Responses to Biological Therapy in Severe Eosinophilic Asthma

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LETTERS TO THE EDITOR

To the Editor:

Severe asthma often remains uncontrolled despite high doses of inhaled corticosteroids and a second controller drug [1,2]. Biological therapy reduces the frequency of severe asthma exacerbations (SAEs) and improves lung function. Consequently, it is a promising therapeutic option. The biologics currently indicated in severe asthma include the following: omalizumab, which blocks IgE; mepolizumab and reslizumab, which target free serum IL-5; benralizumab, which depletes eosinophils by blocking the IL-5 receptor; and dupilumab, which blocks the IL-4/IL-13 axis. All of these agents are more effective in asthma characterized by type 2 inflammation and eosinophilic asthma. However, even in these phenotypes, individual responses vary and are unpredictable. Therefore, patients should be reevaluated during the first year of treatment and continue with treatment only if the response is adequate [3]. The National Institute for Health and Care Excellence defines an adequate response as a ≥50% reduction in the frequency of SAEs or in the corticosteroid dose needed to maintain control [3].

We report the response to biological therapies in 2 patients with severe asthma [1,2]. Given that the presentation of asthma was similar in both cases similarity of their asthma, we initially assumed that the response to therapy would be comparable.

Patient 1 was a 51-year-old woman with a 30-year history of asthma and rhinosinusitis with nasal polyps. She had had adrenal insufficiency for 3 years and required daily treatment with methylprednisolone (5 mg). She had experienced 3 episodes of eosinophilic pneumonia. Total serum IgE was 78-422 kU/L, and her blood eosinophil count varies from 100 to 300/μL. The Asthma Control Test score ranged from 11 to 25. The Asthma Control Test score ranged from 11 to 25.

Patient 2 was a 51-year-old man with a 15-year history of asthma and rhinosinusitis with nasal polyps. Total serum IgE was 202-330 kU/L, sputum eosinophilia was 5%-75%, and the blood eosinophil count was 300-800/μL. The Asthma Control Test score ranged from 8 to 25.

Neither patient was obese (body mass index, 29 and 28). Tests for dyspnea and anxiety were always negative. Both patients were sensitized to perennial allergens (Patient 1 to Cupressus pollen and animal dander and Patient 2 to Dermatophagoides pteronyssinus), had documented bronchial reversibility, and tolerated nonsteroidal anti-inflammatory drugs. They were taking fluticasone (1000 μg), formoterol (40 μg), and tiotropium (5 μg), and adherence to treatment was good. We calculated the annual average number of SAEs by multiplying the number of SAEs by 12 and dividing the result by the number of months in which they were recorded. The duration of treatment was different for each patient depending on the clinical decisions made (Figure). In both cases, 1 year of therapy with omalizumab failed to reduce the number of SAEs (from 4/y to 6.67/y in Patient 1 and from 6.0/y to 7.2/y in Patient 2).

After completion of therapy with omalizumab, Patient 1 experienced an episode of eosinophilic pneumonia. Since starting mepolizumab (100 mg/4 wk) in October 2017, she has been symptom-free, with no SAEs and a 340-mL increase in mean FEV1 value.

Six months after discontinuation of omalizumab, Patient 2 initiated mepolizumab at 100 mg/4 wk. After the first 4 months, SAEs reappeared with their usual frequency. Consistent with data reported elsewhere [4], we decided—together with the patient, the Ethics Committee, and the Pharmacy Department—to increase the dose of mepolizumab to 200 mg/2 wk. During the following 10 months, the SAE rate dropped to 4.8/y (33%), although the patient complained of muscle pain that interfered with his activity. In August 2018, we stopped mepolizumab and started intravenous, weight-adjusted reslizumab (30 mg/4 wk). The initially milder muscle symptoms disappeared, and the SAE rate dropped to 2.4 (66%). However, since initiation of anti–IL-5 therapy, his mean FEV1 values dropped by 237 mL and mean exhaled nitric oxide (ENO) values increased from 70 ppb to 106 ppb. Moreover, SAEs became less sensitive to systemic corticosteroids, thus necessitating longer treatment regimens (from 5 to 10 days).

