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Effects of 24-week add-on treatment with ciclesonide and montelukast on small airways inflammation in asthma

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Trial registration; Registry ID UMIN000001083

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Authors’ contributions

HN recruited and managed the patients, collected, analyzed and interpreted the data, and prepared the manuscript. GP collected, analyzed and interpreted the data, and wrote the draft. HM conceived the study, recruited and managed the patients, collected, analyzed and interpreted the data, and revised the manuscript. TI collected and analyzed data and prepared the part of the draft. AN and II recruited the patients, collected the data, and contributed to the edition of the manuscript. TO and HI performed IOS measurements and collected the data and prepared the part of the draft. TT, TN, and YK measured exhaled nitric oxide and analyzed and interpreted the data, and prepared the part of the draft. MM contributed to the discussion of the data and critical revision of the manuscript.
Abstract

Background: Eosinophilic inflammation of the small airways is a key process in asthma that often smolders in treated patients. The long-term effects of add-on therapy on the persistent inflammation in the small airways remain unknown.

Objective: To examine the effects of add-on therapy with either ciclesonide, an inhaled corticosteroid with extrafine particles, or montelukast on small airway inflammation.

Methods: Sixty patients with stable asthma receiving inhaled corticosteroid treatment were enrolled in a randomized, open-label, parallel comparison study of 24-week add-on treatment with ciclesonide or montelukast. Patients were randomly assigned to 3 groups: ciclesonide (n = 19), montelukast (n = 22) and no add-on as controls (n = 19). At baseline and at weeks 4, 12 and 24, extended nitric oxide analysis; pulmonary function tests, including impulse oscillometry; blood eosinophil counts; and asthma control tests (ACTs) were performed.

Results: A total of 18 patients in the ciclesonide group, 19 in the montelukast group and 15 in the control group completed the study and were analysed. With repeated-measures analysis of variance, ciclesonide produced a significant decrease in alveolar nitric oxide and a significant improvement in ACT scores over time. Montelukast produced significant decreases in alveolar nitric oxide concentrations and blood eosinophil counts over time and slightly improved ACT scores, whereas no such changes were observed in the control group. Alveolar nitric oxide concentrations with ciclesonide and reactance area at low frequencies with montelukast produced greater improvements over time compared with control.

Conclusions: Ciclesonide add-on therapy and montelukast add-on therapy may act differently, but both separately can improve small airway abnormalities and provide better asthma control.
58 Funding; funded by none

59

60 **Key words:** add-on treatment, alveolar nitric oxide, asthma control, ciclesonide, montelukast, small airways
**Introduction**

Asthma is a chronic inflammatory disease of the airways characterized by variable, recurring symptoms and reversible airflow obstruction. The immunohistopathologic features include infiltration of eosinophils and lymphocytes, mast cell activation and epithelial cell injury. To date, pathological, physiologic and radiologic findings have provided sufficient evidence to support not only large but also small airways involvement in inflammation and airflow obstruction, particularly in patients with severe asthma.

Recently, it was found that eosinophilic inflammation of the small airways could be assessed by determining alveolar nitric oxide concentrations. Small airway inflammation as assessed by alveolar nitric oxide concentrations is increased in patients with refractory asthma and those with nocturnal asthma and is associated with disease severity and small airways dysfunction. Of note, 20% of asthmatic patients have increased alveolar nitric oxide concentrations despite treatments with inhaled corticosteroids (ICSs) and long-acting β2 agonists. Alveolar nitric oxide concentrations can also predict a future risk of disease exacerbation.

These findings suggest that, even in apparently stable patients taking ICSs, additional treatment targeting the small airways may lead to reaching total asthma control.

Few studies have evaluated the changes in alveolar nitric oxide concentrations based on either an uncorrected or corrected model of add-on medication for persistent inflammation of the small airways. Previous studies found that oral prednisolone, but not double doses of ICSs, could decrease alveolar nitric oxide concentrations. These results suggest that alveolar nitric oxide concentrations may be resistant to a simple ICS dose elevation. In steroid-naïve patients, however, extrafine particle hydrofluoroalkane–ciclesonide resulted in decreased alveolar nitric oxide concentrations and hydrofluoroalkane–beclomethasone propionate...
improved peripheral airway dysfunction. Collectively, an extrafine particle ICS is expected to decrease alveolar nitric oxide concentrations when they are used as an add-on medication. Leukotriene receptor antagonists (LTRAs) that are administered systemically are another medication that are supposed to decrease alveolar nitric oxide concentrations. Treatment with montelukast for 4 weeks improved small airway obstruction in steroid-naive patients, which resulted in a decrease in regional air trapping. So far published study data of an add-on LTRA to ICS therapy for 3 to 8 weeks with regard to alveolar nitric oxide concentrations have been conflicting. These effects require confirmation with a longer-term study.

