Adjustable maintenance dosing with budesonide/formoterol reduces asthma exacerbations compared with traditional fixed dosing: A five-month multicentre Canadian study

JM FitzGerald, MR Sears, L-P Boulet, AB Becker, et al.


RESULTS: With adjustable dosing, significantly fewer patients experienced exacerbations compared with fixed dosing (4.2% versus 8.9%, P=0.002; number needed to treat=21 [95% CI 13 to 59]). Patients required 36% fewer overall doses of budesonide/formoterol (2.5 versus 3.9 inhalations/day, P<0.001), and total costs per patient were lower (difference over five months CDN$–141 [95% CI $–162 to $–116]). Asthma symptom severity (modified National Heart, Lung, and Blood Institute stage) was maintained or improved in 97% or greater of patients in both groups (pre-run-in to end of treatment). Both treatments were well tolerated.

CONCLUSIONS: Budesonide/formoterol adjustable maintenance dosing provided more effective asthma control than fixed dosing, with a lower overall drug dose and reduced total cost.

Key Words: Adjustable dosing; Asthma; Budesonide; Exacerbation; Formoterol

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Guidelines for asthma care in Canada stress the importance of adapting treatment to the individual needs of patients (1). Effective management includes patient education, avoidance of environmental trigger factors and use of appropriate medications to achieve and maintain control of asthma symptoms. Inhaled corticosteroids (ICSs) offer the best option for anti-inflammatory treatment of asthma and are used regularly to maintain long term asthma control in all but the mildest forms of the disease. The 2001 update to the Canadian Asthma Consensus Guidelines (2) recommends the addition of a long acting beta2-agonist when a low to moderate dose of ICS is insufficient to maintain control.

The Canadian guidelines recommend that asthma medications be used at the lowest dose and frequency required to maintain acceptable asthma control (2). Because patients experience considerable periodic variations in their levels of asthma symptoms, the most appropriate amount of medication required to maintain control differs over time. Fixed dosing combination regimens do not enable patients to modify their dose of maintenance medication, which may lead to overtreatment during periods of control or undertreatment during periods of worsening asthma. Adjustable maintenance dosing regimens provide patients with the flexibility to change their dose of ICS and long acting beta2-agonist as appropriate in order to maintain and prevent loss of control according to their level of symptoms.

All asthma guidelines emphasize the need to develop action plans by which patients can alter their maintenance treatment as an early intervention strategy at the beginning of an asthma exacerbation (3,4). Although action plans are generally useful (5,6), the best approach for combination maintenance therapy is not well defined.

Budesonide (ICS) and formoterol (fast and long acting beta2-agonist) are both effective drugs when inhaled regularly for the treatment of asthma and have a long history of well-tolerated use. Both drugs are effective at low doses and demonstrate dose-response relationships over the ranges used clinically (7-9), making them suitable for adjustable dosing regimens.

The aim of this study was to evaluate whether patients using an adjustable maintenance dosing regimen with budesonide/formoterol in a single inhaler could maintain or improve their asthma control by using lower levels of treatment overall in comparison with patients using a traditional fixed dosing regimen with the same agents.

METHODS

Study design
This was a randomized, open-label, parallel-group study conducted in 95 primary health care and hospital centres in Canada; approximately 70% of patients were seen by primary care physicians. After enrolment (visit 1), patients entered a one-month run-in period, during which they received two inhalations of budesonide/formoterol (Symbicort Turbuhaler) twice daily. The duration of the budesonide/formoterol inhaler used depended on an individual's prestudy ICS dose (ie, patients on a prestudy dose of 250 µg to 400 µg daily used a budesonide/formoterol 100/6 µg metered dose inhaler; patients on a prestudy dose of 500 µg to 1000 µg daily used a budesonide/formoterol 200/6 µg metered dose inhaler). At the end of the run-in period, patients attended the clinic (visit 2) and were randomly allocated sequentially, according to a computer generated schedule, to receive either budesonide/formoterol adjustable maintenance dosing (Table 1) or fixed dosing (two inhalations twice daily) for a period of five months, using the same strength of inhaler as used during the run-in period. Patients were provided with instructions on how to use their medications, and in the case of adjustable dosing, on criteria for stepping up and stepping down their dose according to perceived control of their asthma. All patients attended the clinic again for assessments at one, three and five months after randomization (visits 3, 4 and 5, respectively). Patients used their usual short acting beta2-agonist inhaler (salbutamol or terbutaline) as needed throughout the study.

