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Asthma control in patients receiving inhaled corticosteroid and long-acting beta$_2$-agonist fixed combinations. A real-life study comparing dry powder inhalers and a pressurized metered dose inhaler extrafine formulation

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Abstract

Background: Although patients have more problems using metered dose inhalers, clinical comparisons suggest they provide similar control to dry powder inhalers. Using real-life situations this study was designed to evaluate asthma control in outpatients with moderate to severe persistent asthma and to compare efficacy of fixed combinations of inhaled corticosteroids (ICS) and long acting beta-agonists (LABA).

Methods: This real-life study had a cross-sectional design. Patients using fixed combinations of ICS and LABA had their asthma control and spirometry assessed during regular visits.

Results: 111 patients were analyzed: 53 (47.7%) received maintenance therapy of extrafine beclomethasone-formoterol (BDP/F) pressurized metered dose inhaler (pMDI), 25 (22.5%) fluticasone-salmeterol (FP/S) dry powder inhaler (DPI), and 33 (29.7%) budesonide-formoterol (BUD/F) DPI. Severity of asthma at time of diagnosis, assessed by the treating physician, was comparable among groups. Asthma control was achieved by 45.9% of patients; 38.7% were partially controlled and 15.3% were uncontrolled. In the extrafine BDF/F group, asthma control total score, daytime symptom score and rescue medication use score were significantly better than those using fixed DPI combinations (5.8 ± 6.2 vs. 8.5 ± 6.8; 1.4 ± 1.8 vs. 2.3 ± 2.1; 1.8 ± 2.2 vs. 2.6 ± 2.2; p = 0.0160; p = 0.012 and p = 0.025, respectively) and the mean daily ICS dose were significantly lower.

Conclusions: pMDI extrafine BDP/F combination demonstrated better asthma control compared to DPIs formulated with larger particles. This could be due to the improved lung deposition of the dose or less reliance on the optimal inhalation technique or both.

Keywords: inhaler, fixed combinations, asthma control, extrafine
current therapeutic asthma treatments are efficient in granting good asthma control [3]. However, efficacy results reported in RCTs are often inconsistent with data observed in real-life settings where asthma control still remains an unmet need for the great majority of patients [1,4]. Despite the variety of treatment options available and the periodical update of GINA guidelines since 1995, the evidence indicates suboptimal asthma control, even if improvements are evident when comparing more recent data [1] with previous ones [5], in a great number of patients.

However, RCTs are performed under highly controlled settings and in selected populations of patients. Under these controlled conditions, good inhalation technique is usually checked and granted and the importance of compliance is emphasized. Such a level of adherence is not to be expected in real-life settings. In an effort to find a strategy leading to improved asthma care, the International Primary Care Respiratory Group (IPCRG) examined the main factors contributing to poor control of the disease [6]. Among the several identified reasons, wrong diagnosis, smoking, co-morbidities, individual variation in response to treatment, poor adherence, and poor inhalation technique have been described.

Inhalers for fixed combinations of ICS and LABA include pMDIs and DPIs. It is known that choice of the correct inhaler device is crucial since sub-optimal inhalation techniques can result in reduced drug delivery and efficacy and it should be adjusted to different patients [7,8]. A meta-analysis of RCTs comparing the same drug delivered by different devices concluded that efficacy outcomes do not differ significantly and that pMDIs and DPIs are equally efficacious in clinical settings [9,10], although studies have shown that there are fewer inhalation errors with some DPIs [11-13].

A new fixed combination of BDP/F has been recently developed. The innovative formulation of BDP/F pMDI is characterized by extrafine particle size [14] which results in improved lung deposition and allows for uniform treatment of inflammation and bronchoconstriction throughout the entire bronchial tree [15]. This can translate into beneficial asthma outcomes as demonstrated by a significant increase in asthma control vs. a combination of the same drugs (BDP plus F) given as larger (not extrafine) particles [16]. It has been reported that the extrafine formulation of BDP/F allows for a longer duration of the aerosol plume and a reduced need for hand-to-lung coordination, thus facilitating patient inhaler technique [17]. Furthermore, it has been suggested that pMDI inhalation technique for formulations that produce extrafine particles is less critical than with other MDIs [8], because lung deposition is less affected by inhalation flow [18] and coordination [19]. Moreover, the fast onset of bronchodilation may improve patient adherence [20] and contribute to efficacy.

