Satisfaction levels and asthma control amongst Malaysian asthmatic patients on budesonide/formoterol maintenance and reliever therapy: experience in a real-life setting

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Introduction

Asthma remains a disease that brings considerable morbidity and mortality worldwide despite modern effective treatment.1 Over recent years, there has been increasing evidence to support an approach of using budesonide/formoterol as both maintenance and reliever therapy (Symbicort SMART®, AstraZeneca AB, Sodertalje, Sweden) in chronic persistent asthma. Several studies involving both adult2–4 and pediatric5 patients have shown that such an approach reduces...
severe asthma exacerbations and improves day-to-day asthma control, compared with conventional maintenance inhaled corticosteroid (ICS) or combination ICS and long-acting β₂-agonist (LABA) with as-needed bronchodilator. Although there is a sound scientific basis in support of budesonide/formoterol maintenance and reliever therapy, acceptability and practicality can be best addressed by a real-life effectiveness study. Furthermore, the sociocultural and local healthcare systems may play a significant role here.

Studies have shown that patient satisfaction is greater when ICS and LABA are administered in a single inhaler than separately for maintenance therapy. Although there are two studies reporting on patient satisfaction with such an approach, there is currently no published prospective multicenter study that specifically examines patient satisfaction with budesonide/formoterol maintenance and reliever therapy and its potential for improving treatment adherence and acceptability. Although real-life effectiveness of budesonide/formoterol maintenance and reliever therapy has been well studied elsewhere, it is prudent to evaluate whether such benefit can be translated into a multiethnic population like Malaysia.

With this purpose in mind, we tested our hypothesis in the local Malaysian patient population in a prospective, open-label, single-cohort study in ten major hospitals across Peninsular Malaysia. The primary outcome was change in mean score from baseline in patient-reported satisfaction level using the Satisfaction with Asthma Treatment Questionnaire (SATQ) to the average of 3 months and 6 months. The secondary outcomes were change in mean score from baseline in asthma control level as defined by Asthma Control Questionnaire 5-item Version (ACQ-5) to the average of 3 months and 6 months after starting budesonide/formoterol maintenance and reliever therapy. Physician-assessed asthma control levels defined by Global Initiative for Asthma (GINA), frequency of reliever medication use, and asthma exacerbation rates were also determined.

**Methods**

**Patients**

All adult patients ≥18 years old with an asthma history of ≥6 months treated with at least 3 months of daily ICS were considered for the study. Patients who remained partially controlled or uncontrolled as defined by the GINA guidelines and on daily doses of 200–1000 µg beclomethasone dipropionate (or equivalent) for at least 4 weeks prior to study entry were eligible. Patients currently on budesonide/formoterol maintenance and reliever therapy; those who had used oral, rectal, or parenteral glucocorticoids; those who had respiratory tract infection within 30 days prior or had known or suspected hypersensitivity to budesonide, formoterol, or inhaled lactose; those who were pregnant or lactating; and those diagnosed with chronic obstructive pulmonary disorder or another chronic respiratory or significant medical condition were excluded. Patients were able to discontinue from the study treatment and assessments at any time. Specific reasons for discontinuing from the study were voluntary discontinuation, safety reasons, severe noncompliance, incorrect enrolment, lost to follow-up, or pregnancy. The study protocol was approved by the ethics and research committees of participating centers and conducted in accordance with the Declaration of Helsinki and Good Clinical Practice guidelines (Clinical Trial Registration Number: NCT00576316). Written informed consent was obtained from all recruited patients.

**Study design**

This was a 6-month, open-label, single-cohort, multicenter study conducted in chest clinics of ten urban-based hospitals across Peninsular Malaysia. The study involved a screening/baseline visit where eligibility was assessed and informed consent obtained. Relevant baseline demographic data including educational level, baseline SATQ score, and ACQ-5 score were obtained prior to initiation of budesonide/formoterol maintenance and reliever therapy. Patients were followed up at 3 months and 6 months. Recruited patients were prescribed one or two inhalations of budesonide/formoterol Turbuhaler (160/4.5 µg per inhalation; AstraZeneca AB) twice daily as maintenance treatment, and additional inhalation of budesonide/formoterol Turbuhaler (160/4.5 µg per inhalation) as needed for breakthrough symptoms. Initial dose for maintenance treatment and its subsequent adjustment were left to the clinical judgment of the treating physicians based on GINA-defined treatment objectives. In all visits, relevant data on treatment satisfaction, asthma control level, asthma symptoms, asthma exacerbations, frequency of rescue treatment, current medications, and occurrence of adverse events were collected in a standard case record form.

