Randomised controlled trial of montelukast plus inhaled budesonide versus double dose inhaled budesonide in adult patients with asthma

D B Price, D Hernandez, P Magyar, J Fiterman, K M Beeh, I G James, S Konstantopoulos, R Rojas, J A van Noord, M Pons, L Gilles, J A Leff, for the Clinical Outcomes with Montelukast as a Partner Agent to Corticosteroid Therapy (COMPACT) International Study Group*

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Background: Inhaled corticosteroids (ICS) affect many inflammatory pathways in asthma but have little impact on cysteinyl leukotrienes. This may partly explain persistent airway inflammation during chronic ICS treatment and failure to achieve adequate asthma control in some patients. This double blind, randomised, parallel group, non-inferiority, multicentre 16 week study compared the clinical benefits of adding montelukast to budesonide with doubling the budesonide dose in adults with asthma.

Methods: After a 1 month single blind run in period, patients inadequately controlled on inhaled budesonide (800 µg/day) were randomised to receive montelukast 10 mg + inhaled budesonide 800 µg/day (n=448) or budesonide 1600 µg/day (n=441) for 12 weeks.

Results: Both groups showed progressive improvement in several measures of asthma control compared with baseline. Mean morning peak expiratory flow (AM PEF) improved similarly in the last 10 weeks of treatment compared with baseline in both the montelukast + budesonide group and in the double dose budesonide group (33.5 ± 30.1 l/min). During days 1–3 after start of treatment, the change in AM PEF from baseline was significantly greater in the montelukast + budesonide group than in the double dose budesonide group (20.1 ± 9.6 l/min, p<0.001), indicating faster onset of action in the montelukast group. Both groups showed similar improvements with respect to "as needed" β agonist use, mean daytime symptom score, nocturnal awakenings, exacerbations, asthma free days, peripheral eosinophil counts, and asthma specific quality of life. Both montelukast + budesonide and double dose budesonide were generally well tolerated.

Conclusion: The addition of montelukast to inhaled budesonide is an effective and well tolerated alternative to doubling the dose of inhaled budesonide in adult asthma patients experiencing symptoms and inadequate control on budesonide alone.

C hronic inflammation is recognised as a central component of asthma pathophysiology.1,2 Invading inflammatory cells in lung tissue release a wide variety of mediators and cytokines that contribute to the clinical characteristics of asthma.3 Cysteinyl leukotrienes released from eosinophils and mast cells are important pro-inflammatory asthma mediators which give rise to bronchoconstriction, mucus secretion, increased vascular permeability, smooth muscle hypertrophy, and inflammatory cell infiltration.4

Inhaled corticosteroids (ICS) affect a variety of inflammatory pathways in asthma and represent a gold standard in anti-inflammatory treatment.5 However, for some patients with persistent asthma, ICS as prescribed may fail to achieve adequate control. Increasing the ICS dose is one therapeutic option but clinical trials suggest this option may only help a proportion of patients6 and concerns exist that high dose ICS may be associated with local and systemic side effects.7 These issues have led to trials of adding other agents to ICS rather than increasing the dose, with results suggesting in the patient groups studied that adding other agents such as inhaled long acting β agonists may achieve at least similar benefits to increasing inhaled steroids.8,9 The principle that increasing inhaled steroids is only one option in patients with uncontrolled asthma receiving ICS alone has become accepted in guidelines.2

Research into the pathogenesis of asthma has led to the development of specific anti-inflammatory treatments, including montelukast, which blocks the interaction of cysteinyl leukotrienes with their receptor and resulting downstream events. Since montelukast attenuates leukotriene mediated effects, combination therapy with montelukast and ICS represents a theoretical alternative to increasing the ICS dose in patients inadequately controlled on ICS alone. Although several studies have demonstrated additive effects of montelukast with ICS,10,11 none have compared this effect with higher dose ICS as has been done with other treatments.1 A double blind non-inferiority randomised 16 week study was therefore performed to compare the clinical benefits of adding montelukast to inhaled budesonide with doubling the dose of inhaled budesonide in adult patients who were symptomatic on inhaled budesonide alone.

