ABSTRACT

Purpose: Several markers for eosinophilic inflammation have been proposed to predict response to asthma treatment. However, definitive criteria for treatment decisions have not yet been established. We investigate a potentially useful relatively non-invasive biomarker, eosinophil-derived neurotoxin (EDN), to predict favorable responses to budesonide or montelukast, common treatment for children with asthma.

Methods: Young children (1 to 6 years old) were enrolled in this randomized, parallel, 2-group, open-label trial. Criteria for eligibility included: 1) being symptomatic during the run-in period; and 2) having a serum EDN (sEDN) level ≥ 53 ng/mL, with positive specific immunoglobulin E to house dust mite. Eligible patients were randomly placed into 2 groups: the BIS group received budesonide inhalation suspension (BIS) 0.5 mg once daily; the MONT group received montelukast 4 mg once daily. Ineligible patients were invited to receive montelukast 4 mg once daily (OBS group). Treatment period was 12 weeks.

Results: Asthma control days increased significantly in the BIS and MONT groups (P < 0.000) over the 12-week study period. There was no significant change in sEDN in the BIS group but there was a significant decrease in the MONT group (P < 0.000). Patients in the OBS group with high EDN levels (> 53 ng/mL) showed a significant decrease due to MONT treatment (P = 0.023). Rescue medication usage significantly decreased in the BIS and MONT groups (P < 0.000).

Conclusions: EDN is a useful relatively non-invasive biomarker for predicting responses to montelukast and budesonide treatment of preschool children with beta2-agonist responsive recurrent wheeze and multiple-trigger wheeze (Trial registry at UMIN Clinical Trials Registry, UMIN000008335).

Keywords: Biomarkers; children; eosinophil-derived neurotoxin; asthma

INTRODUCTION

Chronic airway inflammation is a hallmark of asthma; consequently, inhaled corticosteroids (ICSs) are the main treatment. Although leukotriene receptor antagonists (LTRAs) also have anti-inflammatory properties and are often administered, especially to young children, guidelines clearly state ICSs are the preferred option for long-term control of asthma.
preference is based mainly on evidence from clinical trials comparing mean responses between treatment and control groups. However, other trials have demonstrated that there is considerable inter-individual variability in treatment response to ICSs versus LTRAs.\textsuperscript{4,6} It is important for a clinician to collect information that better guides them in selecting medication most likely to achieve a favorable response in each patient (\textit{i.e.}, tailor-made treatment regimens).

Several markers for eosinophilic inflammation or atopy, such as elevated levels of serum immunoglobulin E (IgE) due to house dust mite (HDM), \textit{Dermatophagoides pteronyssinus} (Dp), sensitivity,\textsuperscript{7} sputum eosinophil counts,\textsuperscript{8} exhaled nitric oxide\textsuperscript{9} and serum eosinophil cationic protein (ECP),\textsuperscript{10} have been suggested to predict response to ICSs. However, definitive criteria for treatment decisions are still to be established. In addition, characteristics of airway pathology in preschool children may be different from those in older children and adults,\textsuperscript{11} which may lead to differential treatment responses.

In addition to ECP, elevated levels of another eosinophil degranulation product, eosinophil-derived neurotoxin (EDN), have been noted in asthmatics, with higher levels also found during asthma exacerbation when compared to healthy patients and those with stable asthma.\textsuperscript{12} Therefore, EDN has been suggested as a key marker for disease activity of asthma and other eosinophil-related disease.\textsuperscript{13}

To date, there are no published studies comparing differential responses to the 2 types of anti-inflammatory therapies, namely ICSs and LTRAs, for preschool children with asthma. Furthermore, clinical markers for these possible differential responses have not been defined. Here, we propose a study to seek a useful and relatively non-invasive marker (\textit{i.e.}, serum EDN [sEDN]) to predict favorable responses to nebulized budesonide or montelukast, common treatment options for young children with asthma.