**Patient satisfaction assessment**

Patient satisfaction with their current treatment was assessed using the self-administered SATQ, which consisted of an overall score plus individual scores from four domains of satisfaction (ie, effectiveness of treatment, ease of use, medication burden, and side effects or worries). A higher score indicates greater satisfaction. Changes in mean scores from baseline to the average of the two follow-up visits were used as our primary outcome measure of patient satisfaction.
Asthma control assessment

The self-administered ACQ-5 was used for assessing asthma control. Lower scores indicate better control, and a change of −0.5 or more (the minimal clinically important difference [MCID]) represents clinically important improvement. In our study, the secondary outcome of asthma control was based on changes in mean scores from baseline to the average of the two follow-up visits, and the proportion of patients achieving MCID at 3 months and 6 months. Asthma control was also assessed by treating physicians in accordance with GINA guidelines at each visit.

Data analysis

The data were analyzed on an intention-to-treat basis. Changes in mean scores of SATQ and ACQ-5 were assessed using two-sided paired t-tests, and a P-value of ≤0.05 was considered significant. The Wilcoxon rank-sum test was used for comparisons of nonparametric data.

Calculation of SATQ

Scores for negatively phrased questions were reversed before calculation of the overall and domain scores, such that for all questions a higher score indicated greater satisfaction.

The overall (26 questions) score was calculated for each subject using the following formula:

\[ SATQ \text{ overall score}, x = \frac{\sum_{i=1}^{26} qs_i}{26} \]  

(1)

The domain (relevant questions) score was calculated using the following formula:

\[ SATQ \text{ score}, x_{d1} = \frac{\sum_{i=1}^{n} qs_i}{n} \]  

(2)

The average of SATQ at months 3 and 6 for each subject was calculated by using the following formula:

\[ Average \ SATQ \text{ score at visit 3 and 4}, x = \frac{\sum_{i=1}^{26} \left( (qs \text{ at visit 3}) + (qs \text{ at visit 4}) \right)}{26} \]  

(3)

The overall and domain group mean scores were calculated by the sum of mean scores divided by 195 subjects.

For subjects who discontinued before month 6, data up to the point of discontinuation were used to estimate response to treatment. No MCID has yet been established for the SATQ.

Calculation of ACQ

The ACQ-5 score was the mean of five responses. At least four out of the five questions had to be answered for a valid score. The mean group ACQ-5 scores at baseline, 3 months, and 6 months were calculated using the following formula:

\[ Sum\{\frac{\Sigma_{i=1}^{5} q_s}{5}\}/195 \]  

(4)

Results

A total of 195 patients of the 201 eligible patients who were recruited completed the study. Most of the patients were female, 79% were middle aged (mean age 46 years), and 76% were Malays. More than half the patients were in step three asthma severity requiring two or more controller medications. Thirty-seven percent of patients were on ICS/LABA combination therapy. Slightly over a quarter of these patients were considered by their attending physician to have uncontrolled asthma (Table 1). The proportions of patients judged to have controlled, partly controlled, or uncontrolled asthma at study entry were 0%, 73%, and 27%, respectively.

The average SATQ mean (standard deviation) score between 3 months and 6 months had also significantly improved to 5.5 (0.63) \((P < 0.001)\) from 5.1 (0.76) at baseline. The mean SATQ score was overall significantly improved to 5.4 (0.63) in 3 months \((P < 0.001)\) and to 5.5 (0.63) at

Table 1 Clinic-demographic characteristics (n = 195)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>42 (21.5)</td>
</tr>
<tr>
<td>Mean age, yrs (range)</td>
<td>46 (18–79)</td>
</tr>
<tr>
<td>Ethnicity</td>
<td></td>
</tr>
<tr>
<td>Malay</td>
<td>148 (76)</td>
</tr>
<tr>
<td>Chinese</td>
<td>12 (6.6)</td>
</tr>
<tr>
<td>Indian</td>
<td>34 (17.4)</td>
</tr>
<tr>
<td>Education level</td>
<td></td>
</tr>
<tr>
<td>Primary</td>
<td>41 (21)</td>
</tr>
<tr>
<td>Secondary</td>
<td>106 (54.3)</td>
</tr>
<tr>
<td>Diploma</td>
<td>29 (14.8)</td>
</tr>
<tr>
<td>University degree</td>
<td>19 (9.7)</td>
</tr>
<tr>
<td>Medication types</td>
<td></td>
</tr>
<tr>
<td>ICS</td>
<td>93 (47.18)</td>
</tr>
<tr>
<td>ICS + LABA (2 separate inhalers)</td>
<td>28 (14.36)</td>
</tr>
<tr>
<td>ICS + LABA (1 combined inhaler)</td>
<td>72 (36.92)</td>
</tr>
<tr>
<td>Others</td>
<td>3 (1.54)</td>
</tr>
<tr>
<td>Asthma control level</td>
<td></td>
</tr>
<tr>
<td>Controlled</td>
<td>0 (0)</td>
</tr>
<tr>
<td>Partially controlled</td>
<td>142 (72.82)</td>
</tr>
<tr>
<td>Uncontrolled</td>
<td>53 (27.18)</td>
</tr>
</tbody>
</table>