Patients
The study included outpatients 12 years of age and older with a diagnosis of asthma (10) who had a forced expiratory volume in 1 s (FEV1) of 70% or greater of predicted normal (2 h or less after inhalation of a short acting beta2-agonist or 6 h or less after inhalation of a long acting beta2-agonist). The patients had received an ICS for at least six months before enrolment with a constant daily dose of 250 µg/day to 1000 µg/day over the previous 30 days. For patients to be enrolled, the presence of at least one of the following was required: asthma symptoms on more than two of the previous seven days, use of short acting beta2-agonists on more than two occasions in the previous seven days, or nocturnal awakenings due to asthma on more than two nights during the previous 30 days.

TABLE 1
Adjustable dosing: Criteria for step up and step down

<table>
<thead>
<tr>
<th>Dosing</th>
<th>Criteria for step up and step down</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial step down to one inhalation twice daily*</td>
<td>The patient felt well in his/her asthma and met both of the following in the previous seven days:</td>
</tr>
<tr>
<td></td>
<td>• Releiver medication on two occasions or less</td>
</tr>
<tr>
<td></td>
<td>• No nocturnal awakenings due to asthma</td>
</tr>
<tr>
<td>Step up to four inhalations twice daily for seven or 14 days</td>
<td>On two consecutive days or nights, the patient met any of the following:</td>
</tr>
<tr>
<td></td>
<td>• Releiver medication on three or more occasions/day</td>
</tr>
<tr>
<td></td>
<td>• Nocturnal awakening due to asthma</td>
</tr>
<tr>
<td></td>
<td>• Morning PEF less than 85% of the mean baseline value†</td>
</tr>
<tr>
<td>Step down from four inhalations twice daily to one or two inhalations twice daily (ie, the dose before step up) after a period of worsening asthma (ie, after a step-up period‡)</td>
<td>During the previous two days or nights, the patient met any of the following:</td>
</tr>
<tr>
<td></td>
<td>• No more asthma symptoms than before the worsening of asthma</td>
</tr>
<tr>
<td></td>
<td>• No reliever medication use</td>
</tr>
<tr>
<td></td>
<td>• No nocturnal awakenings due to asthma</td>
</tr>
<tr>
<td></td>
<td>• Morning PEF 85% or greater of the mean baseline value†</td>
</tr>
</tbody>
</table>

*Initial step-down criteria were assessed by the physician at visit 2; patients not meeting step-down criteria at visit 2 were reassessed at visits 3 and 4; the criteria were met if patients did not meet the step-down criteria after 14 days on four inhalations twice daily, they contacted the clinic and further treatment was given at the investigator’s discretion. PEF Peak expiratory flow

†Mean baseline value was calculated from seven available days within the last 10 days of the run-in period. †Patients were able to step down their dose after seven or 14 days of increased dosing (step-up period) if the criteria were met; if patients did not meet the step-down criteria after 14 days on four inhalations twice daily, they contacted the clinic and further treatment was given at the investigator’s discretion.
Patients were excluded from the study if they had a history of a near-fatal asthma attack, significant disease or disorder that could put them at risk during the study, or if they had suffered an exacerbation of asthma in the previous month. Other exclusion criteria included a smoking history of more than 10 pack-years, or recent use of inhaled sodium cromoglycate or nedocromil sodium, leukotriene modifiers, xanthines, parenteral beta2-agonists, inhaled anticholinergics or beta-blockers. Pregnancy, planned pregnancy or lactation were also exclusion criteria. Patients were not eligible for randomization if they suffered an exacerbation of asthma (as defined in the efficacy section below) during the run-in period, if they used a short acting beta2-agonist reliever on three or more occasions per day or if they had nocturnal awakenings due to asthma on two or more consecutive nights over the last seven days of the run-in period. Patients were discontinued from the study if they experienced more than two periods of worsening asthma requiring intervention with oral corticosteroids or if they used additional nonstudy anti-asthma medication, ie, not including study medication, reliever medication or medication related to the treatment of exacerbations.