For this study, we hypothesized that prolonged add-on therapy to the ICS treatment with either ciclesonide or montelukast would have beneficial effects on the persistent inflammation of the small airways and would improve pulmonary function. To test this hypothesis, our primary objectives were to examine the effects of this add-on therapy on alveolar nitric oxide concentrations and to compare its effects on small airways in patients with stable asthma who had not been previously treated with extrafine particle ICSs or LTRAs.
Methods

The full details of the study methods are given in the eMethods. In brief, adult patients with stable asthma who regularly visited our outpatient asthma clinic were enrolled from April 2008 to August 2011. Asthma was diagnosed according to American Thoracic Society criteria \(^{20}\). Patients were included if they were classified as being in treatment steps 2 to 5 of ICS treatment according to the Global Initiative for Asthma guidelines \(^{21}\). These patients had no exacerbations 3 months before enrollment, had alveolar nitric oxide concentrations of 5.0 ppb or higher, and were either never-smokers or ex-smokers who had smoked fewer than 5 pack-years and had stopped more than 1 year before. The threshold level for uncorrected alveolar nitric oxide concentrations was set at 5.0 ppb; this value was the average minus 1 SD of uncorrected alveolar nitric oxide concentrations of 70 patients with asthma taking ICSs in our previous study \(^{22}\).

Exclusion criteria were current or previous use of extrafine particle ICSs or LTRAs. Patients were also excluded if, during the study period, any adverse effects of the add-on therapy or asthma exacerbations, including mild exacerbations, defined as an increased need for rescue use of short-acting \(\beta_2\)-agonists, were noted.

This study was approved by the ethics committees of our institute and was registered in UMIN Clinical Trials Registry (Registry Identified UMIN000001083). Written informed consent was obtained from all participants.

Design and Measurements

This was a randomized, open-label, parallel comparison study of 24-week add-on treatment with either inhaled ciclesonide or montelukast. Patients were randomly assigned to 3 treatment groups: inhaled ciclesonide, 400 \(\mu\)g once daily add-on (ciclesonide group);
montelukast, 10 mg once daily add-on (montelukast group); and control group, who were taking current medication only. At weeks 0 (baseline), 4, 12, and 24 (end of study period) the patients underwent extended nitric oxide analysis and pulmonary function tests, including tests with an impulse oscillometry system (IOS), spirometry, and a nitrogen single-breath wash out test. At the same time points, patients completed an asthma control test (ACT) questionnaire comprising 5 questions with a best possible score of 25 and were given a rhinitis symptom score (RSS), a self-assessment questionnaire comprising 4 questions, the responses to which were ranked on a Likert-type scale with a maximum of 5 points per answer. The RSS was determined based on the Japanese Guideline for Allergic Rhinitis (best score, 20) (eTable 1).

At the start and end of the study period, blood samples were obtained for blood eosinophil counts and serum high sensitivity C-reactive protein, serum eosinophil cationic protein, and serum YKL-40, a chitinase like protein. Blood samples for eosinophil cationic protein determinations were collected in SST tubes (Becton Dickinson, Mountain View, California) and were processed as previously described. YKL-40 levels were determined using an enzyme-linked immunosorbent assay kit (Quidel, San Diego, California) following the manufacturer’s instructions.

Nitric oxide levels were determined with a chemiluminescence analyzer (NOA 280; Sievers, Boulder, Colorado) according to current guidelines, and as previously described alveolar nitric oxide concentrations are provided as noncorrected and corrected values using a trumpet-shaped model with axial back diffusion (eMethods).

After nitric oxide measurements, patients underwent prebronchodilator and postbronchodilator (ie, inhalation of 200 μg of salbutamol) pulmonary function tests.
Spirograms were obtained as recommended by the American Thoracic Society/European Respiratory Society. A nitrogen single-breath washout test was performed only before the inhalation of salbutamol to assess ventilation inhomogeneity by measuring the slope of phase 3 of the nitrogen washout curve ($\Delta N_2$).

Respiratory impedance was determined by IOS using a Jaeger MasterScreen, IOS™ (Erich Jaeger, Hoechberg Germany) that met standard recommendations (eMethods). 16, 22

**Statistical analysis**

For sample size determinations, we originally sought to enroll 90 patients based on previous findings 15, 17, 19. However, as described in the “Results” section, we decided to stop patient enrollment at 60 because of the more frequent occurrence of exacerbations in the control group, although these were mild.