**MATERIALS AND METHODS**

**Subjects**

Children aged 1 to 6 years were enrolled in this study. Because asthma is a heterogeneous disease and especially difficult to diagnose in this age group, the definition of asthma used was based on the Global Initiative for Asthma (GINA)\textsuperscript{2} and Japanese pediatric guidelines.\textsuperscript{14} Major inclusion criteria were as follows: 1) 3 or more recurrent wheezing episodes within a year, with at least one confirmed by a physician, and multiple-trigger wheeze (not just common cold but other triggers such as exercise, crying or laughing); and 2) improvement of wheezing after administration of inhaled or oral short-acting beta\textsubscript{2}-agonists. Minor inclusion criteria include at least one of the following: 1) parental asthma, 2) physician-diagnosed atopic dermatitis (AD), 3) physician-diagnosed allergic rhinitis (AR) or 4) physician-diagnosed food allergy. To be included in the study, a patient must have had both major criteria and at least one of the minor inclusion criteria. Criteria for randomization included patient was: 1) symptomatic during the run-in period; and 2) had a sEDN level $\geq 53$ ng/mL,\textsuperscript{13} with positive specific IgE to HDM (Immulite 2000 3gAllergy $> 0.1$ IU/mL).\textsuperscript{14} Exclusion criteria included one or more of the following: 1) patients who received asthma controller medication, such as ICSs or LTRAs, in the previous 6 months; 2) chronic lung disease, heart or kidney disease or any other congenital disease; and 3) a patient whom the physician regarded as inappropriate for the study (\textit{e.g.}, did not attend the clinic when instructed).
**Study Design**

This was a randomized, parallel, 3-group, open-label trial. Patients who met the enrollment criteria entered a 2-week run-in period, with short-acting beta2-agonists as needed. Participants were then randomly selected to join 1 of the 2 groups: the BIS group received budesonide inhalation suspension (BIS) 0.5 mg once daily for 12 weeks and the MONT group received montelukast 4 mg once daily (orally) for 12 weeks. BIS was administered via a SkyNeb (Medel, Milan, Italy) compressor and nebulizer with a mouthpiece (for older children) or face mask (for children 1-3 years old). Patients that were ineligible for the trial because randomization criteria 2) was not met, EDN < 53 ng/mL and/or negative specific IgE to HDM, yet had symptoms in the run-in period were invited to receive montelukast 4 mg once daily based on recommendation by the Japanese guideline.\(^{14,15}\) They were observed for 12 weeks as exploratory purpose (OBS group). Parents/caregivers of all participants were instructed to record all symptoms and treatment times in an asthma diary. Patients were evaluated at 4-week intervals (Fig. 1) in either a hospital or clinic to have their asthma diary checked and to determine whether a step-up in therapy was needed as per GINA and Japanese guidelines.

Computer randomization of an eligible patient into either the BIS or MONT group was performed. However, adjustments were made so that there was a statistically similar proportion of patients in each group based on age (1-2 years old versus 3-5 years old), sex and specific IgE to HDM (< 10 IU/mL versus ≥ 10 IU/mL).

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**Fig. 1.** Study protocol.

At visit 1, patients who met inclusion criteria were tested for sEDN and HDM-IgE, then entered to run-in period. After 2 weeks of run-in period, the results of sEDN and HDM-IgE were reported to the study physician and the patients who met randomization criteria were randomized to receive budesonide or Montelukast for 12 weeks. Those who did not meet randomization criteria, yet had symptoms during run-in period were invited to receive montelukast. Clinical assessment was made at each visit and with asthma symptom diary recorded by the caregivers. At visit 5, sEDN was measured.

sEDN, serum eosinophil-derived neurotoxin; HDM-IgE, house dust mite-immunoglobulin E; R, randomization.
Caregivers provided written informed consent on behalf of patients. This clinical trial was registered at UMIN Clinical Trials Registry: http://www.umin.ac.jp/ctr/index.htm (UMIN000008335).

