Improvement in 24-hour bronchodilation and symptom control with aclidinium bromide versus tiotropium and placebo in symptomatic patients with COPD: post hoc analysis of a Phase IIIb study

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Background: A previous Phase IIIb study (NCT01462929) in patients with moderate to severe COPD demonstrated that 6 weeks of treatment with aclidinium led to improvements in 24-hour bronchodilation comparable to those with tiotropium, and improvement of symptoms versus placebo. This post hoc analysis was performed to assess the effect of treatment in the symptomatic patient group participating in the study.

Methods: Symptomatic patients (defined as those with Evaluating Respiratory Symptoms [E-RS™] in COPD baseline score ≥10 units) received aclidinium bromide 400 μg twice daily (BID), tiotropium 18 μg once daily (QD), or placebo, for 6 weeks. Lung function, COPD respiratory symptoms, and incidence of adverse events (AEs) were assessed.

Results: In all, 277 symptomatic patients were included in this post hoc analysis. Aclidinium and tiotropium treatment improved forced expiratory volume in 1 second (FEV1) from baseline to week 6 at all time points over 24 hours versus placebo. In addition, improvements in FEV1 from baseline during the nighttime period were observed for aclidinium versus tiotropium on day 1 (aclidinium 157 mL, tiotropium 67 mL; P<0.001) and week 6 (aclidinium 153 mL, tiotropium 90 mL; P<0.05). Aclidinium improved trough FEV1 from baseline versus placebo and tiotropium at day 1 (aclidinium 136 mL, tiotropium 68 mL; P<0.05) and week 6 (aclidinium 137 mL, tiotropium 71 mL; P<0.05). Aclidinium also improved early-morning and nighttime symptom severity, limitation of early-morning activities, and E-RS Total and domain scores versus tiotropium (except E-RS Chest Symptoms) and placebo over 6 weeks. Tolerability showed similar incidence of AEs in each arm.

Conclusion: In this post hoc analysis of symptomatic patients with moderate to severe COPD, aclidinium 400 μg BID provided additional improvements compared with tiotropium 18 μg QD in: 1) bronchodilation, particularly during the nighttime, 2) daily COPD symptoms (E-RS), 3) early-morning and nighttime symptoms, and 4) early-morning limitation of activity.

Keywords: COPD, 24-hour bronchodilation, long-acting muscarinic antagonist, nighttime, symptoms

Introduction

Symptoms of COPD can vary in severity over a 24-hour period, and studies indicate that they are generally worse in the early morning and at nighttime.1–3 Symptoms include chronic cough, sputum production, and breathlessness, which can severely impact on a patient’s daily activities and overall well-being, and have a corresponding high socioeconomic burden.4 Estimates suggest that the frequency of nocturnal symptoms
and symptomatic sleep disturbance may exceed 75% in patients with COPD, and potential long-term consequences may include lung function changes, increased exacerbation frequency, emergence or worsening of cardiovascular disease, impaired quality of life, and increased mortality. It is therefore important that symptoms over the entire 24-hour day are identified and managed appropriately.

In order to provide appropriate therapy, clinical guidelines (Global initiative for chronic Obstructive Lung Disease [GOLD]) suggest that symptoms, airflow limitation, and risk of exacerbations are assessed. Patients are classified into one of four groups according to their symptom burden and risk of exacerbations: A, low risk, less symptoms; B, low risk, more symptoms; C, high risk, less symptoms; or D, high risk, more symptoms; current evidence suggests that bronchodilator treatment may be more effective in those patients who are considered symptomatic (ie, groups B and D).

Bronchodilator therapies are a mainstay of COPD treatment, with two classes of long-acting bronchodilators currently available: long-acting muscarinic antagonists (LAMAs) and long-acting β₂-agonists (LABAs). LAMAs inhibit the action of acetylcholine at muscarinic receptors, while LABAs enhance cAMP signaling through stimulation of β₂-adrenergic receptors, resulting in the relaxation of bronchial smooth muscle. The LAMA aclidinium bromide is a maintenance bronchodilator therapy for adults with COPD.

The efficacy and tolerability results from a Phase IIIb study in patients with moderate to severe COPD, who received either aclidinium 400 μg twice daily (BID), the active comparator tiotropium 18 μg once daily (QD), or placebo have been previously reported. Briefly, following 6 weeks of treatment, patients receiving aclidinium 400 μg BID demonstrated improvements in 24-hour bronchodilation, compared with placebo, that were comparable with tiotropium 18 μg QD. In addition, COPD symptoms significantly improved from baseline with aclidinium, but not tiotropium, compared with placebo. These results were similar to those observed in a prior 2-week Phase IIa trial. Furthermore, a recent real-world study in patients with COPD reported improvements in nighttime and early-morning symptoms, limitation of morning activities, and quality of life over 3 months with aclidinium 400 μg BID, compared with baseline. Since aclidinium has a greater impact on COPD symptoms than tiotropium, and the “more symptomatic” patient groups stand to benefit more from bronchodilator treatment than the “less symptomatic” groups, aclidinium may provide an additional therapeutic benefit over tiotropium in these patients.