Statistical analysis used JMP 6.00 (SAS Institute Inc., Cary, North Carolina) on a per-protocol basis. For non-normally distributed results, comparisons were made by the Kruskal–Wallis test, Fisher exact test or Wilcoxon signed-rank test as appropriate. For normally distributed results, comparisons were made by analysis of variance (ANOVA) and the paired t-test. Two-way repeated-measures ANOVA was used to assess the variations among the 3 treatment modalities and at different time points. For cases with unequal variations in the treatment modalities, only 1-way repeated-measures ANOVA within 1 treatment group was used. For correlation analysis, the Spearman rank-correlation test was used. Data are expressed as mean ± SD. $P \leq 0.05$ were considered statistically significant.
Results

Enrollment, Dropout, and Exacerbation Rates and Baseline Characteristics

Sixty asthmatic patients were enrolled in this study and randomly assigned to the groups: 19 in the ciclesonide group, 22 in the montelukast group, and 19 in the control group (Fig 1). The reasons for patient dropout were as follows: in the ciclesonide group, 1 patient had a possible adverse effect (urticaria); in the montelukast group, 3 patients had possible adverse effects (2 experienced mild gastrointestinal discomfort and they preferred to discontinue use of the medication and 1 patient had mildly elevated transaminase levels); and in the control group, 3 had mild asthma exacerbations and they preferred to intensify medications and 1 patient discontinued ICS treatment following a general practitioner’s advice. As a result 18 patients in the ciclesonide group, 19 in the montelukast group, and 15 in the control group completed the study and were analyzed thereafter (Table 1). For these patients, adherence to the add-on and current medications was satisfactory, which was confirmed by 2 of the authors (H.N. and H.M.) on each visit by checking the residual number of medications.

When the exacerbation frequencies were compared between the 19 patients in the control group and the 41 patients in the add-on therapy groups and assuming that the 5 patients who dropped out for reasons other than exacerbation would complete the protocol without exacerbation, the control group had a significantly higher rate of exacerbation (p = 0.03; by Fisher exact test). The baseline patient characteristics, ICS doses, and biomarkers, including fractional exhaled nitric oxide (FeNO) and alveolar nitric oxide concentrations, were not significantly different among the 3 patients who later experienced mild exacerbations and the other 57 patients.
**ACT scores and RSSs**

By 1-way ANOVA, there was a significant improvement in ACT scores during the treatment period within the ciclesonide group \((p = 0.02; \text{ Fig 2})\), and there was a trend for improvement within the montelukast group \((p = 0.08)\). When subscores for the ACT components were separately analyzed in the ciclesonide group, subscores for ACT question 3 concerning nocturnal symptoms and question 5 for self-rating were marginally and insignificantly improved over time \((p = 0.05 \text{ and } p = 0.06, \text{ respectively})\). Because of the unequal variations among the 3 treatment modalities, we did not conduct 2-way ANOVA for the ACT scores. Details on ACT scores across the treatment steps are presented in eTable 2.

Among the 3 groups, neither the proportions of patients with allergic rhinitis nor their baseline RSSs differed. However, a significant difference was seen in the time trends for RSS among the 3 treatment modalities \((p = 0.004; \text{ eFig 1})\); in particular, using 2-way ANOVA, significant differences were seen for the symptom of nasal obstruction \((p = 0.046)\). When comparing 2 different treatment modalities in a post hoc analysis, the montelukast group exhibited a significantly better time trend for the RSS than the control group \((p < 0.001)\) and a trend for better scores than the ciclesonide group \((p = 0.07; \text{ eFig 1})\). A significant increase in RSS over time was found only in the montelukast group \((p < 0.001, \text{ by 1-way ANOVA})\).

There were no associations between changes in ACT or RSS from baseline to the end of the treatment period and changes in alveolar nitric oxide concentrations or corrected alveolar nitric oxide concentrations in either treatment group.

**Nitric Oxide Results**
No significant differences were found in the time trends for FeNO at an expiratory flow rate of 50 mL/s among the 3 treatment modalities or within each of the groups (results not shown).

The time trends for uncorrected alveolar nitric oxide concentrations were significantly different among the 3 treatment groups (p = 0.048, by 2-way ANOVA). When comparing 2 different treatment modalities in a post hoc analysis, the ciclesonide group had a greater decrease in alveolar nitric oxide concentrations over time than the control group (p = 0.03, by 2-way ANOVA). By 1-way ANOVA, alveolar nitric oxide concentrations in the control group did not change during the study period, whereas in both of the add-on treatment groups, alveolar nitric oxide concentrations significantly decreased over time (p = 0.01 for the ciclesonide and montelukast groups; Fig 3).

For corrected alveolar nitric oxide concentrations, 1-way ANOVA showed that there was an insignificant decrease over time in the ciclesonide group (p = 0.06).

