bullous...
pemphigoid is in remission with oral prednisolone in a dose of 5 mg/day. Stoppage of oral steroids followed by a reduction of omalizumab dosing to 150 mg every 4 weeks is planned, if the patient stays in remission.

The antigens targeted by the pathogenic antibodies responsible for bullous pemphigoid are the 230-kDa bullous pemphigoid antigen (BP-230) within basal keratinocytes and the 180-kDa bullous pemphigoid antigen, type XVII collagen (BP-180), within the basement membrane zone.[1,2] However, in mouse models, it has been shown that IgG antibodies alone do not account for all the features of bullous pemphigoid, as dermal eosinophil infiltration and purely spontaneous blistering could not be induced by IgG antibodies.[1] Moreover, the pathogenicity of IgE autoantibodies in humans has been shown and the first case of bullous pemphigoid treated with omalizumab was reported by Fairley et al. in 2009.[3]

The mechanism of action of omalizumab, a recombinant humanized IgG1 monoclonal antibody against human IgE, is not completely understood in the treatment of bullous pemphigoid. However, it is thought that omalizumab prevents binding of IgE to its receptor and hence inhibits the activation of mast cells that are increased in lesions of bullous pemphigoid. In addition, omalizumab downregulates IgE receptors and circulating eosinophils. Another hypothesis is that omalizumab is able to induce eosinophil apoptosis and the downregulation of pro-inflammatory cytokines.[2]

The standard therapy for severe bullous pemphigoid is systemic steroids plus a potentially steroid-sparing agent such as azathioprine, mycophenolate mofetil and methotrexate. Compared with these agents, omalizumab has a relatively selective effect.[1] Therefore, it may be a favorable alternative therapy for bullous pemphigoid patients, especially those in whom immunosuppressive therapy is contraindicated, as in our patient. The hepatitis C viral RNA load of our patient has stayed stable during 9 months of omalizumab therapy. Improvement of the lesions on omalizumab therapy, exacerbation of the lesions when omalizumab became unavailable for 2 months and the disappearance of the lesions with re-administration are all supporting evidence for the efficacy of omalizumab therapy in bullous pemphigoid. Moreover, our observation shows that 300 mg omalizumab monthly may be an effective therapy for bullous pemphigoid even if the IgE level is very high. This is in contrast to previous reports, in which the omalizumab doses were adjusted according to body weight and baseline IgE levels of the patients.[1-3] Its onset of action is rapid and our patient improved within 1 week.

Adverse effects of omalizumab treatment include headache, pyrexia, infections, abdominal pain and also thrombocytopenia. Data about omalizumab-induced thrombocytopenia are limited and conflicting.[5] Although our patient had a normal platelet count (168,000/uL) before omalizumab therapy, thrombocytopenia occurred 5 days after injection of the drug. Platelet counts ranged between 110,000 and 130,000/uL during treatment but these values did not require the cessation of omalizumab.

In summary, omalizumab may be an effective and relatively safe therapeutic option for a subset of patients with bullous pemphigoid who do not respond to or have contraindications to standard treatment. Its onset of action is rapid and 300 mg/month may be sufficient to suppress blistering in most patients. Thrombocytopenia is a known side effect and complete blood counts need to be checked regularly during treatment.

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Sir,

Dermatophytosis is among the most common skin diseases affecting millions of people worldwide. We are facing an unprecedented increase in the number of recurrent tinea infections in our daily practice. In the absence of susceptibility tests and studies, it is difficult to comment whether these recurrences represent true resistance to common antifungals or are due to other reasons.

A 23-year-old man presented with a 2-year history of recurrent, reddish itchy lesions involving the face, neck, trunk, lower extremities, gluteal and inguinal regions. Initially, the lesions were limited to the gluteal and inguinal regions and used to resolve following treatment with oral fluconazole and topical steroid-antifungal combination therapy. During subsequent episodes, similar lesions appeared on the trunk, lower extremities and face; these were treated with over-the-counter medications, only to recur. He underwent therapy with systemic terbinafine and topical eberconazole for 2 weeks at a local hospital and achieved complete remission but his lesions recurred 1 week after completing the course. He was otherwise healthy and denied any history of recurrent bacterial or viral infections. There was no history of atopy, diabetes or usage of immunosuppressive drugs. No history of contact with domesticated or wild animals was present and there was no other significant family history.

At presentation, he had multiple erythematous papules and a few plaques with minimal scaling over the lower abdomen, bilateral groins, upper thigh, buttocks [Figure 1] and a few on the right mandibular angle, right side of the neck and on the left lower extremity [Figure 2]. Routine laboratory investigations were within normal limits. Septate and branching hyphae were observed on direct microscopic examination (potassium hydroxide mount) of scales obtained by scraping the lesions [Figure 3] and fungal culture showed the growth of *Trichophyton rubrum*.

The patient was prescribed oral itraconazole and topical eberconazole followed by oral ketoconazole and topical amorolfine for 2 weeks each, but the lesions recurred after partial resolution. Finally, oral isotretinoin (20 mg/day) and itraconazole (200 mg/day) along with topical sertaconazole was given for a period of 1 month. Following this treatment, the skin lesions resolved completely [Figure 4] and a repeat potassium hydroxide mount showed no fungi. A follow-up examination 6 months later showed no recurrence of lesions.

In the absence of antifungal susceptibility testing on isolated strains of fungi, it is inappropriate to label this patient as a case of drug-resistant dermatophytosis. The mechanism of therapeutic success with isotretinoin,