CASE REPORT

Omalizumab as a Successful Therapy in Normocomplementemic Urticarial Vasculitis: A Series of Four Patients and Review of the Literature

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Urticarial vasculitis is an eruption characterized by inflamed itchy or painful red papules or plaques that resemble urticaria but last longer than 24 hours and heal with residual pigmentation or purpura. Histopathologically, urticarial vasculitis presents as leukocytoclastic vasculitis with perivascular infiltrate and fibrin deposits. The treatment options are oral antihistamines, oral corticosteroids, dapsone, colchicine and hydroxychloroquine. We report four cases with normocomplementemic urticarial vasculitis who were treated with omalizumab and a brief review of the literature on the use of omalizumab in normocomplementemic urticarial vasculitis.

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-Keywords-
Omalizumab, Urticarial vasculitis

INTRODUCTION

Urticarial vasculitis (UV) is a clinicopathological entity that is characterized by the coexistence of urticaria with leukocytoclastic vasculitis, perivascular infiltrate of lymphocytes, neutrophils and eosinophils, as well as fibrin deposits that are histopathologically typical features of leukocytoclastic vasculitis. Symptoms of UV are frequently pain and burning rather than itching and resolve with persistent hyperpigmentation. UV can be classified into normocomplementemic UV (NUV) and hypocomplementemic UV (HUV) depending on the levels of complement protein in blood. Recommended treatments for UV are oral antihistamines (OAH) and systemic immunosuppressant drugs such as oral corticosteroids, colchicine, dapsone and hydroxychloroquine. Omalizumab is a recombinant humanized monoclonal anti-immunoglobulin (IgE antibody that has been approved for the treatment of chronic urticaria. There have been recent reports on the use of omalizumab for the treatment of UV.

Here we describe four cases of UV treated with omalizumab with persuasive results. We also review previous reports of patients with UV treated with omalizumab.

CASE REPORT

Case 1

A 41-year-old female patient presented with pruritic and painful skin rash typical of urticaria on her upper and lower extremities for the last year. The lesions were erythematous infiltrated wheals lasting for longer than 24 hours that resolved with postinflammatory hyperpigmentation (Fig. 1). Histopathologic examination of the skin revealed leukocytoclastic vasculitis (Fig. 2). The patient was treated with OAH for a month without improvement which was then followed by colchicine 0.5 mg three times a day. After three months, the patient experienced relapse of skin rashes and was then treated with hydroxychloroquine 200 mg twice daily for three months without significant im-
provement and switched to dapsone 50 mg twice a day. After unsuccessful treatment with dapsone for 8 months, omalizumab 300 mg subcutaneously (s.c.) once every four weeks was initiated. After the first application, complete remission of the UV was achieved with a urticaria control test (UCT) score of 16. The patient is still under treatment with omalizumab 300 mg s.c. every four weeks with sustained remission and no apparent adverse effect after 8 injections. We received the patient’s consent form about publishing all photographic materials.

**Case 2**

A 34-year-old female patient with a 2-year history of UV was previously treated with OAH for at least 6 months without improvement. Oral prednisone 16 mg/day had achieved remission but with early relapse after drug cessation. After colchicine 1.5 mg/day started, the condition had worsened. Omalizumab 300 mg s.c. once every four weeks was initiated which provided complete remission of the UV after the first injection and UCT scored as 15.

She remained stable after 12 injections of omalizumab 300 mg monthly and the treatment switched to omalizumab 300 mg per 6 weeks. The patient is still under treatment with sustained remission.

**Case 3**

A 26-year-old male patient with a 3-year history UV was previously treated with OAH at least 6 months without improvement which was then followed by cyclosporine 100 mg twice a day. After one month of cyclosporine treatment, the patient experienced exacerbation of skin lesions. Omalizumab 300 mg s.c. once every four weeks was initiated which provided complete remission of the UV after the first application and UCT scored as 16. The patient is still under treatment with omalizumab 300 mg s.c. every four weeks with sustained remission after 14 injections.

**Case 4**

A 58-year-old female patient with one-year history of UV,

| Case no. | Age (yr) | Sex | DD (mo) | Angioedema | Biopsy | DIF | C3 | ANA | Previous drugs (mg/d) | NOI | TCR (wk) |
|---------|---------|-----|---------|------------|--------|-----|----|-----|------------------------|-----|----------|
| 1       | 41      | F   | 12      | No         | LV     | N/A | N  | (−) | Rupatadin 10            | 8   | 4        |
|         |         |     |         |            |        |     |    |     | Colchicine 1.5          |     |          |
|         |         |     |         |            |        |     |    |     | Hydroxychloroquine 400  |     |          |
|         |         |     |         |            |        |     |    |     | Dapsone 100             |     |          |
| 2       | 34      | F   | 24      | Yes        | LV     | Positive C1q+ | N  | (−) | Cetirizin 10           | 12  | 4        |
|         |         |     |         |            |        |          |    |     | Prednisone 16           |     |          |
|         |         |     |         |            |        |          |    |     | Colchicine 1.5          |     |          |
|         |         |     |         |            |        |          |    |     | Levocetirizin 5         |     |          |
|         |         |     |         |            |        |          |    |     | Cyclosporine 200        |     |          |
| 3       | 26      | M   | 36      | Yes        | LV     | N/A  | N  | (−) | Evacetazone 10          | 14  | 4        |
| 4       | 58      | F   | 12      | Yes        | LV     | Positive C3+C1q+ | N  | (−) | Omalizumab 300 mg s.c.   | 5   | 4        |