The aim of this observational study was to evaluate the level of asthma control in a group of patients with moderate or severe persistent asthma (at the time of diagnosis) and to compare, in a real-life setting, efficacy outcomes achieved with extrafine BDP/F delivered via pMDI vs. ICS-LABA combinations (FP/S and BUD/F) delivered via DPIs.

Methods
Study design and patients
This real-life study had a cross-sectional design and involved a group of asthmatic outpatients from the Department of Pulmonology, Semmelweis University Budapest, Hungary. Patients were consecutively recruited during their regular visits to the outpatient department from May 2008 until August 2008. Male and female asthma patients aged above 18 years were eligible for inclusion in the study if: 1) they had a diagnosis of moderate or severe persistent asthma confirmed by chest physician; 2) diagnosis occurred at least 6 months before the beginning of the study; 3) they were treated with fixed combination of ICS and LABA as maintenance therapy; 4) they did not change their medication within the last 4 weeks prior to start of study; 5) they had no asthmatic exacerbation within the last 6 weeks prior to start of study. Patients were excluded if they had any other chronic disease (except chronic rhinitis), extreme obesity, or if any acute disease occurred within the last 6 weeks prior to start of study. Patients were enrolled regardless of their smoking history or habit. Asthma severity was evaluated at the time of diagnosis and assessed by the treating physician.

The study was approved by the institutional Ethics Committee of the Department of Pulmonology, Semmelweis University. No consent was obtained from the participants as it was a non-interventional retrospective data analysis of the real-life data collected on the usual visits of patients. No intervention was done only for the sake of the study.

Assessment of asthma control
Asthma control was assessed by patients using a questionnaire (Table 1) developed on the basis of the GINA Guidelines. A score ≤ 4 indicated well controlled asthma, a score >14 reflected uncontrolled asthma, and intermediate scores were indicative of partially controlled asthma. Patients were asked to fill in the questionnaire during their regular clinic visits.

To determine if other factors may influence the level of asthma control, the following information was
collected from patient records: date of diagnosis, smoking status, history of allergy, asthma maintenance and reliever therapy.

Spirometric measurements were performed and data on forced expiratory volume at 1 s (FEV1), peak expiratory flow (PEF), forced expiratory flow between 25 and 75% (FEF 25-75%), maximum expiratory flow (MEF) 25%, and MEF 75% were collected. Lung function was analyzed using an electronic spirometer (PDD-301/s, Piston, Budapest, Hungary) according to the American Thoracic Society (ATS) guidelines [21].

Statistical Analysis
Data were analyzed with the MEDCALC statistical program package. Normal distribution was checked and tested formally by the Kolmogorov-Smirnov test. Differences between groups were tested by parametric or non-parametric tests as appropriate. The Student t test was used to test differences in case of normal distribution between the groups. Kruskal-Wallis and Mann-Whitney U tests were used for statistical comparison of the groups in case of non-normal distribution. Chi-square test was used where indicated. The level of significance was set at 0.05.

Results
Clinical data of the patients
The study included 111 patients (81 women and 30 men). For the purpose of the study, patients were divided in two groups depending on the inhaler device they were using (pMDI or DPI). No significant difference in patient characteristics between these two groups was found (Table 2). The severity of patients was similar in both groups. Lung function values measured at the time of diagnosis, on the basis of which patients were characterized as moderate or severe asthmatics, were not available. Lung function values that are indicated in Table 2 were measured during patient visits (using regular asthma maintenance therapy). No significant difference was detected between the two treatment groups in terms of spirometric values.

Asthma control results
The proportions of patients with different levels of asthma control in the two study groups and in the whole study population are shown in Table 3 and on Figure 1. Comparing the two device types, a significantly higher proportion of controlled patients was observed in the pMDI group when compared to DPI group (p = 0.031; Chi-square test)