METHODS

Study population

Patients enrolled in the study were non-smokers or ex-smokers (stopped for at least 6 months and <12 pack year history) diagnosed with asthma for >1 year, aged 15–75 years, who were not optimally controlled as judged by investigators.
in spite of a regular ICS prescription at doses of 600–1200 µg/day for budesonide, beclometasone, triamcinolone, flunisolide, and 300–800 µg/day for fluticasone. Patients were required to have forced expiratory volume in 1 second (FEV₁) values ≥50% predicted at visits 1 and 3, together with ≥12% improvement in FEV₁, after β agonist administration, and symptoms requiring β agonist treatment of at least 1 puff/day during the last 2 weeks of the run in period.

Patients were excluded if they had other active pulmonary disorders, respiratory infection within 3 weeks of visit 1 or during the run in period, treatment in an emergency setting within 2 months of visit 1, systemic corticosteroid treatment within 1 month, cromones or leukotriene receptor antagonists within 2 weeks, long acting β₂ agonist within 1 week (astemizole 3 months), or long acting β agonists or anticholinergic agents within 24 hours. The study was approved by the appropriate ethical review committees and each patient gave written informed consent.

**Study protocol**

The study included a 4 week run in period during which patients were switched to budesonide Turbhaler (800 µg/day (200 µg, two puffs twice daily). After 1 week single blind montelukast placebo was added; β agonist use and daytime symptoms were assessed during this period to determine eligibility for randomisation and to establish baseline values. Patients were randomised to one of two treatment groups for 12 weeks. Group 1 (MONT-BUD) received montelukast 10 mg/day (one tablet at bedtime) in addition to budesonide 800 µg/day and group 2 (BUD1600) received budesonide 1600 µg/day (800 µg twice daily) while receiving oral placebo montelukast. Budesonide in both groups was identical in appearance. Patients were instructed to withhold inhaled β agonist (for 6 hours) and short acting antihistamines (within 24 hours) before clinic visits (every 4 weeks).

Morning peak expiratory flow (AM PEF) was the prespecified primary end point. Other prespecified end points included initial treatment effect (days 1–3 AM PEF) and time course of morning PEF during the first 14 days of double blind treatment, daily self-reported β agonist use, daytime symptoms, nocturnal awakenings, asthma exacerbations, asthma free days (defined as any day free of oral corticosteroid use, emergency care, nocturnal awakenings, with use of ≤2 puffs of β₂ agonist), peripheral blood eosinophil counts, asthma specific quality of life, and resource utilisation. Patients assessed daytime asthma symptoms in the evening before bedtime using a validated diary card containing four questions (scored from 0 to 6 where 0 is best). Patients also recorded nocturnal awakenings and overnight β agonist use. A day with an asthma exacerbation was defined as a day with: a decrease from baseline in AM PEF of >20% or AM PEF <180 l/min or an increase in β agonist use of >70% (minimum increase of two puffs), or an increase in symptom score of ≥50%, or an asthma attack (worsening of asthma requiring an unscheduled visit to the doctor’s office, emergency room, admission to hospital, or treatment with oral corticosteroids). Patients were allowed to use short acting β₂ agonists on an “as needed” basis but were encouraged to use only the amount required. At baseline and week 12 (or on withdrawal from the study) patients completed a validated self-administered asthma specific quality of life questionnaire (questions scored from 1 to 7 where 1 is worst).

**RESULTS**

**Patients**

A total of 1192 subjects were screened and 889 patients randomised, 448 to the MONT-BUD group and 441 to the BUD1600 group; 46 patients withdrew after randomisation (20 MONT-BUD, 26 BUD1600) because of clinical adverse events (n=20), protocol deviations (n=10), lost to follow up (n=8), withdrew consent (n=6), or personal reasons (n=2). There were no clinically meaningful differences between the groups in baseline characteristics (table 1), incidence of concomitant diseases, or use of concomitant drug treatments (data not shown).