EDN measurement
EDN levels in each patient were measured twice: at −2 weeks (during the run-in period) and at the 12-week time point. EDN was measured using an enzyme-linked immunosorbent assay (ELISA) following the manufacturer’s protocol (MBL, Woburn, MA, USA), and the results were expressed in nanograms per milliliter (ng/mL). This ELISA detects human EDN with a minimum detection limit of < 0.62 ng/mL, a maximum detection limit > 300 ng/mL, and no cross-reaction with ECP. All assays were run in duplicate, and the mean value was used for statistical analyses.

Virus detection
At initial enrollment and when exhibiting asthma or upper respiratory infection symptoms at regular visits, nasopharyngeal swabs were collected and total RNA from common respiratory viruses was extracted and identified using a multiplex PCR system (Seeplex RV12 Ace Detection; Seegene, Seoul, Korea) that identifies adenovirus (AdV) A-E, coronavirus (CoV) 229E/NL63/OC43, parainfluenza virus (PIV) 1-4, rhinovirus (RV) A-C, influenza virus (IFV) A and B, respiratory syncytial virus (RSV) A and B, bocavirus (BV) 1-4, metapneumovirus (MPV), and enterovirus.

Assessment of outcome
Primary outcome was the number of asthma control days (ACD). An ACD was defined as a day with no beta-agonist rescue use; use of oral corticosteroids for asthma; daytime symptoms; nighttime awakenings; unscheduled health care visits, emergency department visits, or hospitalizations for asthma; or school absenteeism for asthma. ACD in both groups (BIS and MONT) were compared in 2 ways: 1) within group analyses comparing ACD at the beginning of the treatment period (0 week) to ACD at the other 3-time points (4, 8 and 12 weeks); and 2) between group analyses comparing ACD in each group at each time point (e.g., BIS ACD at 0 week versus MONT ACD at 0 week). An ACD value was calculated for each group during each time period. For example, ACD at 0 week was calculated during the 2-week run-in period (i.e., −2 to 0 week), ACD at 4 weeks was calculated during the time period from 0 to 4 weeks, etc. The first comparison used 2-way analysis of variance (ANOVA), followed by Sidak’s multiple comparisons test. The second comparison used the Wilcoxon matched-pairs signed rank test.

A secondary outcome of this study was to compare changes in sEDN in all patients with sEDN ≥ 53 ng/mL and HDM sensitivity (BIS and MONT groups) and in those patients whose sEDN was < 53 ng/mL, were not sensitive to HDM or both (OBS group). Serum was taken from each patient at −2 weeks, sEDN was measured, and then this value was used at 0 week (termed “Pre”) to randomize each patient into either the BIS or the MONT group. Also, sEDN was measured at 12 weeks (termed “Post”). “Pre” and “Post” sEDN levels were then compared in each patient using the Wilcoxon matched-pairs signed rank test.

Another secondary outcome of this study was changes in rescue medication usage with treatment. Comparisons were done in 3 ways: 1) within each group (BIS and MONT), usage at run-in (−2 weeks) was compared to usage at the end of the treatment period (12 weeks); 2) each subject in each group was analyzed over the study period (−2 to 12 weeks) to determine
any change in usage; 3) usage in BIS group was compared to usage in MONT group at each
time point (−2, 0, 4, 8 and 12 weeks). For each endpoint, stratified analysis was performed
based on the virus type detected.

**Statistical analysis**
Within group analyses of ACD at multiple time points during the entire study period (−2 to
12 weeks) were compared using 2-way ANOVA, followed by Sidak’s multiple comparisons
test. This was only done for the BIS and MONT groups. Between-group analyses of ACD
(BIS versus MONT groups) were done using the Wilcoxon matched-pairs signed rank test.
Within group analyses of treatment effect comparing “Pre” (measured during run-in period
−2 to 0 week) and “Post” (measured at 12 weeks) measurements of changes in sEDN for all 3
groups (BIS, MONT and OBS) were done using the Wilcoxon matched-pairs signed rank test.
Each group was then subgrouped according to presence or absence of any virus and change
in sEDN with treatment from “Pre” to “Post” was analyzed using the same statistical test.
Rescue medication usage within each group (BIS and MONT) and between each group were
also compared. Data that were normally distributed were presented as mean values, unless
otherwise noted. *P* values < 0.05 were considered statistically significant.