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### Table 1 Questionnaire for the assessment of asthma control

| Question                                                                 | Score |
|-------------------------------------------------------------------------|-------|
| 1. Have daytime asthma symptoms occurred during the last week (dyspnoea, shortness of breath, wheezing)? |       |
| a. No                                                                   | 0     |
| b. Yes, maximum twice a week                                             | 1     |
| c. Yes, more than twice a week                                           | 5     |
| 2. Have night-time symptoms occurred OR have you been woken up by asthma symptoms during the last week? |       |
| a. No                                                                   | 0     |
| b. Yes, but I am not sure if it was asthma                              | 1     |
| c. Yes                                                                   | 5     |
| 3. Have you felt any limitation during physical activity during the last week? |       |
| a. No                                                                   | 0     |
| b. Yes, but only during hard physical work                               | 1     |
| c. Yes, during normal daily activities                                   | 5     |
| 4. How frequently have you used reliever therapy during the last week(Ventolin/Berotec/Berodual/Salbutamol/Astrovent/Bricanyl/Symbicort reliever)?* |       |
| a. Never                                                                | 0     |
| b. Twice or less                                                         | 1     |
| c. More than twice                                                      | 5     |
| 5. Have you had an acute exacerbation/attack of asthma that resulted in emergency department/hospitalization since you started using this maintenance therapy? |       |
| a. No                                                                   | 0     |
| b. Yes                                                                  | 5     |
| Total score:                                                             |       |
| CONTROLLED ASTHMA = 0-4 points                                           |       |
| PARTIALLY CONTROLLED ASTHMA = 5-14 points                               |       |
| UNCONTROLLED ASTHMA = more than 14                                      |       |

* Ventolin/Berotec/Berodual/Salbutamol/Astrovent/Bricanyl/Symbicort were all MDI formulations.
Mean asthma control score was significantly lower for the extrafine pMDI group as compared to the DPI group \( (p = 0.016) \) indicating a better asthma control in extrafine pMDI-treated patients (Figure 2). When looking at single domains of asthma control, mean daytime symptom score was significantly lower in the extrafine pMDI group as compared to the DPI group \( (p = 0.012) \), demonstrating better symptom control in patients using extrafine pMDI (Figure 2). The same was true for rescue medication use that was significantly lower in the extrafine pMDI group \( (p = 0.025; \text{Figure 2}) \). Scores representing night-time symptoms and physical activity limitation did not differ significantly between the two groups.

Notably, the mean daily ICS dose was significantly lower in the extrafine pMDI group as compared to the DPI group, suggesting an overall better asthma control with a lower steroid load \( (p < 0.001; \text{Figure 3}) \).

Asthma control levels detected by the questionnaire adopted in this study were in good agreement with the control levels assessed by using the current GINA guidelines. According to GINA-based control assessment (that includes lung function values as well), 38.74% of the patients proved to be controlled, 38.74% and 22.52% partially controlled and uncontrolled, respectively.

### Table 2 Clinical characteristics and lung function parameters in different treatment groups

|                          | BDP/F extrafine pMDI (n = 53) | BUD/F and FP/S DPI (n = 58) |
|--------------------------|-------------------------------|----------------------------|
| Male, n (%)              | 15 (28.3%)                    | 15 (25.8%)                 |
| Age, years (mean and range) | 48 (19-84)                   | 47 (18-86)                 |
| Current smokers, n (%)   | 20 (37.7%)                    | 19 (32.8%)                 |
| Allergies, n (%)         | 40 (75.5%)                    | 41 (70.6%)                 |
| Patients using SABA for as needed medication, n (%) | 53 (100%)                   | 50 (86.2%)                 |
| Patients using BUD/F for as needed medication, n (%) | 0 (0%)                       | 8 (13.8%)                  |
| Initial diagnosis of mild persistent asthma, n (%) | 0 (0%)                       | 0 (0%)                     |
| Initial diagnosis of moderate persistent asthma, n (%) | 45 (84.9%)                   | 49 (84.5%)                 |
| Initial diagnosis of severe persistent asthma, n (%) | 8 (15.1%)                    | 9 (15.5%)                  |
| FEV1,(mean ± SE, % of predicted) | 81.71 ± 17.53               | 79.38 ± 17.71              |
| PEF (mean ± SE, % of predicted) | 78.47 ± 19.92               | 75.34 ± 22.12              |
| FEF_25-75 (mean ± SE, % of predicted) | 62.35 ± 26.18               | 56.52 ± 23.3               |
| MEF_75 (mean ± SE, % of predicted) | 71.53 ± 28.38               | 71.41 ± 29.01              |
| MEF_25 (mean ± SE, % of predicted) | 58.2 ± 26.12                | 53.74 ± 24.23              |

SABA = short acting beta2-agonist; BDP = beclomethasone dipropionate; BUD = budesonide; FP = fluticasone propionate; F = formoterol; S = salmeterol. In pMDI group spirometry data were not available in 2 patients.