**Pulmonary Function Tests**

None of the spirometry indices, ΔN₂, or IOS indices of respiratory resistance at 5 Hz (Rrs₅), respiratory resistance at 20 Hz (Rrs₂₀), or respiratory reactance at 5 Hz (Xrs₅) revealed any difference among the 3 treatment modalities during the treatment period regardless of prebronchodilator or postbronchodilator conditions. No significant changes were observed within any of the 3 groups (data not shown).

A significant difference was found in the time trends for the reactance area (AX) among the 3 treatment modalities (p = 0.04, by 2-way ANOVA). The AX levels in the montelukast group improved over time when compared with the control group (p = 0.05, by 2-way ANOVA; Fig 4). For Rrs₅–Rrs₂₀, 2-way ANOVA was not used because of the unequal variations among
the 3 treatment modalities; however, 1-way ANOVA revealed that there was a trend for a change
over time in the ciclesonide group (p = 0.09).

Although there were associations between corrected alveolar nitric oxide concentrations
and IOS indices of AX or Rrs$_5$-Rrs$_{20}$ at baseline (r = 0.30, p < 0.05 for both, n = 52), no
associations were found between changes in pulmonary function data from baseline to the end of
the treatment period and changes in alveolar nitric oxide concentrations or corrected alveolar
nitric oxide concentrations in either treatment group.

**Blood Test Results**

Blood samples were obtained at baseline and at the end of the treatment period to determine
blood eosinophil counts and serum levels of eosinophil cationic protein, high sensitivity C-
reactive protein, and YKL-40. No significant changes were found in these tests results between
the beginning and the end of the treatment period, except for the montelukast group in which the
eosinophil counts significantly declined after treatment (2.9 ± 2.2% at 24 weeks)(p = 0.02, paired
$t$ test).
To the best of our knowledge, this is the first long-term study that clarified the benefits and potential role of add-on therapy with either ciclesonide of extrafine particle ICS or montelukast in steroid-treated patients with stable asthma. Ciclesonide may have attenuated smoldering inflammation of the small airways and significantly improved asthma control over time. Montelukast ameliorated the remnant dysfunction of the small airways and reduced nasal symptoms and blood eosinophil counts. To a lesser extent than ciclesonide, montelukast also improved smoldering inflammation of the small airways.

Alveolar nitric oxide concentration is an established marker of small airway inflammation and is correlated with eosinophil counts in bronchoalveolar lavage fluid. In the ciclesonide group, alveolar nitric oxide concentrations significantly decreased over time when compared with the control group and the ciclesonide intragroup analysis. Our data confirmed earlier findings of the effects of 5-week treatment with ciclesonide on alveolar nitric oxide concentrations in steroid-naïve patients and reinforced the advantage of extrafine particle ICSs to treat smoldering inflammation of the small airways, even in patients already taking ICSs. There remains the possibility that the addition of ciclesonide to the patients’ current medication may have exerted anti-inflammatory effects via the increase in the total amount of ICS, which may have suppressed the remnant inflammation throughout the airways. However, this is unlikely because FeNO at 50 mL/s did not change over time. Taken the results of the previous short-term study and current study together, ciclesonide would be capable of treating the small airways potentially because of its particles size, which was sufficiently small to reach the peripheral airways.
In contrast to uncorrected alveolar nitric oxide concentrations, corrected alveolar nitric oxide concentrations only showed a trend toward being decreased in the ciclesonide group (p = 0.06, 1-way ANOVA). Although corrected alveolar nitric oxide concentrations reflect airway dysfunction \(^{22,29}\), as do alveolar nitric oxide concentrations, corrected alveolar nitric oxide concentrations do not reflect disease severity \(^{14,22}\) or asthma control status \(^{29}\). It is also not increased during asthma exacerbations in adults \(^{30}\), a finding that is in contrast to several lines of evidence for alveolar nitric oxide concentrations. Although alveolar nitric oxide concentrations are contaminated with bronchial nitric oxide, potentially from small conducting airways where diffusion begins to replace bulk flow, our findings on alveolar nitric oxide concentrations imply that relatively small airways, albeit not actual peripheral airways, are still important in the management of asthma.

Studies of add-on medication using LTRAs that have evaluated changes in alveolar nitric oxide concentrations in persistent inflammation of the small airways reported inconsistent findings. Previous add-on studies of montelukast to fluticasone \(^{18}\) or fluticasone and salmeterol treatment \(^{12}\) did not find any significant benefits for montelukast with regard to decreases in alveolar nitric oxide concentrations after montelukast add-on therapy. However, these earlier studies were relatively short-term, with treatment periods of only 3 to 4 weeks. Yasui et al. investigated pranlukast use in patients with stable asthma and found significant decreases in both corrected and uncorrected alveolar nitric oxide concentrations after 8-week crossover of add-on therapy with pranlukast \(^{19}\). In agreement with that study, we found that alveolar nitric oxide concentrations in the montelukast group decreased during the 24-week add-on period, although these levels were not significantly different from the control group. As with the ciclesonide group, FeNO at 50 mL/s did not change over time. These findings indicate that add-on treatment
with LTRAs for longer than 8 weeks suppresses the remnant inflammation in the small airways.