The study was conducted in accordance with the principles of the Declaration of Helsinki. Institutional Review Board approval was obtained for each centre and all patients (or their parents or guardians if younger than 18 years of age) gave written, informed consent.

**Efficacy assessments**

The primary efficacy variable was the proportion of patients experiencing asthma exacerbations, defined as one or more of the following: need for additional inhaled and/or oral corticosteroids due to worsening asthma, emergency department treatment due to worsening asthma, an asthma-related serious adverse event (SAE) or study withdrawal because of the need for added asthma maintenance therapy. The additional use of an ICS refers to patients in both treatment arms prescribed an ICS for 10 to 14 days for treatment of worsening asthma. Additional ICS use for 15 or more days was classified as ‘added asthma maintenance therapy’. An additional analysis considered severe exacerbations, defined as one or more of the following: need for oral (but not inhaled) corticosteroids due to worsening asthma, emergency department treatment due to worsening asthma, or an asthma-related SAE.

Investigators assessed asthma symptom severity at visits 1 and 5 using a modification of the National Heart, Lung, and Blood Institute severity stage definitions (Table 2) (11). During visit 1, demographic details were recorded, clinical assessments were made (including asthma symptom severity status and FEV₁), and training was given on the use of the Turbuhaler inhaler and the measurement of peak expiratory flow (PEF). Thereafter, eligible subjects recorded their use of budesonide/formoterol and reliever medication, nocturnal awakenings, unscheduled health care contacts due to asthma and days off from work or school due to asthma in a diary. Patients recorded morning PEF each day during the run-in period to determine a baseline value (mean of seven available days within the last 10 days of the run-in period), but subsequently only during periods of asthma worsening. Patients were instructed to contact the investigator if the morning PEF decreased by more than 30% from baseline on two consecutive days, a maximum daily reliever dose was used on two consecutive days (or more than the maximum reliever dose was used on one day), symptoms were not relieved, or there was sudden worsening of shortness of breath. Maximum daily reliever doses were terbutaline Turbuhaler 3 mg (12 years and older), salbutamol inhalation aerosol 800 µg (18 years and older) or 400 µg (12 to 17 years old), salbutamol Ventodisk or Rotahaler (GlaxoSmithKline Inc, Canada) 1600 µg (18 years and older) or 800 µg (12 to 17 years old), or generic equivalents.

During clinic visits, diary records were reviewed and the patient was interviewed to identify asthma exacerbations and adverse events (AEs). The level of treatment and occurrence of exacerbations were identified objectively according to clearly defined and unambiguous criteria, and records were subject to systematic and regular audit to confirm that this process was completed accurately. Compliance was monitored via diary records and encouraged at each clinic visit. At visits 2, 4 and 5, patients 18 years of age and older were asked to assess their overall satisfaction with a single score on a seven-point scale (ranging from ‘a great deal worse’ [score –3] to ‘a great deal better’ [score +3]).