**Efficacy**

Both treatment groups had significant and progressive improvements in AM PEF compared with baseline over 12 weeks (fig 1). The improvement in AM PEF over the last 10 weeks of the 12 week treatment period with the addition of montelukast was at least as effective as doubling the budesonide dose (33.5 l/min vs 30.1 l/min in the MONT-BUD and NUD1600 groups, respectively; 95% CI=−12.9 to 4.8 for the difference). The distribution of response which was similar for both treatment arms is shown in fig 2. These findings were consistent across a variety of subgroups including sex, age,
race, prior corticosteroid dose, baseline FEV₁, baseline β agonist use, and concomitant allergic rhinitis or nasal polyps. The initial change from baseline in AM PEF during the first 3 days of treatment was rapid in onset and significantly greater in the MONT-BUD group than in the BUD1600 group (20.1 l/min vs 9.6 l/min; 95% CI –17.6 to –4.3, p<0.001, fig 3). Beta agonist use (–0.63 vs –0.44 puffs/day, 95% CI for difference 0.03 to 0.44, p<0.05) and daytime symptom improvement (–0.21 vs –0.14, 95% CI for difference –0.02 to 0.19, p>0.05) followed the same pattern.

AM PEF increased progressively during the last 10 weeks of the double blind treatment period in the MONT-BUD group (1.88 l/min per week, p<0.001) and in the BUD1600 group (2.41 l/min per week, p<0.001) but remained unchanged during the run in period (p=0.096). There was no evidence that the increase in AM PEF levelled off by the end of the study in either treatment group (p>0.05).

Progressive improvements over the 12 weeks after randomisation compared with no improvement during the run in

| Table 1 Baseline characteristics of randomised patients |
|---------------------------------|----------------|----------------|------|
| Characteristic                  | Montelukast + budesonide 800 µg/day (n=448) | Budesonide 1600 µg/day (n=441) | Total (n=889) |
| Age (years) *                   | 43 (14)        | 43 (14)        | 43 (14)       |
| Age range                       | 15–74          | 15–75          | 15–75         |
| Sex (% female)                  | 59             | 61             | 60            |
| Race (%)                        | White 77.2     | Black 0.4      | Asian 5.4     |
|                                 | Other 17.0     |                |               |
| Prestudy ICS dose, actual (µg/day)* | 730 (238)     | 543.8 (220.1) (n=112) | 778.2 (257.4) (n=103) |
| Prestudy fluticasone dose (µg/day)* | 578.0 (187.3) (n=108) | 775.2 (215.6) (n=109) | 807.3 (230.1) (n=234) |
| Prestudy beclomethasone dose (µg/day)* |                |                |                |
| Prestudy budesonide dose (µg/day)* |                |                |                |
| Prestudy triamcinolone dose (µg/day)* |                |                |                |
| Prestudy flunisolide dose (µg/day)* |                |                |                |
| Age first treated for asthma (years) * | 26 (17)       | 26 (18)        | 26 (18)       |
| Asthma duration (years) *       | 18 (14)        | 26 (15)        | 17 (14)       |
| Morning PEF (l/min) *           | 385 (130)      | 383 (133)      | 384 (131)     |
| Daily β agonist use (puffs/day) * | 2.7 (2.4)      | 2.7 (2.2)      | 2.7 (2.3)     |
| Nocturnal awakenings (median % of days) | 12.3           | 13.8           | 13.3          |
| Quality of life score *         | 4.7 (1.1)      | 4.7 (1.1)      | 4.7 (1.1)     |
| Asthma affected work/school (%) | 49.7           | 46.4           | 48.0          |
| Days missed from work/school due to asthma in previous year * | 22.7 (52.3) | 20.2 (46.0) | 21.5 (49.3) |
| Oral corticosteroid treatment in previous year (% of patients) | 39.5 | 43.4 | 41.4 |
| Number of visits with healthcare provider due to worsening asthma in previous year (for patients with at least one visit) * | 4.9 (5.1) | 4.6 (5.4) | 4.8 (5.3) |

*A mean (SD) values.
ICS=inhaled corticosteroids; PEF=peak expiratory flow; FEV₁=forced expiratory volume in 1 second.