### Subgroup of smoking patients

Current or previous smoking was common in asthmatic patients enrolled in this study (35%), with a mean smoking history of 19.0 ± 3.6 pack/years. No gender difference between smokers was noted. While smoking status did not influence steroid dose used, it significantly increased the proportion of uncontrolled patients as compared to never smokers \( (11\% \text{ vs. } 23\%, p < 0.05) \). Asthma control in smokers is summarized in Table 4.

### Discussion

This is the first real-life study comparing the clinical effects of BDP/F extrafine to other combinations. Previously published RCTs found that there were no differences between BDP/F extrafine pMDI and BUD/F or FP/S in terms of lung function, use of rescue medications, exacerbations and safety [22,23]. However, these studies do not reflect routine use, during which compliance or each patient’s ability to use the device may influence the result. In addition to the improved lung deposition and distribution, there is less reliance on using an optimal inhalation technique for pMDIs formulated with extrafine particles [8]. Although these were not assessed in this study, the result would include these elements. Previous studies have shown that, when

### Table 3 Proportion of patients with different levels of asthma control

|                          | BDP/F extrafine pMDI (n = 53) | BUD/F and FP/S DPI (n = 58) | All (n = 111) |
|--------------------------|-------------------------------|----------------------------|---------------|
| Controlled               | 30 (56.60%)*                  | 21 (36.21%)                | 51 (45.94%)   |
| Partially controlled     | 17 (32.08%)                   | 26 (44.83%)                | 43 (38.74%)   |
| Uncontrolled             | 6 (11.32%)                    | 11 (18.96%)                | 17 (15.32%)   |

BDP = beclomethasone dipropionate; BUD = budesonide; FP = fluticasone propionate; F = formoterol; S = salmeterol.* \( p = 0.031 \text{ vs. BUD/F and FP/S.} \)
Control status by Inhalation type

Figure 1 Proportion of patients with different levels of asthma control in the two study groups; *p = 0.031 pMDI vs. BUD/F and FP/S. BDP-beclomethasone dipropionate; BUD- budesonide; FP-fluticasone propionate; F-formoterol; S-salmeterol.

Figure 2 Daytime symptom score, rescue medication use score and asthma control total score achieved by BDP/F extrafine pMDI or BUD/F and FP/S DPI; *p = 0.012, #p = 0.025, and ^p = 0.016, respectively (Mann-Whitney Test for BDP/F vs. BUD/F and FP/S). BDP-beclomethasone dipropionate; BUD- budesonide; FP-fluticasone propionate; F-formoterol; S-salmeterol.
inhalers emitting large particles are used, the magnitude
of inhalation technique errors for a pMDI and DPI are
similar [12].

The results show that asthma control was significantly
better in patients receiving BDP/F extrafine pMDI for-
mulation compared to the DPI group. Daytime symp-
toms and use of rescue medications were significantly
lower in the BDP/F extrafine pMDI group, further sup-
porting a greater control of asthma in a real-life setting.
Asthma is an inflammatory disorder affecting both cen-
tral and peripheral airways [24,25], which justifies the
clinical need for anti-asthmatic medication which is also
able to reach peripheral airways. The inverse correla-
tion on particle size and pulmonary deposition is well
known [26]. Of the fixed combinations of ICS currently
available on the market, BDP/F extrafine pMDI drug
particles are approximately half the size in diameter
compared to FP/S and BUD/F DPIs [17]. BDP/F extra-
fine pMDI is the only fixed combination which has
been demonstrated to reach not only large, but also
small airways when administered in asthmatic patients
[15]. Accordingly, the improved drug delivery, which
enables a more uniform treatment of inflammation and
bronchoconstriction throughout the entire bronchial
tree, contributes to the higher level of asthma control
achieved with BDP/F extrafine combination in this
study. In other words, extrafine BDP/F can reach and
treat distal areas of the lung left untargeted by larger
particle sized medications, and this translates into a
greater clinical benefit for patients. A significantly
superior asthma control with the BDP/F extrafine com-
bination vs. the combination of the same drugs (BDP
plus F) given as larger particles, non extrafine agents,
has been found in a large 6-month RCT carried-out in
asthmatic patients [16]. However, in this study, the defi-
nition of asthma control was established according to
Zetterström [27] and was therefore different from the
GINA guidelines, which is a limitation of our analysis.
No differences in lung function parameters were
observed among treatments. This finding is not surpris-
ing considering that the spirometric tests performed in
this study do not provide comprehensive evaluation of
the entire bronchial tree as they are unable to properly
reflect small airway abnormalities.