**Health economic data**

Health economic analysis was conducted from a societal perspective, capturing medication and health care resource use (direct costs), and days of lost productivity (absences from work or school, including caregivers) (indirect costs) collected from patient diaries and case report forms. The use of health care services was multiplied by their unit costs obtained from public sources (medication costs [12,13], physician visits [14], hospitalization and emergency department visits [15], and health care costs associated with acute asthma [personal communication, Ontario Nursing Association, April 2002]). Costs of lost productivity were calculated for patients by multiplying the number of days absent from work or school by the standard daily wage for each type of employment: CDN$120.33/day (the average industrial aggregate wage) for full-time workers (16); one-half of that amount ($60.17) for part-time workers; and the Ontario minimum wage ($54.80/day) for those not engaged in paid work (eg, full-time students or homemakers). Unemployed patients were assigned Canadian Federal Employment Insurance benefits of CDN$66.18/day (55% of full-time earnings).
Safety variables

AEs, SAEs (ie, death, life-threatening event, inpatient hospitalization or prolongation of existing inpatient hospitalization, persistent or significant disability, congenital abnormality or birth defect, or important medical event) and discontinuations due to AEs were identified by the investigators during the study from patient diaries and interviews at clinic visits. The investigators decided whether there was a causal relationship between treatment and the SAEs.

Statistical analysis

Comparisons of efficacy variables between treatment groups used an intention-to-treat approach. All patients who received at least one dose of study medication were included in the safety population. Two-sided tests were used, and P values of 0.05 or less were considered to be statistically significant. All proportions were calculated from the inverse of the absolute risk reduction for each treatment group during the randomized treatment period 

The total cost for each patient (direct plus indirect costs) was calculated from the inverse of the absolute risk reduction for each treatment group during the randomized treatment period. The costs were compared between the two treatments using the CMH test. The numbers of asthma exacerbations and severe exacerbations were compared between the two treatments using a Wilcoxon test. The numbers of asthma exacerbations and severe exacerbations were compared between the two treatment groups using an analysis of covariance model with fixed factors and the daily baseline count as a covariate. Changes in asthma symptom severity category and the overall treatment evaluation rate ratio, 0.43, P=0.001. The use of oral corticosteroids, of exacerbations by 57% compared with fixed dosing (exacerbation rate ratio 0.43, P=0.001). To prevent one patient from experiencing an exacerbation, the number needed to treat with adjustable dosing was 21 (95% CI 13 to 59). Only five patients had multiple exacerbations (one on adjustable dosing, four on fixed dosing). Overall, use of adjustable dosing reduced the mean number of exacerbations by 57% compared with fixed dosing (exacerbation rate ratio 0.43, P=0.001). The use of oral corticosteroids, the most common criterion for defining an exacerbation in both treatment groups, was more frequent with fixed dosing than with adjustable dosing. Emergency department treatment was required in approximately 2% of patients in both groups. The frequencies of all categories of exacerbations are shown in Figure 2. Of the eight patients categorized as having an exacerbation by virtue of receiving added maintenance therapy, in only one case was this a result of using additional ICSs for 15 days or longer. In some cases, patients met more than one criterion for identifying an exacerbation. The time to first exacerbation was significantly longer for patients on adjustable dosing than for those on fixed dosing (P=0.001) (Figure 3).

A smaller proportion of patients in the adjustable dosing group than in the fixed dosing group had one or more severe exacerbations and use of reliever medication, the change from the daily baseline count (ie, improved, maintained and worsened) were compared between the two treatments using the CMH test.
exacerbations (3.6% [n=18] versus 6.3% [n=31], P=0.06; CMH odds ratio 0.57 [95% CI 0.31 to 1.03]). Use of adjustable dosing reduced the mean number of severe exacerbations by 47% compared with fixed dosing (rate ratio 0.53, P=0.02).

### Use of study medication

In the adjustable dosing group, 464 patients (93%) decreased their maintenance dose to one inhalation twice daily at least once during the study; 76% of patients stepped down their dose within one week of random assignment. Twenty per cent of patients stepped up their dose at least once during the study (13.2% once, 4.2% twice, 2.4% three or more times). The most common step-up period was seven days.

In the adjustable dosing group, 441 patients (88%) met the criteria for step down in dose, compared with 430 patients (87%) in the fixed dosing group. Also, 156 patients (32%) on adjustable dosing met the criteria to increase their dose at least once during the randomized treatment period, compared with 139 patients (28%) on fixed dosing. These differences between treatment groups were not statistically significant.