Figure 1 Morning peak expiratory flow (AM PEF) over the 12 week treatment period (solid line=montelukast + budesonide 800 µg daily, dashed line=budesonide 1600 µg daily). Data represent the mean AM PEF measured before administration of study medication.

Figure 2 Distribution of response in mean change from baseline in morning peak expiratory flow (AM PEF) in (A) patients treated with montelukast + budesonide 800 µg daily and (B) those treated with budesonide alone in a dose of 1600 µg/day.
period were evident in terms of as needed β agonist use and daytime symptom score (fig 4). Changes from baseline in β agonist use were –0.78 and –0.75 puffs per day for the MONT-BUD and BUD1600 groups, respectively (p=0.510), and in daytime symptom score were –0.34 and –0.35 in the two groups (p=0.908). Patients in both treatment groups improved and were not significantly different with respect to improved nocturnal awakenings with asthma (from 12.3% to 2.3% and from 13.8% to 3.9% of nights in the MONT-BUD and BUD1600 groups, respectively, p=0.353), median days with asthma exacerbations (6.7% v 6.3%, p=0.781), proportion of patients requiring oral steroids or admission to hospital (1.6% v 1.9%, p=0.387), proportion of patients with respiratory infection, asthma worsening, and headache.

### Safety

Both treatment regimens were generally well tolerated with no significant differences in the number of patients with adverse events, drug related adverse events, serious adverse events, or discontinuing treatment because of adverse events. There were significantly fewer investigator diagnosed respiratory adverse events in the MONT-BUD group than in the BUD1600 group (11.6% v 16.6% of patients, p<0.05). In the MONT-BUD group 166 patients (37.1%) experienced an adverse event compared with 182 patients (41.3%) in the BUD1600 group. The most common adverse events were upper respiratory infection, asthma worsening, and headache.
DISCUSSION

This double blind randomised 16 week study in asthma patients symptomatic while receiving budesonide 800 µg/day has shown that the addition of montelukast to budesonide produced comparable and substantial improvements in asthma control compared with doubling the dose of budesonide. However, during the first 3 days of treatment montelukast + budesonide was associated with a faster onset of action, as evidenced by a significantly greater change in AM PEF and reduction in β agonist use. In addition, both groups showed comparable and progressive improvements in other end points including as needed β agonist use, daytime symptom score, nocturnal awakenings, asthma exacerbations, asthma free days, blood eosinophil counts, and asthma specific quality of life.

An interesting finding of this study was the progressive steady increase in AM PEF during the course of the 12 week treatment period in both study groups. Indeed, at the end of 12 weeks of treatment a clear plateau had not been reached. This is surprising for patients in whom montelukast had been added to the treatment regime, although these findings are consistent with a recent 6 week study by Virchow and colleagues who also found that AM PEF continued to rise on a week by week basis following treatment of asthma patients with high dose zafirlukast and high dose ICS, while no such improvement occurred in the placebo arm.13 This does, however, differ from a previous study of montelukast + ICS where an early plateau was reached.1 One explanation for this difference is that our study population might have had more severe asthma with persisting symptoms and impaired lung function in spite of a higher prestudy ICS dose, much as in the study by Virchow. This suggests a mechanism of action that is not seen in patients with milder asthma, possibly similar to that seen with higher dose inhaled steroids. The clinical implication of these results, taken with the data from Virchow et al, is that in patients with more difficult asthma we need to consider longer trials of treatment of at least 12 weeks if using a leukotriene antagonist.

