Combined Infliximab and Rituximab in Necrotising Scleritis

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Abstract

Introduction

Granulomatosis with polyangiitis (GPA), previously known as Wegener’s granulomatosis, is a multi-system autoimmune disease, often associated with anti-neutrophil cytoplasm antibodies (ANCA), which commonly involves the eye as well as the respiratory tract and kidneys. Ophthalmic complications occur in 50–60% of patients and typically take the form of necrotising scleritis or granulomatous orbital involvement [1]. Ophthalmologists need to be aware that these conditions are potentially sight-threatening – some 90% of patients with necrotising scleritis develop ocular complications and 50% develop significant visual loss [2]. Suboptimal treatment of either condition can lead to serious visual consequences.

Cyclophosphamide and corticosteroids have been the cornerstones of treatment for severe GPA but are associated with treatment-related morbidity and mortality [3]. More recent treatment options include the biological agent rituximab, which is a monoclonal antibody directed against CD20, a cell surface antigen expressed by many subtypes of B cells, although not by plasma cells. We have recently shown that rituximab can be effective in refractory ophthalmic GPA and induce long-lasting disease...
remission, but it takes up to 6 months to work, meaning that additional agents may be required to provide disease control in the interim [4].

Infliximab belongs to a different class of biological agents, being a monoclonal antibody directed against tumour necrosis factor-alpha (TNF-α). It is licensed for the treatment of both rheumatoid arthritis and Crohn’s disease in the United States and Europe [5] and has also demonstrated efficacy in the treatment of the ANCA-associated vasculitides (AAV), such as GPA, although a recent study in which infliximab was added to standard therapy in patients with active AAV failed to demonstrate any benefit compared to standard therapy alone [6]. Infliximab has been used successfully in the treatment of refractory scleritis [7, 8].

We report here a patient with necrotising scleritis associated with GPA in whom infliximab was used for short-term steroid-sparing while rituximab took effect. This enabled disease control without requiring an extended period of high-dose corticosteroid administration or the concurrent use of cyclophosphamide, neither of which was desirable as a result of his comorbidities and previous treatment history.

Case Report

A 71-year-old man was referred to the Hammersmith Hospital Multidisciplinary Vasculitis Clinic with necrotising scleritis in the left eye (fig. 1). He had previously been diagnosed with GPA on the basis of characteristic renal disease associated with a raised ANCA titre and the presence of anti-PR3 antibodies. He had previously been treated with cyclophosphamide, and his cumulative dose of cyclophosphamide was approximately 20 g. He was initially treated with corticosteroids at a dose of 1 mg/kg and a cycle of rituximab (MabThera, Hoffmann-La Roche, Ltd.) given as 2 intravenous doses of 1 g two weeks apart. After 3 weeks, his necrotising scleritis was still active, despite his B cell count having fallen to 0 cells/mm³. We were reluctant to continue his high-dose corticosteroids as he had osteoporosis confirmed on DEXA scanning, and treatment with cyclophosphamide was not felt desirable in view of his cumulative dose. Therefore, infusions of infliximab (Remicade, Merck & Co.) were arranged and given as doses of 5 mg/kg at baseline, 2, 6 and 12 weeks. His necrotising scleritis was controlled by week 6 (fig. 2), and his systemic corticosteroids were reduced to 5 mg prednisolone/day by week 12. He had his final infusion of infliximab at this point and thereafter was maintained on 5 mg prednisolone/day alone for the following 12 months.

Discussion

We report a case of necrotising scleritis secondary to GPA, successfully treated with infliximab whilst waiting for rituximab to induce disease remission. Importantly, this enabled cyclophosphamide to be avoided in the acute phase, as rituximab can take up to 6 months to work [4, 5]. This was especially important as the patient had had significant previous exposure to cyclophosphamide, increasing his risk of malignancy with further treatment. Infliximab rather than etanercept was used as the latter has previously been proven not to be effective for the maintenance of remission in patients with GPA [5, 9]. In addition, retrospective studies of patients with uveitis treated with TNF-α blockade have indicated infliximab to be the more effective treatment in ophthalmic disease [10].

There was some concern that the use of two biological agents simultaneously could lead to life-threatening infection or malignancy, but there is little in the literature that
addresses this directly. One recent registry-based study of patients being treated with biological agents for rheumatoid arthritis showed their rate of hospitalised infection to be 4.6 per 100 person-years, increasing to 7.0 per 100 person-years for the group that had ‘switched’ from a different biological agent [11], the situation most analogous to the one here. Interestingly, the rate for rituximab was higher at 10.4 episodes per 100 person-years in this study, falling to 7.1 in switchers; the authors suggested that this might represent higher-risk patients being channelled towards rituximab therapy. Encouragingly for the treating clinician, analysis showed that patient characteristics were a more powerful determinant of the rate of serious infections than specific biological drugs or patterns of switching drugs. Similarly, a meta-analysis of patients treated with TNF-α blockade in rheumatoid arthritis did not demonstrate an increased risk of malignancy [12].

Nevertheless, this case report illustrates the potential usefulness of infliximab in gaining rapid disease control and allowing time for rituximab to take effect, at least in some patients with necrotising scleritis. It also illustrates the increasing treatment options which are available to treat patients with refractory ocular inflammation, and supports the management of such patients in a multidisciplinary setting so that close contact with appropriate specialists can enable these treatments to be taken advantage of by patients in a safe and controlled environment.

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Fig. 1. An active area of necrotising scleritis is visible in the superotemporal quadrant of the left eye, despite 1 mg/kg/day prednisolone and 2 recent doses of rituximab.

Fig. 2. Six weeks after the institution of infliximab therapy, the necrotic area can be seen to have revascularised and healed.
References