**RESULTS**

Flow chart for screening and randomization into study groups (BIS and MONT) is in **Fig. 2.**

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**Fig. 2.** Patient flow.
R, randomization; BIS, budesonide inhalation suspension; MONT, eligible and receiving montelukast; OBS, not
eligible and receiving montelukast; EDN, eosinophil-derived neurotoxin; HDM, house dust mite.
Demographics

ANOVA followed by the Bonferroni multiple comparison test were used to determine any statistical differences in demographic data among the 3 groups (i.e., BIS, MONT, and OBS; Table 1). Age ($P < 0.05$) and the proportion of those with AD ($P < 0.05$) in the OBS group were significantly lower compared to the other groups. The age difference between the OBS groups and the other 2 groups was expected as subjects in the BIS and MONT groups were age-matched. All other demographic variables were statistically similar among the 3 groups.

Laboratory data

Laboratory data at entry can be found in Table 2. HDM-IgE and send levels in the OBS group were significantly lower when compared to the other 2 groups. This was expected because to be included in the OBS group a patient had to have a sEDN level < 53 ng/mL, no sensitivity to HDM or both. The proportion of the MONT group (77.3%) with a detected virus was significantly greater ($P = 0.005$) than that found in the other 2 groups (BIS [38.1%], OBS [37.3%]).

Virus detection

PCR detection of virus was as follows: RV (A, B, C) = 29; RSV A = 3, RSV B = 6, total = 9; CoV 229E/NL63 = 7; CoV OC43 = 2, total = 9; PIV 1 = 0, PIV 2 = 3, PIV 3 = 4, PIV 4 = 1, total = 8; IFV A = 6, IFV B = 0; AdV = 2; enterovirus = 1; MPV = 0; BV = 0.

ACD

ACD increased significantly in each group ($P < 0.000$) from the beginning of the study (0 week) to the end of the study (12 weeks). However, when comparing ACD of each group at each time point, there were no significant differences (Fig. 3). Both treatment options had a statistically similar effect on ACD over the 12-week treatment period. The following are ACD values for each group at each time point: 1) BIS group (0 week = 14.5, 4 weeks = 23.7, 8 weeks = 23.5, 12 weeks = 23.8); 2) MONT group (0 week = 16.3, 4 weeks = 23.8, 8 weeks = 24.7, 12 weeks = 26.6).

### Table 1. Demographics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>BIS group (n = 21)</th>
<th>MONT group (n = 22)</th>
<th>OBS group (n = 51)</th>
<th>Total (n = 94)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (mon)</td>
<td>39.33 (20–63)</td>
<td>40.77 (16–69)</td>
<td>31.35* (9–69)</td>
<td>35.34 (9–69)</td>
</tr>
<tr>
<td>Male/female (% male)</td>
<td>6/15 (28.6)</td>
<td>9/13 (40.9)</td>
<td>18/33 (25.5)</td>
<td>33/61 (35.1)</td>
</tr>
<tr>
<td>Food allergy</td>
<td>8 (38.1)</td>
<td>7 (31.8)</td>
<td>13 (25.5)</td>
<td>28 (29.8)</td>
</tr>
<tr>
<td>Atopic dermatitis</td>
<td>9 (42.9)</td>
<td>7 (31.8)</td>
<td>8 (15.7)†</td>
<td>24 (25.5)</td>
</tr>
<tr>
<td>Body weight (grams)</td>
<td>3,128 (2,505–3,827)</td>
<td>3,041 (2,386–4,172)</td>
<td>3,028 (1,658–3,690)</td>
<td>3,053 (1,658–4,172)</td>
</tr>
<tr>
<td>Sibling, mean (number with at least 1 sibling, %)</td>
<td>1.11 (15, 75.0)</td>
<td>1.38 (18, 90)</td>
<td>1.16 (36, 80.0)</td>
<td>1.20 (61.9)</td>
</tr>
<tr>
<td>Nursery school</td>
<td>16 (76.19)</td>
<td>3 (13.64)</td>
<td>8 (15.68)</td>
<td>17 (18.09)</td>
</tr>
<tr>
<td>Pet</td>
<td>6 (28.57)</td>
<td>3 (13.64)</td>
<td>8 (15.68)</td>
<td>17 (18.09)</td>
</tr>
<tr>
<td>Family smoker</td>
<td>11 (57.89)</td>
<td>8 (40.0)</td>
<td>27 (60.0)</td>
<td>46 (48.94)</td>
</tr>
<tr>
<td>Parental asthma</td>
<td>10 (52.63)</td>
<td>9 (42.86)</td>
<td>26 (57.78)</td>
<td>45 (47.87)</td>
</tr>
</tbody>
</table>