Overall, patients on adjustable maintenance dosing used significantly fewer inhalations of budesonide/formoterol during the randomized treatment period than patients on fixed dosing (mean 2.51 versus 3.92 inhalations/day; difference –1.41

### Table 3

Demographic and baseline characteristics in a study comparing adjustable and fixed maintenance dosing regimens of budesonide/formoterol in asthma

<table>
<thead>
<tr>
<th></th>
<th>Adjustable maintenance dosing group (n=499)</th>
<th>Fixed maintenance dosing group (n=496)</th>
<th>Total (n=995)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age in years (mean ± SD, range)</td>
<td>40.6±16.3 (12 to 85)</td>
<td>42.8±16.8 (12 to 96)</td>
<td>41.7±16.6 (12 to 96)</td>
</tr>
<tr>
<td>Number of men (%)</td>
<td>205 (41.1)</td>
<td>186 (37.5)</td>
<td>391 (39.3)</td>
</tr>
<tr>
<td>FEV₁ (L) after bronchodilator*</td>
<td>2.97±0.89 (1.1 to 8.1)</td>
<td>2.86±0.85 (0.9 to 5.7)</td>
<td>2.91±0.88 (0.9 to 8.1)</td>
</tr>
<tr>
<td>FEV₁ (% predicted normal)</td>
<td>93.0±15.6 (66.0 to 198.0)</td>
<td>92.5±15.9 (52.0 to 173.0)</td>
<td>92.7±15.7 (52.0 to 198.0)</td>
</tr>
<tr>
<td>PEF (L/min) during run-in (mean ± SD, range)</td>
<td>454±108 (217 to 770)</td>
<td>447±108 (186 to 773)</td>
<td>451±108 (186 to 773)</td>
</tr>
<tr>
<td>Classification of asthma symptom severity (n, %)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Severe persistent</td>
<td>9 (1.8)</td>
<td>7 (1.4)</td>
<td>16 (1.6)</td>
</tr>
<tr>
<td>Moderate persistent</td>
<td>210 (42.1)</td>
<td>221 (44.6)</td>
<td>431 (43.3)</td>
</tr>
<tr>
<td>Mild persistent</td>
<td>280 (56.1)</td>
<td>267 (53.8)</td>
<td>547 (55.0)</td>
</tr>
<tr>
<td>Mild intermittent</td>
<td>0 (0)</td>
<td>1 (0.2)</td>
<td>1 (0.1)</td>
</tr>
<tr>
<td>Treatment allocation – patients stratification (n, %)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Budesonide/formoterol 100/6 µg</td>
<td>140 (28)</td>
<td>129 (26)</td>
<td>269 (27)</td>
</tr>
<tr>
<td>Budesonide/formoterol 200/6 µg</td>
<td>359 (72)</td>
<td>367 (74)</td>
<td>726 (73)</td>
</tr>
<tr>
<td>Prestudy ICS dose per day (mean ± SD, range)</td>
<td>318±177 (100 to 400)</td>
<td>333±174 (200 to 400)</td>
<td>325±176 (100 to 400)</td>
</tr>
<tr>
<td>Budesonide/formoterol 200/6 µg</td>
<td>690±200 (500 to 1600)</td>
<td>685±192 (500 to 1000)</td>
<td>677±196 (500 to 1600)</td>
</tr>
<tr>
<td>Prestudy asthma therapy (n, %)</td>
<td>263 (53)</td>
<td>278 (56)</td>
<td>541 (54)</td>
</tr>
<tr>
<td>ICS only</td>
<td>236 (47)</td>
<td>218 (44)</td>
<td>454 (46)</td>
</tr>
<tr>
<td>ICS plus long acting beta₂-agonist†</td>
<td>0.4</td>
<td>0.6</td>
<td>1.0</td>
</tr>
</tbody>
</table>

*Measured at visit 1; †Includes combination inhalers. FEV₁ Forced expiratory volume in 1 s; PEF Peak expiratory flow; ICS Inhaled corticosteroids