Table 4 Asthma control in smoker subgroups

|                      | BDP/F extrafine pMDI (n = 20) | BUD/F and FP/S DPI (n = 19) | All (n = 39) |
|----------------------|-------------------------------|----------------------------|--------------|
| Controlled           | 12 (60%)                      | 8 (42.1%)                  | 20 (51.3%)   |
| Partially controlled | 5 (25%)                       | 5 (26.3%)                  | 10 (25.6%)   |
| Uncontrolled         | 3 (15%)                       | 6 (31.6%)                  | 9 (23.1%)    |

BDP = beclomethasone dipropionate; BUD = budesonide; FP = fluticasone propionate; F = formoterol; S = salmeterol.
In our real-life study, current or former smokers were not excluded since they represent a large proportion of asthmatic patients. Generally, the level of asthma control in smokers was similar to that in non smokers; however, the proportion of uncontrolled patients was significantly increased in these patients as compared to never smokers. Smoking asthmatics are mostly excluded from clinical trials, therefore only sparse data are available on asthma control in current or former smoker asthmatic patients.

Notably, these results were obtained with a lower mean daily ICS dose indicating that BDP/F extrafine delivers a greater efficacy per μg of steroid when compared to BUD/F and FP/S. This is of interest considering that reduced compliance in asthmatic patients is often due to steroid phobia, an issue that can be decreased by offering a medication with a low corticosteroid dose [28].

In the studied population, 45.9% of patients had controlled asthma, 38.7% were partially controlled, and 15.3% were uncontrolled. These findings are in agreement with recent observational trials focusing on asthma control. A study performed across 10 different European countries showed that asthma is suitably controlled in 45% of non-inhaled corticosteroid users and only in 15% of patients using ICS [1]. Similarly, a web-based survey to assess the prevalence of uncontrolled asthma in the United States reported that symptoms were successfully controlled in 45% of patients [29].

This real-life observational study shows that GINA-based asthma control is achieved in a good proportion of asthmatic patients treated with fixed combinations of ICS and LABA. Patients treated with BDP/F extrafine pMDI achieved a greater level of asthma control as compared to patients treated with FP/S or BUD/F larger particle combinations, suggesting that differences in inhaler devices and formulations can have an impact on important clinical outcomes and should be therefore taken into consideration when managing patients with asthma.

Conclusions

In asthmatic patients with comparable disease severity, the use of extrafine BDP/F combination formulated in a pMDI demonstrated better asthma control than using DPI fixed dose combinations formulated with larger particles. This could be due to the improved lung deposition and distribution of the dose or less reliance on the optimal inhalation technique or both.

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Authors’ contributions

WM, GG, TL participated in conception and design of the study, acquisition, analysis and interpretation of data, as well as in drafting of the manuscript. EN, LGY participated in study design, analysis and interpretation of data and have been involved in drafting of the manuscript. GN and AB participated in drafting and reviewing the manuscript. HC provided external advice and received regular updates. He participated in the design and analysis of the study as well as involvement with drafting of the paper. All authors read and approved the final manuscript.

Conflict of interest declaration

Henry Chrystyn has no shares in any pharmaceutical companies. He has received sponsorship to carry out studies, together with consultant arrangements with Almirall, AstraZeneca, Boehringer Ingelheim, Chiesi, GlaxoSmithKline, Innovata Biomed, Meda, Naop Pharmaceuticals, Orion, Teva, Trudell and UCB. Research sponsorship has also been received from grant awarding bodies (EPSRC and MRC).

Lilla Tamasi, Veronika Müller, Gabriella Galffy and György Losonczy have no shares in any pharmaceutical companies. All of them have consultant arrangements with AstraZeneca, Boehringer Ingelheim, Chiesi, GlaxoSmithKline, MSD, Pfizer and Nycomed. Andrea Bizi and Gabriele Nicolini are employees of Chiesi Farmaceutici S.p.A.

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