*Age in OBS group significantly lower ($P < 0.05$); †% Atopic dermatitis significantly less ($P < 0.05$).

<table>
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<th>OBS group (n = 51)</th>
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</thead>
<tbody>
<tr>
<td>sEDN (ng/mL)</td>
<td>106.01 (53.5–201)</td>
<td>119.45 (54–201)</td>
<td>52.68 (14.7–201)</td>
<td>80.22 (14.7–201)</td>
</tr>
<tr>
<td>HDM-IgE</td>
<td>112.62 (0.27–302)</td>
<td>130.16 (0.1–501)</td>
<td>21.66 (&lt;0.1–249)</td>
<td>66.89 (0.09–501)</td>
</tr>
<tr>
<td>Detection of any virus</td>
<td>8 (38.1)</td>
<td>17 (77.3)†</td>
<td>19 (37.3)</td>
<td>44 (46.8)</td>
</tr>
</tbody>
</table>

Data are shown as number (%) or number (range). BIS, budesonide inhalation suspension; MONT, eligible and receiving montelukast; OBS, not eligible and receiving montelukast.

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Table 2. Laboratory data

<table>
<thead>
<tr>
<th>Characteristics</th>
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</tr>
</tbody>
</table>

Data are shown as number (%) or number (range). BIS, budesonide inhalation suspension; MONT, eligible and receiving montelukast; OBS, not eligible and receiving Montelukast; sEDN, serum eosinophil-derived neurotoxin; HDM-IgE, house dust mite-immunoglobulin E. *% of OBS group with virus was significantly greater ($P = 0.005$).
Changes in sEDN with treatment

There was no significant change in sEDN in the BIS group (Fig. 4A), but there was a significant decrease found in the MONT group ($P < 0.000$) (Fig. 4B). When the OBS group was divided into those with low EDN (< 53 ng/mL) (Fig. 4C) and those with high EDN (> 53 ng/mL) (Fig. 4D), only patients with high EDN exhibited a significant decrease due to MONT treatment ($P = 0.023$), regardless of HDM sensitization.

Virus detection

Forty-four (46.8%) study subjects were positive for any virus tested (Table 2). When each treatment group (BIS, MONT was subgrouped according to virus-positivity, measured for sEDN “Pre” and “Post”), then changes in sEDN ($\Delta$EDN) were compared between virus-positive and -negative subjects in the BIS and MONT groups, and $\Delta$EDN was significantly larger in the virus-negative MONT subgroup (Supplementary Fig. S1B) and mean $\Delta$EDN was also larger in the virus-negative BIS subgroup, but not statistically significant (Supplementary Fig. S1B). Changes in ACD were not different between virus-positive and -negative subgroups in the BIS (Supplementary Fig. S2A) and MONT (Supplementary Fig. S2B) groups.