The results in the group who received an increased dose of budesonide are equally intriguing, suggesting that maximal benefit is not achieved even by 12 weeks. They also appear to run counter to the idea of a very flat dose-response curve for inhaled steroids.15 It is interesting that additional benefits of high dose ICS have also been observed in several other studies.5 7 10 16 The reason for this discrepancy may be similar to that for the montelukast group. These were symptomatic patients with asthma which was significantly reversible in six of the seven clinical trials of inhaled ICS.15 However, the inclusion of patients with a maintained component of asthma and their exacerbation rate both in and out of hospital may have had a significant impact on our ability to observe differences in infrequent adverse events including those related to higher doses of ICS.

One potential limitation of the present study is the absence of a placebo group. However, during the last 2 weeks of the placebo run in period all patients received inhaled budesonide and a daily montelukast placebo tablet in a single blind fashion. Subsequently, on day 1 of the treatment phase patients were unaware that the montelukast placebo had been switched to active drug or that the ICS dose was doubled. The flat baseline, faster onset of action in the MONT-BUD group, and abrupt increase in AM PEF observed in both treatment groups is consistent with a true therapeutic effect rather than a placebo effect or regression to the mean, which would have been expected to manifest itself as a gradual rise in AM PEF beginning during the placebo run in period and continuing throughout both treatment periods of the study. In view of the flat baseline and persistent symptoms, it would also have been difficult to justify ethically maintaining patients on treatment that was not improving their asthma.

Patients with persistent asthma symptoms are typically managed by increasing the dose of ICS or adding a second therapeutic agent. Increasing the dose of ICS may be associated with a number of potential side effects and higher doses may not necessarily result in more effective control of asthma symptoms for all patients.1 19 24 26 International guidelines therefore recommend that the ICS dose should be minimised whenever possible.2 The addition of a second controller agent with a complementary mechanism of action may therefore be appropriate. The findings of the present study suggest that the addition of montelukast to ICS offers comparable asthma control to doubling the dose of ICS with a faster onset of action, and might lessen the potential risk of side effects associated with long term administration of high dose ICS.

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APPENDIX

The members of the montelukast COMPACT study group are as follows: Argentina: R Alchapor, C E Baena-Cagnani, A M Lopez, L J
Authors’ affiliations

D B Price, University of Aberdeen, Aberdeen, UK

D Hernandez, Hospital Civil de Guadalajara, Guadalajara, Mexico

P Magyar, Department of Pulmonology, Semmelweis University, Budapest, Hungary

J Fiterman, Pulcs Medical School, Porto Alegre, Brazil

K M Beek, Pulmonary Division, University Hospital, Mainz, Germany

I G James, Spring House Surgery, Bolton, Lancashire, UK

S Konstantopoulos, Pneumology Clinic, Medical School University Hospital of Ioannina, Greece

R Rojas, Tucuman National University, San Miguel de Tucumán, Argentina

J A van Noord, Atrium Medical Center, Department of Pulmonology Heerlen, The Netherlands

M Pons, Ospedale Civico, Lugano, Switzerland

L Gilles, J A Leff, Merck and Co Inc, Whitehouse Station, New Jersey, USA

David Price and Jan Leff undertook the initial study design, protocol production, co-led the evaluation, co-wrote the data analysis plan, reviewed the statistics report and prepared the manuscript for publication. Leon Gilles contributed to the study design, co-wrote the data analysis plan, undertook the statistical analysis and contributed to the manuscript. All other authors reviewed and contributed to study design, protocol review, patient recruitment, data analysis review and input into the manuscript. David Price, Leon Gilles and Jan Leff are guarantors for the paper.

Conflicts of interest: David Price either through his role at the University of Aberdeen or personally has received grants, honoraria or educational support from the UK NHS R&D programme, 3M Pharmaceuticals, Abbot Laboratories, Altana, AstraZeneca, GlaxoSmithKline, Ivax, Merck, Sharpe and Dohme, Novartis, Schering Plough and Trinity Pharmaceuticals. He does not possess any pharmaceutical shares.

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