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**Figure 2** Categories of exacerbations in each treatment group during the study. Patients who met criteria for an exacerbation in more than one category are counted once in each of these categories. *Patients who took both inhaled corticosteroids (ICSs) plus oral steroids or ICSs plus long acting beta₂-agonists; †Any asthma therapy required for long term asthma maintenance; ‡Percentage of patients with at least one exacerbation; *P=0.002. ICS Inhaled corticosteroids; SAE Serious adverse event

**Figure 3** Time to first exacerbation (Kaplan-Meier plot)
During the randomized treatment period, the mean dose of budesonide taken by patients on adjustable dosing was 36% lower than that of patients on fixed dosing (435 µg versus 682 µg).

Secondary efficacy variables

There was a marked and statistically significant shift to a lower asthma symptom severity status (modified NHLBI severity stage) in both treatment groups (P<0.001) from visit 1 (pre-run-in) to visit 5 (end of treatment period). Asthma symptom severity status was maintained or improved in 97% or greater of patients in each treatment group. At the end of the treatment period, approximately one-half of the patients in both groups were categorized as having mild intermittent asthma. There was no significant difference between the two treatment groups with regard to the degree of improvement.

The mean use of short acting beta 2-agonist reliever medication (adjustable dosing, 0.38 occasions/day; fixed dosing 0.43 occasions/day) and frequency of nocturnal awakenings (adjustable dosing, 2.5% nights; fixed dosing, 2.3% nights) was low in both groups, and there were no significant differences between the groups with regard to changes from the run-in period. At the end of the study, 378 patients (86%) on adjustable dosing and 389 patients (88%) on fixed dosing expressed feeling the same or better due to treatment; there was no significant difference between the groups.

Health economic data

Over the randomized treatment period, total asthma medication costs, direct costs and total costs were all lower with adjustable dosing than with fixed dosing (Table 4). The reduction in total cost in favour of adjustable dosing (difference in cost per patient over five months CDN$–141 [95% CI –$162 to –$116]) was largely due to the lower use, and corresponding 36% lower cost, of budesonide/formoterol by patients on adjustable dosing than by those on fixed dosing. All other differences in treatment costs were small.

Safety assessments

The total number of AEs was similar in each treatment group (adjustable dosing, n=827; fixed dosing, n=884). Individual AEs occurred with similar frequencies in both treatment groups. There were only 19 SAEs during the study: four SAEs in three nonrandomized patients, six SAEs in five patients (1.0%) on adjustable dosing and nine SAEs in seven patients (1.4%) on fixed dosing. After randomization, there was only one asthma-related SAE (fixed dosing group), which was classified as an exacerbation. There were no deaths during the study.

**DISCUSSION**

This large, multicentre study was the first study in Canada to evaluate adjustable maintenance dosing with budesonide/formoterol in a single inhaler for the management of asthma exacerbations. The study demonstrated that patients, the majority of whom were seen by primary care physicians, can use adjustable maintenance dosing to maintain better control of their asthma, using a lower overall dose at a reduced cost compared with a traditional fixed dosing regimen using the same drugs. It is important to note that the use of adjustable dosing was associated with a significant reduction in the
Budesonide/formoterol adjustable dosing in asthma

Budesonide/formoterol adjustable dosing is consistent with the recommendations for maintenance treatment in the Canadian Asthma Consensus Guidelines (1,2), ie, establish and then maintain control with the lowest effective dose of medication. Adjustable dosing provided a convenient means of reducing medication to a lower dose, while maintaining acceptable asthma control. Although patients on adjustable dosing used 36% fewer inhalations of budesonide/formoterol than patients on fixed dosing, they did not use more reliever medication; nocturnal awakenings were also similarly low in both groups. Furthermore, the improved exacerbation control achieved with the use of budesonide/formoterol adjustable dosing was significantly less costly than fixed dosing. Both treatments were well tolerated and there were no safety issues identified in the study. The roles played by the increased dose of the individual components of the budesonide/formoterol single inhaler in reducing exacerbations are unknown; further studies are needed to identify the relative importance of increasing the ICS and long acting beta2-agonist components alone or in combination. In a further Canadian study by FitzGerald et al (21), doubling the maintenance dose of budesonide was no better than continuing the maintenance dose. However, a recent study examining the effect of doubling the dose of budesonide, formoterol or the combination of budesonide plus formoterol on the increased protection against indirect bronchoconstrictor challenge from adenosine, indicated that a significant increase in the level of protection was only apparent after a double dose of the combination (22). Thus, the simultaneous increase in the dose of budesonide/formoterol may be an important attribute of this treatment approach.