Rescue medication usage

Another secondary outcome of this study was change in rescue medication usage with treatment. Comparisons were done 3 ways: 1) within each group (BIS and MONT), usage at run-in (~2 weeks) was compared to usage at the end of the treatment period (12 weeks); 2) each subject in each group was analyzed over the study period (~2 weeks to 12 weeks) to determine any change in usage; 3) usage in the BIS group was compared to usage in the MONT group at each time point (~2, 0, 4, 8 and 12 weeks). There was a significant decrease ($P < 0.000$) in rescue medication usage within each group (BIS and MONT) and in each subject in each group when analyzed over the study period (~2 to 12 weeks) ($P = 0.002$). However, there were no significant differences found when comparing usage in the BIS group to that in the MONT group at each time point (~2, 0, 4, 8 and 12 weeks).
DISCUSSION

This study clearly demonstrated that preschool children with beta2-agonist responsive recurrent wheeze and multiple-trigger wheeze, a high EDN level, and HDM sensitivity responded well to both montelukast and budesonide. As expected, montelukast treatment significantly reduced sEDN levels. Unexpectedly, BIS treatment did not. Both treatments also resulted in significant decreases in rescue medication usage over the treatment period. In terms of clinical outcomes, montelukast was as effective as budesonide in the biomarker-defined (i.e., mite sensitized, eosinophilic asthma) subpopulation of preschool asthmatics.

Asthma is a heterogeneous disease with many phenotypes. The recognition of a high EDN/HDM-sensitive phenotype that exhibits a favorable response to standard asthma treatments is important. Of the 100 patients screened for eligibility, nearly half (43%) were found to be of this phenotype, which is considerable. The elevated levels of EDN and other eosinophil degranulation products found in symptomatic asthma would suggest airway inflammation is characterized not only by an increase in the number of eosinophils, but also an increase in airway eosinophil degranulation. Therefore, it has been suggested that the secretory
activity of eosinophils — a combination of concentration of eosinophils and their propensity to release degranulation products — may be a key marker for disease activity and is more accurately assessed by eosinophil degranulation products such as EDN. EDN is released almost exclusively by eosinophils; therefore, any change in EDN levels would be a direct reflection of changes in eosinophilic inflammation. Significant changes in EDN levels between disease phases have been noted in previous studies. EDN may therefore be useful in predicting therapeutic outcomes and is a welcome addition to a clinician’s ability to effectively treat and monitor asthma. This was further demonstrated in our study as montelukast treatment significantly reduced elevated EDN levels in the MONT group, which has also been shown in other asthma studies and even in patients with post-RSV bronchiolitis recurrent wheeze. Among the subjects excluded from the main study group (OBS group), the subgroup with no HDM sensitivity but elevated EDN experienced a reduction in their EDN levels as well. Montelukast was the chosen treatment for the OBS group because it has been reported that ICS is less effective in non-atopic populations. Furthermore, from our clinical experience montelukast is preferred over ICS by both physicians and patients in Korea and Japan.

Treatment-related reduction in EDN levels may yield actual patient benefits. In our study, subjects receiving montelukast treatment over 12 weeks had not only a significant increase in the number of ACD compared to the beginning of the study, but also had a statistically similar number of ACD at each time point (4, 8 and 12 weeks) when compared to those receiving BIS treatment (BIS group). This is an interesting finding considering a recent Cochrane Review of 65 studies directly comparing the safety and efficacy of anti-leukotrienes and ICSs as monotherapy found ICS treatment to be superior in a number of respects including symptom-free days. However, it should be noted that no sample size calculation was done for our study. This point will be discussed further in the study limitations section. The BIS group also showed a significant increase in ACD over the 12-week treatment period.