DD: duration of disease, DIF: direct immunofluorescence examination, ANA: antinuclear antibody, NOI: number of omalizumab injections, TCR: time to complete response, F: female, M: male, LV: leukocytoclastic vasculitis, N/A: nonavailable, N: normal.
was previously treated with OAH at least 6 months without improvement. According to our previous experiences with UV, the patient was started on omalizumab 300 mg s.c. every four weeks. After the first application, complete remission of the UV was achieved and UCT score increased from 3 to 12. The patient is still treated with omalizumab 300 mg s.c. every four weeks with sustained remission after 5 injections.

The diagnosis of UV in all of the above cases was made based on the clinical, histopathological and direct immunofluorescence examination findings (Table 1).

**DISCUSSION**

UV is a leukocytoclastic vasculitis with wheals that resemble urticaria but last more than 24 hours. It represents a continuum of disease, ranging from urticaria with minimal vasculitis, to life- or organ-threatening systemic vasculitis with minimal urticaria.

UV can be divided into NUV and HUV depending on the levels of complement protein in blood. HUV has systemic manifestations such as angioedema, arthralgia, fever, abdominal or thoracic pain, renal or pulmonary disease, episcleritis and uveitis. NUV usually presents with less systemic symptoms and without underlying diseases therefore are considered as idiopathic. NUV might also be considered as urticaria showing signs of vasculitis in histopathology because in a series of 83 patients with chronic urticaria 10 exhibited histologic features of UV.

Omalizumab has been designed to recognise and attach to a specific structure (C3 domain) on circulating human IgE. This prevents IgE binding to high affinity receptors (FcεRI) on the surface of mast cells and basophils, thus reducing inflammatory cell activation including B-cells. It also causes eosinophil apoptosis, increases interleukin (IL)-2, IL-3, interferon-gamma and tumor necrosis factor-alpha and reduces IL-4. The mechanism of action of omalizumab in UV is not fully understood, it may be associated with downregulation of FcεRI receptors and suppression of proinflammatory cytokines and therefore interfering the generation of urticarial wheals. Blocking anti-IgG IgEs which are responsible for the development of immune complexes on the vessel walls may also be considered as a possible mechanism of action. In summary, omalizumab may be effective in UV through its reduction of IgE and complement levels in our patients were normal, therefore they are considered as NUV. The efficacy of omalizumab in these patients is not unexpected when we consider NUV as “urticaria with minimal vasculitis.”

HUV syndrome (HUVS) is a rare immune complex mediated disease which has distinct pathophysiology and outcomes. Aurich et al. reported that omalizumab is not effective in HUVS. This suggests the pathophysiology of HUVS is significantly different from chronic spontaneous urticaria (CSU) and NUV.

Omalizumab has been shown to be effective on the symptoms of UV in case reports but there are no prospective clinical studies of omalizumab for UV (grade D recommendation). A case report by Ghanizafar and Thomsen described a male patient with UV who failed to respond to oral prednisolone and dapsone therapy. He was then treated with omalizumab 300 mg once every four weeks and after one month a complete remission of the disease and symptoms were observed. Ramirez Del Pozo et al. reported a female patient with UV and systemic lupus erythematosus. The patient was unresponsive to OAH, oral corticosteroids and azathioprine and was started on omalizumab which provided a rapid relief of symptoms.

Another case report by Varriachi et al. described a female patient with Churg-Strauss syndrome, asthma and UV. She was treated with OAH, oral corticosteroids and azathioprine and was started on omalizumab 300 mg s.c. every two weeks as an addition to her pre-treatments. The patient reported a significant improvement after 6 months of treatment with omalizumab. A case report from Spain describing three female patients with CSU difficult to control with associated vasculitis noted that omalizumab was a successful treatment option for the patients in the study. Lately, Fueyo-Casado et al. reported a female patient with ten year history of chronic recurrent UV. She was treated with OAH at least 6 months, oral prednisolone and oral cyclosporine without improvement. Dapsone and antileucotrienies had only been minimally effective. She was then started on omalizumab 300 mg s.c. every 4 weeks which showed clinical improvement within the first month and complete remission within the fifth month.

Omalizumab 300 mg per a month provided symptom control in our cases within one month with a notable increase in UCT scores. Some authors use 300 mg per a month and others 300 mg per 6 weeks but there is no consensus on the dose and intervals; but applying the same dose and interval scheme as CSU seems rationale. Our report is important because we noted complete remission of the UV lesions within one month. It shows that mechanism of action of omalizumab in NUV may be similar to its mechanism of action in CSU.

As many patients with UV experience major side effects of the standard treatment and cannot be treated successfully, omalizumab gives us promising results with its safety pro-
file in the treatment of NUV.

In conclusion, omalizumab may be useful in the treatment of vasculitic urticaria, especially with normal complement levels. This observation needs to be validated with prospective, randomized, placebo-controlled trials.

CONFLICTS OF INTEREST

The authors have nothing to disclose.

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