The benefits of adjustable dosing compared with fixed dosing on exacerbation control in this study were increasingly apparent over time, suggesting that adequate anti-inflammatory treatment was maintained despite less use of budesonide/formoterol in the adjustable dosing group. However, further studies are needed to determine whether the benefits of treatment shown over the six months of this study would be maintained over a longer time period. Additionally, longer term studies may provide answers as to whether adjustable maintenance dosing with budesonide/formoterol has benefits for airway remodelling and reduction of inflammatory markers.

The scientific principle supporting the role of adjustable dosing for asthma medication is central to all asthma guidelines. These guidelines, in general, support the need for an asthma education program to complement any pharmacological intervention, and central to all of these programs is the use of a written action plan (6,23). Simple, predominantly symptom-driven action plans that provide a simple, variable dose schedule and enable a rapid increase in maintenance medication during the early stages of an exacerbation – such as the adjustable dosing regimen in the present study – would seem appropriate.

In summary, this study demonstrates that an adjustable maintenance dosing regimen for budesonide/formoterol in a single inhaler can be used effectively in a primary care setting to provide better asthma control than a traditional fixed dosing regimen, with a lower overall dose of medication at a lower cost.
CONCLUSIONS

Adjustable maintenance dosing with budesonide/formoterol in a single inhaler allows patients the flexibility to modify their dose to maintain control of their asthma symptoms at a similar level as with traditional fixed dosing; however, there is significantly lower overall drug use at a lower cost. Patients on budesonide/formoterol (Symbicort®) adjustable maintenance dosing were significantly less likely to suffer an asthma exacerbation, as defined by the need for additional medical intervention, than patients on fixed dosing. Both adjustable and fixed dosing regimens were well tolerated. Budesonide/formoterol adjustable maintenance dosing is consistent with Canadian asthma treatment guidelines because it provides effective asthma control at an appropriately reduced dose of medication.

ACKNOWLEDGEMENTS:

REFERENCES

ERRATUM


In the method section of the French abstract of the article, “Adjustable maintenance dosing with budesonide/formoterol reduces asthma exacerbations compared with traditional fixed dosing: A five month multicentre Canadian Study”, adjustable dosing (posologie variable) was incorrectly described. The correct French abstract along with the original English abstract is republished below. We apologize for any inconvenience this may have caused.


BACKGROUND: Adjustable maintenance dosing with budesonide/formoterol in asthma.

OBJECTIVES: To compare adjustable and fixed maintenance dosing regimens of budesonide/formoterol in asthma.

METHODS: This was an open-label, randomized, parallel-group, multicentre, Canadian study of asthma patients (aged 12 years or older, postbrachial dialator forced expiratory volume in 1 s 70% or greater of predicted normal). Following a one-month run-in on budesonide/formoterol (100/6 µg or 200/6 µg metered doses, two inhalations twice daily), 995 patients were randomly assigned either to continue on this fixed dosing regimen or to receive budesonide/formoterol adjustable dosing (step down to one inhalation twice daily if symptoms were controlled or temporarily step up to four inhalations twice daily for seven or 14 days if asthma worsened). The primary efficacy variable was the occurrence of exacerbations (requiring oral or inhaled corticosteroids, emergency department treatment, serious adverse events or added maintenance therapy because of asthma). The patients were divided into two groups, one with fixed dosing and the other with adjustable dosing.