Given that both patients had undergone conventional treatment and had similarly high eosinophil counts and SAE rates, they were candidates for anti–IL-5 therapy. However, their response to mepolizumab differed radically. Mepolizumab was completely successful in Patient 1, who remains symptom-free after 1 year and has improved lung function. Patient 2 experienced an initial 4-month improvement with mepolizumab, although the frequency of SAEs returned to pretherapy values. In premarketing trials, intravenous mepolizumab at 750 mg and 75 mg reduced the...
SAEs and blood eosinophil counts. Nevertheless, the 75-mg dose was significantly less effective in reducing sputum eosinophil counts [5] and was equivalent to the subcutaneous 100-mg dose that was eventually marketed [6]. It is possible that the administration of antibody at suboptimal doses leads to immune-complex formation that would constitute a local reservoir of IL-5, thus perpetuating bronchial eosinophilic inflammation [7]. Therefore, we increased subcutaneous mepolizumab to 200 mg/2 wk. The frequency of SAEs dropped by 33%, although the patient experienced disabling muscular symptoms.

Intravenous reslizumab also blocks free IL-5; however, its dose is adjusted for body weight, thus enabling higher doses. Since both dosing and route of administration can modify the clinical response to anti–IL-5 therapy [4,7], we changed high-dose mepolizumab for intravenous reslizumab. After 8 months of treatment, the frequency of SAEs fell by 66%. However, we consider this response to be unsatisfactory, as both mean ENO and FEV₁ values worsened after initiation of anti–IL-5 therapy and daily bothersome symptoms of asthma reappeared.

It is speculated that respiratory epithelium can be so damaged in some patients with severe asthma that its response to stimuli involves release of epithelial-derived alarmins (thymic stromal lymphopoietin, IL-33, IL-25) that can promote local eosinopoiesis, thus rendering systemic anti–IL-5 therapies ineffective [7]. In the near future, we will try to control asthma in Patient 2 by completely blocking eosinophils with benralizumab.

ENO values were much higher in Patient 2 and may account for the difference in response to anti–IL-5 therapy. However, Patient 1 was taking daily oral corticosteroids, to which ENO is highly sensitive. Since ENO production is associated with type 2 and epithelial cells, dupilumab, which blocks the IL-4/IL-13 axis, could be an alternative in Patient 2, although we do not know the effect of the initial dupilumab-induced increase in eosinophil count [8]. In theory, we could combine dupilumab and benralizumab.

In contrast with clinical trials, where inclusion criteria are well controlled, real-world practice has not yet yielded sufficient data to generate indicators that predict the response to biological therapy for severe asthma. Blood eosinophils, ENO values, and SAE rate are weak markers that do not take into account the heterogeneity of asthma or the not necessarily parallel change in its different facets (functional, clinical, inflammatory), as was the case in Patient 2, who experienced functional deterioration despite experiencing fewer SAEs. The lack of data enabling us to presume the superiority of any antiasthma biologic means that we need to obtain such information empirically from daily clinical practice. We are therefore indebted to the patients we treat. According to Drazen and Harrington [9], it is essential to have independent comparative studies performed by well-established public institutions in which therapies are supplied cost-free by pharmaceutical companies. Only then can we be sure to prescribe the most appropriate treatment. Finally, the lower sensitivity of Patient 2 to corticosteroids for treatment of SAEs
should be assessed in the context of eosinopenia induced by anti–IL-5-therapy and should lead us to revisit the role of oral corticosteroids [10].

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**Conflicts of Interest**

MJ Zavala reports personal fees from GSK. JM Olaguibel reports personal fees from AstraZeneca and GSK and grants from Sanofi. The remaining authors declare that they have no conflicts of interest.

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