Notes: ICS = inhaled corticosteroids; LABA = inhaled long-acting β₂-agonist; Others includes theophylline and montelukast; Unless otherwise specified: Global Initiative for Asthma (GINA) 2008 Guidelines.
There were also significant improvements of the mean SATQ scores in the four SATQ domains. The effectiveness score increased from 5.7 (1.00) to 6.3 (0.73), \( P < 0.001 \). The ease of use score increased from 6.0 (0.87) to 6.3 (0.65), \( P < 0.001 \). The burden of medicine score increased from 4.3 (1.13) to 4.6 (0.79), \( P < 0.05 \). The side effects or worries score increased from 4.0 (1.40) to 4.3 (1.21), \( P < 0.001 \) (Figure 1). There was no significant difference between the 3 months and 6 months for both the overall and individual domain mean SATQ scores; most improvement in overall satisfaction was evident by 3 months.

The mean SATQ overall scores improved significantly on Symbicort single inhaler therapy whether subjects were previously on ICS inhaler alone (5.06 vs 5.49, \( P < 0.0001 \)), ICS/LABA as separate (4.86 vs 5.54, \( P = 0.0002 \)), or combination inhaler (5.40 vs 5.53, \( P = 0.0442 \)) (Figure 2).

At 3 months, the proportion of controlled, partly controlled, and uncontrolled asthma had changed to 54%, 36%, and 10%, respectively, with a shift in favor of better control. At 6 months, the proportions were 67%, 23%, and 10%, respectively, with a further shift toward better control (Figure 6).

At baseline, 161 (80%) patients reported one or more asthma exacerbations in the preceding 12 months. At 3 months and 6 months, only 30 (15%) patients reported one or more asthma exacerbations over the past 3 months, respectively. At baseline, 163 (81%) patients reported use of rescue medication one or more times per week. At 3 months and 6 months, only 90 (45%) and 74 (37%), respectively, had used rescue medication one or more times per week (Figure 7). A total of 146 (73%) patients were treated with two inhalations of budesonide/formoterol twice a day at
study entry. At 3 months and 6 months, 33% and 34% of patients were stepped down by treating physicians to one inhalation of budesonide/formoterol twice a day as maintenance therapy.

Adverse events that were considered causally related to use of Symbicort SMART were mostly local side effects of IHC. They were pharyngitis (n = 5), voice hoarseness (n = 1), dry throat (n = 1), oral thrush (n = 1), disturbed taste in the mouth (n = 1), sneezing (n = 1), and skin pruritis (n = 1).

Discussion

In this open-label, real-life practice study, we found that our patients with persistent asthma were overall more satisfied with budesonide/formoterol maintenance and reliever therapy than with their previous treatment regime. Satisfaction was observed in all the individual domains of treatment effectiveness, ease of use, burden of medicine, and side effects or worries. The level of asthma control based on ACQ-5 was also significantly improved. Physician-assessed asthma control, asthma exacerbations, frequency of reliever use, and maintenance dosing were similarly reported to be better after budesonide/formoterol maintenance and reliever therapy. These achievements were apparent at 3 month follow-up and occurred in the local context of multiethnic Malaysian patients of different hospitals and varying educational levels. The assessment of control by physician discretion reflects real-life practice in using international guidelines.

Our mean change in overall SATQ score at 6 months was 0.4. This compares favorably with previous studies using such an approach.\textsuperscript{10,11} Although all individual domains of satisfaction were significantly increased compared with previous treatment regimes, the greatest increase occurred in the domain of effectiveness. This subjective perception of improved efficacy is supported by objective measures of improved asthma control in our study, such as reliever use and ACQ scores. Improvement in satisfaction may be attributable to the improved efficacy of treatment by the addition of an LABA. However, more than half of the subjects were already receiving LABA either as combination or as add-on prior to initiation of budesonide/formoterol maintenance and reliever therapy. This group still demonstrated significant improvement in SATQ overall scores (Figure 2). In nearly 10% of the patients, maintenance treatment was successfully reduced from two inhalations twice a day to one inhalation twice a day at 3 months and 6 months, respectively. This observation lends support to the clinical efficacy of budesonide/formoterol maintenance and reliever therapy as a treatment strategy, consistent with other findings.

