Intravenous immunoglobulin therapy for asthma: time for a closer look?

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The need for a pooled intravenous immunoglobulin preparation capable of passive transference of broad based humoral immunity was first recognised following the description of congenital agammaglobulinemia by Bruton. It was subsequently recognised that this treatment, originally intended to restore immune deficiency, actually appeared to have therapeutic effects in diseases involving immune effector mechanisms, most notably "autoimmune" diseases such as immune thrombocytopenia. Intravenous immunoglobulin therapy is currently being evaluated for a possible therapeutic benefit in many other such diseases including rheumatoid arthritis, systemic lupus erythematosus, systemic vasculitis, myasthenia gravis, and others.

It is increasingly recognised that asthma is associated with chronic, cell-mediated inflammation of the bronchial mucosa in which cytokine products of activated T cells, especially those which are implicated in selective eosinophilia, accumulate, play a prominent role. Evidence suggests that glucocorticoids ameliorate asthma at least partly through inhibition of activated T cells and elaboration of their cytokine products. For this reason there has been interest in the investigation of other "anti-inflammatory" or "immunosuppressive" agents for their possible therapeutic benefits in asthma. Since many of these agents have potentially serious unwanted effects, attention has generally been focused on those asthmatic patients who continue to have severe disease despite properly administered, maximal topical glucocorticoid therapy and additional continuous systemic therapy. With such patients it is perceived that the benefits of amelioration of the disease and reduction or abolition of systemic glucocorticoid therapy, with its well recognised hazards, might outweigh the risks.

The possible efficacy of intravenous immunoglobulin therapy for asthma has been investigated in two open uncontrolled studies on children. Mazer and Gelfand treated eight children who required continuous systemic glucocorticoid therapy for asthma control with intravenous immunoglobulin in a dose of 1 g/kg as a 6% solution on and a beneficial effect was noted. The mean serum total IgE concentrations were significantly reduced. The mean serum total IgE concentrations before and after four doses of intravenous immunoglobulin at an intended dosage of 1 g/kg on a single day monthly for five months. Treatment with the high dosages used by Mazer and Gelfand was considered "too expensive". This dosage had to be decreased to 0.5 g/kg after the first infusion because it caused severe headache, and then progressively increased (mean dosage 0.8 g/kg). Even at these lower dosages, infusions were frequently accompanied by fever and rigors. During the infusion period six patients were able to reduce dosages of intravenous immunoglobulin from a mean of 720 mg/day to 400 mg/day) whereas three were not. Histamine bronchial reactivity was also reduced (mean PC20 0.33–1.23 mg/ml), as were total symptom scores. These differences, by comparison with the "reference" group, were not maintained after 10 additional months of follow up. Mean serum IgG concentrations before (11.6 g/l) and four months after (11.4 g/l) the infusions were almost identical, as were serum total IgG concentrations. In addition, serum concentrations of IgG anti-IgE antibodies and eosinophil cationic protein were unchanged. These authors concluded that the effects of intravenous immunoglobulin therapy were "small and temporary, and the treatment complicated and expensive". They also hypothesised that some of the improvement observed in the study of Mazer and Gelfand might have reflected the effects of immunoglobulin replacement, since they considered the mean serum IgG concentration before treatment in these children (5.85 g/l) to be abnormally low, and a beneficial effect of intravenous immunoglobulin therapy had previously been demonstrated in asthmatic children with hypogammaglobulinaemia.

A further open study of the effects of intravenous immunoglobulin therapy in two severe glucocorticoid-dependent adult asthmatic patients is reported in this issue of Thorax. These patients were labelled "glucocorticoid insensitive" since they had severe airways obstruction (with high spontaneous variability) despite high dosages of systemic glucocorticoids with a poor clinical response to further escalation of therapy. In addition, peripheral blood T cells from these patients were said to show "decreased sensitivity" to dexamethasone in vitro. Treatment of both patients with intravenous immunoglobulin (2 g/kg on a single day monthly on four or six occasions) allowed a marked reduction in the dose of oral glucocorticoids accompanied by improvements in FEV1 and variability of PEF. In one of these patients bronchial biopsy specimens taken before and after four doses of intravenous immunoglobulin showed reductions in the numbers of T cells and cells expressing the T cell activation marker CD25 in
the bronchial mucosa, while flow cytometric analysis of peripheral blood T cells showed reductions in the percentages of both CD4 and CD8 cells expressing the activation markers CD25 and HLA-DR. The serum concentration of total IgE was reduced, while that of total IgG was increased. In addition, these authors showed increased binding of the cytokine IL-8 to IgG autoantibodies in the serum of this patient following treatment. The serum concentration of eosinophil cationic protein was also reduced. Again, unwanted effects of therapy were presumably sufficiently mild as not to merit specific documentation.

How might intravenous immunoglobulin exert an anti-asthma effect? Mazer and Gelfand postulated that intravenous immunoglobulin therapy might represent a form of passive immunotherapy, interfering with IgE-mediated reactions. In support of this, both they and Jakobsson and colleagues demonstrated reduced immediate skin prick test reactivity to a variety of allergens in their patients following intravenous immunoglobulin therapy, although this was not accompanied by RAST inhibition in vitro or evidence of increased binding of circulating IgE to IgG autoantibodies. The mechanism of this phenomenon, as well as its possible relevance to amelioration of asthma, therefore remains obscure. Intravenous immunoglobulin has been shown to abrogate activation of both T and B cells in vitro. Furthermore, soluble CD4, CD8, and HLA molecules have been identified in intravenous immunoglobulin preparations which may act to inhibit presentation of antigen to T cells. Circumstantial evidence for inhibition of T cell activation by intravenous immunoglobulin in vivo has been provided by the present study of Vrugt and colleagues, although it would be unwise to infer a “cause and effect” relationship with asthma amelioration from observations on a single patient. Finally, this study raises the interesting possibility that intravenous immunoglobulin preparations might contain cytokine autoantibodies, which might inhibit the activities of these cytokines on inflammatory effector cells such as eosinophils.

It is self-evident that uncontrolled, open label studies such as those discussed here are impossible to interpret since they do not allow for the power of the placebo effect and take no account of spontaneous variability in asthma severity. Furthermore, little is known about possible spontaneous temporal variability of inflammatory cell numbers and their activation states in the peripheral blood and bronchial mucosa of asthmatic subjects. Nevertheless, these studies arguably justify the need for larger placebo controlled trials of the possible benefits of intravenous immunoglobulin therapy in glucocorticoid dependent asthma. It is to be hoped that the cost of such trials does not prove prohibitive.
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