In addition, our intervention study that covered the 2 seasons for allergic rhinitis (spring and autumn) provided additional evidence of the established benefit of montelukast on allergic rhinitis and justified a role for LTRA in the therapy for patients with stable asthma with concomitant allergic rhinitis, even those with minimal symptoms.

Symptoms and airway obstruction are integral to the definition of asthma, and represent important components for assessing asthma control in both clinical practice and clinical trials. Therefore, one of the end points in our study was ACT scores. Despite the disadvantage in adherence to inhalation use, as reported previously on adherence to LTRA of 67.7% vs. 33.8% for ICS, ACT scores significantly improved over time in the ciclesonide group. In addition, there was a marginal improvement in the subscore of ACT question 3 concerning nocturnal symptoms in the ciclesonide group. To date, a number of studies have confirmed that eosinophilic inflammation worsens in patients with nocturnal asthma, particularly in the peripheral airways. Lehtimaki et al reported that nocturnal symptoms in asthmatic patients were related to higher alveolar nitric oxide concentrations. These results are in accordance with our results showing that ciclesonide add-on treatment reduced inflammation in the small airways, as assessed by alveolar nitric oxide concentrations, and improved nocturnal symptoms, as assessed by ACT subscores. Care must be taken when interpreting these findings, however, because the minimally important difference in ACT scores that reflects a clinically meaningful change is considered to be 3 points, and the increase in ACT composite scores in our ciclesonide group did not achieve this. Despite this minimal change, these statistically significant changes would still favour add-on therapy for patients with seemingly stable asthma.
We did not find any significant changes in spirometry function results or ΔN₂ between the control and therapy add-on groups, which may seem unexpected. Spirometry is not sensitive enough to detect early small airway involvement because the small airways are pathways of very low resistance and only contribute to approximately 10% of the total airway resistance. Instead of using ΔN₂, ventilation heterogeneity within conductive and acinar airways could have been separately assessed using a nitrogen multiple-washout test. Another possible reason could be that our patients had already good pulmonary function, so that changes in alveolar nitric oxide concentrations were not reflected in the airway function. However, in the montelukast group, the AX significantly decreased over time when compared with the control group, as was found in our previous intervention study in steroid-naïve patients. Montelukast may have reversed remodeling in the airway walls by reducing airway smooth muscle layer thickening and subepithelial fibrosis in long-term treatment, as has been shown in an animal model. More significant findings might be expected in extended studies in a larger number of patients.

A limitation of our study was that it was a parallel, open-label, and unblinded study, which might have influenced subjective measures, such as asthma symptoms and rescue use of short-acting β₂ agonists. Another issue is the use of 2 different inhalers for corticosteroids, although we achieved good adherence in the ciclesonide group. In future studies with more patients and longer treatment periods, this issue could be resolved.

In addition, we may have missed some patients with occult inflammation in the small airways by excluding those with alveolar nitric oxide concentrations less than 5 ppb, given that some patients who have high FeNO and low alveolar nitric oxide concentrations exhibit paradoxical increases in alveolar nitric oxide concentrations after treatment, possibly because of dilatation of constricted small airways from terminal to respiratory bronchioles. However, by
setting this threshold for alveolar nitric oxide concentrations during patient enrollment, the
changes of alveolar nitric oxide concentrations in this study could be simply interpreted.

Finally, from the ethical standpoint, we stopped enrollment at 60 patients because of a
higher, albeit mild, exacerbation rate in the control group, which was consistent with the finding
that elevated alveolar nitric oxide concentration was associated with risk of asthma
exacerbation. Thus, some of the insignificant findings, particularly of the pulmonary function
data in this study, may be due to lesser statistical power. Lack of associations between the
changes in alveolar nitric oxide concentrations and changes in pulmonary function data or ACT
scores might be another issue. However, we did not set the sample size to seek significant
associations between changes in alveolar nitric oxide concentrations and any other clinical
indices because of their potentially large variations during the treatment period, although
alveolar nitric oxide concentrations, pulmonary function, and ACT were intuitively thought to
behave in parallel. Despite these limitations, the current findings of a decrease in alveolar nitric
ox ide concentrations with add-on treatment are sufficient to be used as a future reference when
intensifying treatment with extrafine particle ICS or LTRA add-on therapy, even in patients with
seemingly stable asthma who are receiving ICS treatment but still have evidence of small
airways inflammation as assessed by alveolar nitric oxide concentrations.