With regard to increased satisfaction on ease of use and burden of medicine, budesonide/formoterol maintenance and reliever therapy are likely to have contributed to this, because the treatment requires only a single inhaler, not multiple inhalers, as the treatment regime. Studies that looked at patient preference have shown that, for asthmatic patients, a rapid and effective relief of symptoms is their highest concern.\textsuperscript{16,17} Our findings of improved ease of use and reduced burden of medicine with such an approach suggest that such a “simplified” approach does not compromise on patient preference for rapid and effective treatment of asthma symptoms. Formoterol, the long-acting bronchodilator component, has been shown to provide symptom relief as rapid as any other short-acting β₂-agonist.\textsuperscript{18} Our findings also suggest that budesonide/formoterol maintenance and reliever therapy can successfully be taught and learnt in patients.
of varying levels of education who were previously on conventional regimes that used separate inhalers for regular and rescue use. Patient satisfaction was demonstrated early in the course of treatment, and it is therefore important that practitioners recognize this to avoid treatment discontinuation and to improve compliance.

The use of patient reported satisfaction level and asthma control as key outcome measures in an open-label study such as ours warrants careful interpretation. This is highlighted by a recent asthma study showing that subjective measurements based on patients’ perspectives does not reliably differentiate between active and inert treatments.\(^{19,20}\) This is particularly relevant in our study where patient satisfaction can be affected in more ways than the active treatment alone. Furthermore, the absence of a comparator treatment in our study can compromise further the validity of our findings.

Although there are three studies that reported previously on patient satisfaction with such an approach, to our knowledge, our study is the first major multicenter study that specifically looked at patient satisfaction on budesonide/formoterol maintenance and reliever therapy. In a small, local, Malaysian, real-life, effectiveness study of severe persistent asthmatics, Loh et al\(^9\) demonstrated that most patients have

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**Figure 5** Change in ACQ-5 score by pre-study asthma medication. 

**Abbreviations:** ACQ-5, Asthma Control Questionnaire 5-item; ICS, inhaled corticosteroid; LABA, long-acting \(\beta_2\)-agonist.

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**Figure 6** Proportion of patients assessed by physicians as having uncontrolled, partly controlled and controlled asthma (according to Global Initiative for Asthma, GINA) at baseline and after Symbicort SMART at 3 and 6 months.

**Figure 7** Proportion of patients reporting \(\geq 1\) use of their reliever medication in the preceding week.

**Abbreviations:** SABA, short-acting \(\beta_2\)-agonist; prn, when required.
found that budesonide/formoterol maintenance and reliever therapy met their expectations for effectiveness, simplification of treatment, and ease of use. In the same study, the patients on budesonide/formoterol maintenance and reliever therapy were also shown to experience reduced frequency of rescue treatment, hospital admission rates, and increased forced expiratory volume in 1 second, compared with the conventional approach of using separate maintenance and reliever inhalers. A recently published Swiss postmarketing survey10 of 420 physicians and 2035 patients also reported that general patient satisfaction was high and that mean ACQ-5 score of asthma control improved by more than three-fold the defined MCID.

Although patient satisfaction is often assumed by means of an effective treatment regime, our findings clearly show such an association also exists with regard to ease of use and reduced burden of medicine. From the patients’ perspectives, there was also reduced concern about side effects and worries using this approach compared with previous treatment regimes, indicating that the approach is likely to be well received. This is highly relevant because medication nonadherence is a serious problem among asthmatic patients, where patient satisfaction and preference have significant influence.21 Furthermore, our study suggests the acceptance of budesonide/formoterol maintenance and reliever therapy in a real-life busy clinical setting of an Asian multiethnic country like Malaysia and in patients with varying educational achievements. This is important because cultural influences play important roles in overall asthma management,22 and studies conducted in the West should not be assumed to be translational in their findings. Currently, studies on real-life effectiveness of such an approach in other Asian countries are being pursued,23 and the increasing bulk of evidence will help to confirm or refute the consistency of the effectiveness of budesonide/formoterol maintenance and reliever therapy across cultures and countries.

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Disclosure
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