In previous asthma studies and AD studies, sEDN levels have correlated well with symptom scores. It is known that eosinophils release EDN and other degranulation products when activated; in particular, peripheral blood eosinophils were found to release approximately 17%-28% of their total intracellular EDN. Furthermore, Sedgwick et al. demonstrated that intracellular EDN is also increased in asthmatic patients, while ECP is not. Montelukast is an LTRA, capable of binding with cysteine leukotriene (cysLT) receptors expressed on eosinophils and disrupting their autocrine release. CysLTs can induce eosinophil degranulation, which then leads to release of a number of mediators including EDN, ECP and cysLTs. This may explain the reduction in EDN levels in the MONT group and in patients in the OBS group with high EDN levels. Corticosteroid treatment may or may not eliminate cytokine-primed, EDN-increased and longer-surviving eosinophils resulting in decreased EDN levels. We observed no significant decrease in EDN levels after budesonide treatment in our study, which was contrary to our initial hypothesis that budesonide would also reduce EDN levels. This may be because budesonide, an ICS, acts locally, whereas montelukast acts systemically. Eosinophils develop principally in the bone marrow and are recruited from the circulation, where they migrate to inflammatory sites, such as the airways, and then participate in immunologic processes. Once activated, the eosinophil releases EDN and other granular proteins. Changes in granular protein levels signal changes in the turnover and activity of the eosinophil. Taken together, eosinophil production, recruitment and activation are a systemic process.

All patients in this study were also tested for virus infection as it is well known that respiratory viral infection is the major cause of asthma exacerbation in children.
detected virus in about half of the treated subjects and reduction in sEDN levels was significantly larger in the virus-negative subgroup of the MONT group, indicating that virus infection may have hindered the effect of montelukast (Supplementary Fig. S1). However, EDN levels still significantly decreased in virus-positive MONT subgroups (data not shown), which may have resulted in the same levels of improvement in ACD in the virus-positive and -negative subgroups (Supplementary Fig. S2).

Study limitations included: 1) the open-label design, which could have affected results through the placebo effect; 2) measurements for both secondary outcomes (i.e., sEDN level and rescue medication usage) were done only 2 times, once at the beginning of the study for baseline and a second time at the end of the study, so it may have been more beneficial to measure serially; 3) patient recruitment was through a few different healthcare centers, which may have introduced spectrum bias and affected study results; 4) no sample size calculation was done, so that the number of eligible patients for the study (n = 43) may have been too small to accurately assess the association between predictor (i.e., elevated sEDN and positive IgE for HDM) and outcome (significant improvement in ACD, sEDN and rescue medication following treatment).

Recent meta-analyses\(^28\) of Green et al.’s study\(^8\) and 2 others\(^29,30\) bolster the hypothesis that treatment-based eosinophil indices is more efficacious (i.e., reduces the frequency and severity of exacerbations) than treatment based on clinical symptoms and other traditional objective measures of lung function such as peak expiratory flow and spirometry. Tailoring patient treatment based on the direct and objective measurement of airway inflammation resulted in significantly less frequent and severe asthma exacerbations and, consequently a trend towards fewer asthma-related hospital admissions.\(^28\) Knowing both the inflammatory profile and the clinical categorization of the patient develops a clearer phenotype of the asthma patient and appears to contribute to improved asthma care.\(^27,31\) Using a more accurate biomarker for eosinophilic inflammation in the management of asthma may produce even better results.\(^32,33\) It has been suggested by many studies that EDN is a more accurate biomarker than eosinophil numbers or percentages.\(^11,12,22,32\) Future studies of EDN as a biomarker for the diagnosis, treatment and monitoring of conditions associated with airway inflammation — such as recurrent wheeze, multiple-trigger wheeze and asthma — are needed.

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SUPPLEMENTARY MATERIALS

Supplementary Fig. S1
Decrease in EDN levels (ΔEDN) from visit 1 (enrollment) to visit 4 (after 12 weeks of treatment) in BIS group (A) and MONT group (B).

Click here to view
Supplementary Fig. S2
Changes in ACD per week in virus-positive (virus+) and virus-negative (virus−) subgroups in BIS group (A) and MONT group (B).

Click here to view

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https://e-aair.org


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