We conclude that ciclesonide and montelukast may act differently but that both
separately can improve small airway abnormalities (eTable 3). By coadministration of these
medications, cumulative effects on inflammation and small airways function can be expected and
should be clarified in a future study. We can achieve additional benefits by treating inflammation
of the small airways in patients with stable asthma to reach the ultimate asthma treatment goal:
ideal control.
Acknowledgments

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20. Standards for the diagnosis and care of patients with chronic obstructive pulmonary disease (COPD) and asthma. This official statement of the American Thoracic Society was adopted by the ATS Board of Directors, November 1986. Am Rev Respir Dis. 1987;136: 225-244.


Figure legends

Figure 1. Registration and randomization

Figure 2. Asthma control test (ACT) scores in the 3 study groups. *Significant changes in ACT scores within the ciclesonide add-on group (p = 0.02, by 1-way analysis of variance).

Figure 3. Alveolar nitric oxide concentrations in the 3 study groups. *Significant difference in the time trends for alveolar nitric oxide concentrations among the 3 treatment modalities (p = 0.048, by 2-way analysis of variance [ANOVA]). †Significant changes in alveolar nitric oxide concentrations in the ciclesonide add-on group (p = 0.03 vs the control group, by 2-way ANOVA) (p = 0.01, by 1-way ANOVA). ‡Significant changes within montelukast add-on group (p = 0.01, by 1-way ANOVA).

Figure 4. Reactance area (AX) levels in the 3 study groups. *Significant difference in the time trends for AX levels among the 3 treatment modalities (p = 0.04, by 2-way analysis of variance [ANOVA]), †posthoc analysis between the montelukast add-on and control groups (p = 0.05, by 2-way ANOVA).
Figure 1

60 Entry

Pre-intervention
- +Ciclesonide 19
  - 4 wks
    - +Ciclesonide 19
    - 12 wks
      - +Ciclesonide 18
      - 24 wks
        - +Ciclesonide 18
  - +Montelukast 22
    - 4 wks
      - +Montelukast 22
      - 12 wks
        - +Montelukast 20
        - 24 wks
          - +Montelukast 19
  - Control (no add-on) 19
    - 4 wks
      - Control 18
      - 12 wks
        - Control 17
        - 24 wks
          - Control 15

Randomization

1 drop out due to urticaria
2 drop out due to mild gastro-intestinal discomfort
1 drop out due to elevation of transaminase
1 drop out due to protocol violation
2 drop out due to mild exacerbation of asthma
Figure 2

ACT scores

Montelukast
Ciclesonide
Control

*
Figure 3

Montelukast
Ciclesonide
Control

CANO (ppb)

*
†
‡

0 4 8 12 24 weeks
Figure 4

AX (kPa L⁻¹)

*  †

Montelukast

Ciclesonide

Control

weeks 0 4 12 24

AX (kPa L⁻¹)

.4 .6 .8 1.0 1.2 1.4 1.6
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<th>Control group (n = 15)</th>
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</tr>
<tr>
<td>Rrs5, kPa sL⁻¹</td>
<td>0.43 ± 0.15</td>
<td>0.40 ± 0.13</td>
<td>0.40 ± 0.14</td>
</tr>
<tr>
<td>Rrs20, kPa sL⁻¹</td>
<td>0.35 ± 0.11</td>
<td>0.31 ± 0.09</td>
<td>0.33 ± 0.10</td>
</tr>
<tr>
<td>Rrs5-Rs20, kPa sL⁻¹</td>
<td>0.08 ± 0.05</td>
<td>0.09 ± 0.07</td>
<td>0.07 ± 0.07</td>
</tr>
<tr>
<td>Xrs5, kPa sL⁻¹</td>
<td>-0.14 ± 0.06</td>
<td>-0.14 ± 0.06</td>
<td>-0.14 ± 0.07</td>
</tr>
<tr>
<td>AX, kPa L⁻¹</td>
<td>0.67 ± 0.51</td>
<td>0.78 ± 0.91</td>
<td>0.71 ± 0.68</td>
</tr>
<tr>
<td>ACT score</td>
<td>22.7 ± 2.5</td>
<td>23.2 ± 2.3</td>
<td>23.2 ± 2.7</td>
</tr>
<tr>
<td>Rhinitis symptom score</td>
<td>16.9 ± 2.1</td>
<td>16.4 ± 2.1</td>
<td>17.2 ± 1.7</td>
</tr>
<tr>
<td>Blood eosinophils, %</td>
<td>5.3 ± 3.9</td>
<td>4.7 ± 2.9</td>
<td>4.0 ± 2.5</td>
</tr>
<tr>
<td>Serum ECP, μg/L</td>
<td>16.6 ± 17.6</td>
<td>11.4 ± 11.1</td>
<td>15.0 ± 15.5</td>
</tr>
<tr>
<td>Serum hsCRP, mg/dL</td>
<td>0.21 ± 0.37</td>
<td>0.10 ± 0.20</td>
<td>0.14 ± 0.17</td>
</tr>
<tr>
<td>Serum YKL-40, ng/dL</td>
<td>115.2 ± 86.0</td>
<td>123.2 ± 83.7</td>
<td>93.2 ± 116.4</td>
</tr>
</tbody>
</table>