RESULTS: With adjustable dosing, significantly fewer patients experienced exacerbations compared with fixed dosing (4.0% versus 8.9%, P=0.002; number needed to treat=21 [95% CI 13 to 59]). Patients required 36% fewer overall doses of budesonide/formoterol (2.5 versus 3.9 inhalations/day, P<0.001), and total costs per patient were lower (difference over five months CDN$–141 [95% CI –$162 to –$116]). Asthma symptom severity (modified National Heart, Lung, and Blood Institute stage) was maintained or improved in 97% or greater of patients in both groups (pre-run-in to end of treatment). Both treatments were well tolerated.

CONCLUSIONS: Budesonide/formoterol adjustable maintenance dosing provided more effective asthma control than fixed dosing, with a lower overall drug dose and reduced total cost.

Key Words: Adjustable dosing; Asthma; Budesonide; Exacerbation; Formoterol

Traitement d'entretien à posologie variable de budesonide et de formoterol : plus efficace pour réduire les exacerbations d'asthme que le traitement d'entretien courant à posologie fixe. Étude multicentrique, menée au Canada, pendant cinq mois

CONTEXTE : Le traitement d'entretien à posologie variable de budesonide et de formoterol au moyen d'un seul aérosol (Symbicort, AstraZeneca, Lund, Sweden) pourrait s'avérer une formule pratique pour équilibrer l'asthme tout en utilisant la posologie la plus faible possible de médicaments.

OBJECTIF : Comparer l'efficacité des traitements à posologie variable et à posologie fixe de budesonide et de formoterol pour la maîtrise de l'asthme.

MÉTHODE : Il s'agit d'une étude multicentrique, menée au Canada, avec randomisation et groupes parallèles, au su des parties, auprès de patients asthmatiques (âge : 12 ans et plus; volume expiratoire maximal par seconde après utilisation d'un bronchodilatateur : 70 % ou plus du volume normal prévu). Après une phase de présélection comportant la prise régulière de budesonide et de formoterol (100 µg/6 µg OU 200 µg/6 µg à raison de 2 bouteilles, 2 fois par jour) au moyen d'un inhalateur à poudre avec une séche pendant un mois, 995 patients ont été dirigés au hasard vers le traitement à posologie fixe (celui déjà en cours) ou vers le traitement à posologie variable (mélange de budesonide et de formoterol : réduire à 1 bouteille, 2 fois par jour; si soulagement des symptômes ou augmentation temporaire à 4 bouteilles, 2 fois par jour; pendant 7 ou 14 jours si intensification de l'asthme). Le principal critère d'évaluation de l'efficacité du traitement était le nombre d'exacerbations (nécessitant des corticostéroïdes par voie orale ou en aérosol, un traitement au service d'urgence ou l'adjonction d'un traitement d'entretien ou encore entraînant des complications graves).

RÉSULTATS : Un nombre significativement moins élevé de patients soumis au traitement à posologie variable ont connu des exacerbations que de patients soumis au traitement à posologie fixe (4.0 % contre 8.9 %; p=0.002; nombre nécessaire de patients à traiter : 21 [IC à 95 % : 13-59]). Les patients ont eu besoin de moins de doses de budesonide et de formoterol dans l'ensemble (36 %; 2.5 contre 3.9 bouteilles par jour; p=0.001) et il en a coûté moins cher au total par patient (écart sur 5 mois: –141 $ [IC à 95 % : –162 $ à –116 $]. Le degré de gravité des symptômes d'asthme, selon la classification modifiée du National Heart, Lung and Blood Institute, s'est maintenu ou a diminué chez au moins 97 % des patients dans les deux groupes (depuis la phase de présélection jusqu'à la fin du traitement). Les deux traitements ont été bien tolérés.

CONCLUSION : Le traitement d'entretien à posologie variable de budesonide et de formoterol a été avéré plus efficace pour équilibrer l'asthme que le traitement à posologie fixe, tout en étant associé à une diminution du nombre total de doses et du coût total par patient.