Data are presented as number or mean ± SD, except for IgE, which is presented as median (range); p > 0.05 for all characteristics according to the analysis of variance, the Kruskal-Wallis test or Fisher’s exact test. 1) According to the 2006 Global Initiative for Asthma guidelines, 2) Atopy was determined based on the presence of specific serum IgE antibodies to at least 1 common inhalant allergen, including cat dander, dog dander, weed pollens, grass pollens, molds, or house dust mite, 3) equivalent to fluticasone propionate
E-Supplement material

Effects of 24-week add-on treatment with ciclesonide and montelukast on small airways inflammation in asthma

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* HN and GP equally contributed to this study

Trial registration: Registry ID UMIN000001083

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Methods

Adult patients with stable asthma who regularly visited our outpatient asthma clinic were enrolled from April 2008 to August 2011. Asthma was diagnosed according to American Thoracic Society criteria based on a history of recurrent episodes of wheezing and chest tightness, with or without cough, and documented airway reversibility with a bronchodilator or hyperresponsiveness to inhaled methacholine e1.

Nitric oxide levels were determined with a chemiluminescence analyzer (NOA 280; Sievers, Boulder, Colorado) according to current guidelines and as previously described e2. The analyzer was daily calibrated with gas without nitric oxide and a standard concentration of 640 ppb nitric oxide. Lower detection limit for nitric oxide was 2 ppb. The concentrations were determined using a data analysis program (NOA Analysis™ Software; Sievers). Seated patients inserted a mouthpiece, inhaled orally to total lung capacity, exhaled immediately against a resistance and maintained mouth pressure at 20 cm H2O, displayed on a pressure gauge. The steady-state nitric oxide plateau was taken as the fractional exhaled nitric oxide (FeNO) value.

By varying expiratory resistances, we measured FeNO levels at 3 expiratory flows of 50, 100 and 200 mL/s in that order. Alveolar nitric oxide concentrations are provided as non-corrected e3 and corrected values using trumpet-shaped model and axial back diffusion e2, e4.

After nitric oxide measurements, patients underwent prebronchodilator and postbronchodilator (ie, inhalation of 200 μg of salbutamol) pulmonary function tests. Respiratory impedance was determined by impulse oscillometry system (IOS) followed by spirometric test and a nitrogen single-breath washout test. Forced vital capacity, forced expiratory volume in 1 second, and forced midexpiratory flow were determined using a ChestGraph HI-701 spirometer (Chest MI Corp., Tokyo, Japan). Spirograms were obtained in triplicate, and the best of 3 reproducible measurements was recorded, as recommended by the American Thoracic Society/European Respiratory Society e5. A nitrogen single-breath washout test was performed only before the inhalation of salbutamol to assess ventilation inhomogeneity by measuring the slope of phase 3 of the nitrogen washout curve.
Respiratory impedance was determined using a Jaeger MasterScreen, IOS (Erich Jaeger, Hoechberg Germany), which met standard recommendations \(^{6}\). In brief, rectangular mechanical impulses containing a continuous power spectrum ranging from 0 to 100 Hz, generated by a loudspeaker at intervals of 0.2 second, were applied to the respiratory system through a mouthpiece during tidal breathing. The resulting pressure and flow signals were measured next to the mouthpiece and were analyzed for amplitude and phase differences using a fast Fourier transform to determine respiratory resistance (Rrs) and respiratory reactance (Xrs) of the total respiratory system. To reduce loss of energy in the upper airways, the chin and cheeks were supported by the patients’ hands. As proxies for peripheral airway function, we used the negative frequency dependence of Rrs between 5 and 20 Hz (Rrs5-Rrs20), Xrs at 5 Hz (Xrs5), and reactance area (AX) that is the integral of Xrs from 5 Hz to the resonant frequency at which Xrs crosses zero \(^{2,7}\).
References

e1. Standards for the diagnosis and care of patients with chronic obstructive pulmonary disease (COPD) and asthma. This official statement of the American Thoracic Society was adopted by the ATS Board of Directors, November 1986. Am Rev Respir Dis. 1987;136:225-244.


eFigure legends

eFigure 1. Rhinitis symptom scores (RSS) in the 3 study groups. *Significant difference in the time trends for RSS among the 3 treatment modalities (p = 0.004, by 2-way analysis of variance [ANOVA]).
†Significant changes in RSS in montelukast add-on group (p < 0.001 vs control group, by 2-way ANOVA) (p < 0.001, by 1-way ANOVA).
eFigure 1

RS scores

Montelukast
Ciclesonide
Control

*  
†


<table>
<thead>
<tr>
<th>Rhinitis symptom score</th>
<th>Scoring</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. Mean number of episodes of paroxysmal sneezing in a day</td>
<td></td>
</tr>
<tr>
<td>1. ≥ 21 times</td>
<td>2. 20-11 times</td>
</tr>
<tr>
<td>B. Mean number of episodes of nasal discharge a day</td>
<td></td>
</tr>
<tr>
<td>1. ≥ 21 times</td>
<td>2. 20-11 times</td>
</tr>
<tr>
<td>C. Nasal blockage</td>
<td></td>
</tr>
<tr>
<td>1. completely obstructed all day</td>
<td></td>
</tr>
<tr>
<td>2. severe nasal blockage causing prolonged oral breathing in a day</td>
<td></td>
</tr>
<tr>
<td>3. severe nasal blockage causing occasional oral breathing in a day</td>
<td></td>
</tr>
<tr>
<td>4. nasal blockage without oral breathing</td>
<td></td>
</tr>
<tr>
<td>5. not obstructed / no symptoms</td>
<td></td>
</tr>
<tr>
<td>D. Disturbance of daily activity (troubles with work, study, household work, sleep, going out, etc)</td>
<td></td>
</tr>
<tr>
<td>1. impossible</td>
<td></td>
</tr>
<tr>
<td>2. painful and complicating daily life</td>
<td></td>
</tr>
<tr>
<td>3. intermediate between 2) and 4)</td>
<td></td>
</tr>
<tr>
<td>4. few troubles</td>
<td></td>
</tr>
<tr>
<td>5. not disturbed at all</td>
<td></td>
</tr>
</tbody>
</table>
**Table 2.** Asthma control test (ACT) scores and distribution of control status at baseline according to the treatment steps

<table>
<thead>
<tr>
<th>Treatment steps</th>
<th>2 and 3</th>
<th>4 and 5</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ciclesonide (n = 18)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ACT scores</td>
<td>23.1 ± 1.9</td>
<td>16</td>
<td>NS</td>
</tr>
<tr>
<td>total/good/no control (n)</td>
<td>7/10/0</td>
<td>0/0/1</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Montelukast (n = 19)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ACT scores</td>
<td>23.3 ± 2.4</td>
<td>22.3 ± 2.3</td>
<td>NS</td>
</tr>
<tr>
<td>total/good/no control (n)</td>
<td>7/7/2</td>
<td>1/2/0</td>
<td>NS</td>
</tr>
<tr>
<td>Control (n = 15)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ACT scores</td>
<td>23.7 ± 1.7</td>
<td>21.3 ± 5.5</td>
<td>NS</td>
</tr>
<tr>
<td>total/good/no control (n)</td>
<td>5/7/0</td>
<td>1/1/1</td>
<td>NS</td>
</tr>
</tbody>
</table>

Data are presented as mean ± SD.

Control status is defined as total when ACT score was equal to 25 points, good when ACT score was 20 or higher, no control when ACT score was less than 20.

NS; no significant difference by Wilcoxon rank-sum test or χ² test.
**eTable 3. Summary of the results**

<table>
<thead>
<tr>
<th></th>
<th>Ciclesonide add-on</th>
<th>Montelukast add-on</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>FeNO</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>Alveolar nitric oxide concentrations vs other groups</td>
<td>Significant decrease vs controls</td>
<td>NS</td>
<td>-</td>
</tr>
<tr>
<td>within the treatment modality</td>
<td>Decreased</td>
<td>Decreased</td>
<td>NS</td>
</tr>
<tr>
<td>Corrected alveolar nitric oxide concentrations within the treatment modality</td>
<td>Insignificantly decreased</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>AX vs other groups</td>
<td>NS</td>
<td>Significant decrease vs controls</td>
<td>-</td>
</tr>
<tr>
<td>Blood eosinophils within the treatment modality</td>
<td>NS</td>
<td>Decreased</td>
<td>NS</td>
</tr>
<tr>
<td>ACT within the treatment modality</td>
<td>Improved</td>
<td>Insignificantly improved</td>
<td>NS</td>
</tr>
</tbody>
</table>

ACT: asthma control test  
AX: reactance area at low frequencies  
NS: no significant difference or no significant changes