This study reports the findings of a post hoc analysis, which focused on the response in the symptomatic patient group. The key objective of this analysis was to identify any differences in 24-hour lung function and symptom control between treatment with aclidinium 400 μg BID and tiotropium 18 μg QD in this population of patients.

**Methods**

**Study design and patients**

**Overall study**

This was a randomized, double-blind, double-dummy, placebo- and active comparator-controlled, multicenter Phase IIIb study in patients with moderate to severe COPD (ClinicalTrials.gov identifier: NCT01462929). Full details of the study design and inclusion/exclusion criteria have been published previously. Briefly, patients with COPD aged ≥40 years with a smoking history (current or previous) of ≥10 pack-years were eligible to enter the study. Patients with moderate to severe COPD (for whom long-acting bronchodilators are recommended) had post-salbutamol forced expiratory volume in 1 second (FEV₁) ≥30% and <80% of the predicted normal value, FEV₁/forced vital capacity <70%. Use of long-acting bronchodilators other than the investigative treatment was not permitted. Use of salbutamol pressurized metered dose inhaler (100 μg/puff) was permitted as relief medication as needed (except ≤6 hours before each visit). Patients were permitted to continue use of oral sustained-release theophylline (use of other methylxanthines was not permitted), inhaled corticosteroids, and oral or parenteral corticosteroids (equivalent to ≤10 mg/day or 20 mg every other day of prednisone) if treatment was stable ≥4 weeks prior to screening, except ≤6 hours before each visit. Oxygen therapy (except ≤2 hours before each visit) was permitted. After a screening visit, patients underwent a 2- to 3-week run-in period to assess disease stability. Eligible patients were randomized (2:2:1) to receive aclidinium bromide 400 μg BID in the morning and evening via the Genuair™/Pressair® (registered trademark of AstraZeneca group of companies; for use within the USA as Pressair® and as Genuair™ within all other licensed territories) multidose dry powder inhaler, tiotropium 18 μg QD in the morning via the HandiHaler®, or placebo for 6 weeks.

The study was approved by an independent ethics committee at each site (Table S1) and was conducted in accordance with the Declaration of Helsinki, the International Conference on Harmonisation, and Good Clinical Practice guidelines. All patients provided written informed consent.
Post hoc analysis
This post hoc analysis assessed symptomatic patients, defined as those patients with an Evaluating Respiratory Symptoms in COPD (E-RS:COPD™ [The EXACT™ and E-RSTM are owned by Evidera. Permission to use these instruments may be obtained from Evidera {exactpro@evidera.com}]; formerly known as EXAcerbations of Chronic pulmonary disease Tool [EXACT])-RS baseline score ≥ 10 units. This threshold was chosen based on data indicating that an E-RS score ≥10 units differentiated between asymptomatic (GOLD groups A and C) and symptomatic (GOLD groups B and D) patients.11

Assessments and endpoints
Lung function
Lung function was assessed over 24 hours post-dose on day 1 and at week 6. The primary endpoint was change from baseline in normalized FEV₁ area under the curve (AUC) over 24 hours post-morning dose (AUC[0–24]) at week 6. The secondary endpoint was change from baseline in normalized FEV₁ AUC over the nighttime period (AUC[12–24]) at week 6. An additional lung function endpoint was change from baseline in morning pre-dose (trough) FEV₁.

COPD symptoms
Every evening, patients completed the 14-item EXACT (recall period of “today”) via electronic diaries and daily COPD symptoms scores were derived using E-RS scoring algorithms. The E-RS uses the 11 respiratory symptom items from the 14-item EXACT and assesses both overall daily respiratory COPD symptoms (RS-Total score; score range, 0–40, with higher scores indicating more severe symptoms) and specific respiratory symptoms using three subscales (RS-Breathlessness [score range, 0–17], RS-Cough and Sputum [score range, 0–11], and RS-Chest Symptoms [the sum of three items related to chest congestion/discomfort; score range, 0–12]).12,13 E-RS Total and domain scores were assessed at baseline and over the 6-week study duration. Patients who achieved a clinically meaningful improvement from baseline (E-RS Total score ≥–2 units) were considered to be responders; this responder definition was proposed based on results from three randomized controlled trials.14 Responder status was assessed over the 6 weeks of the study.

To assess the severity of early-morning and nighttime symptoms, an additional COPD symptoms questionnaire developed by the study sponsor was completed by patients each morning via electronic diaries (5-point scale: 1= “did not experience symptoms”; 5= “very severe”) and included individual morning symptoms of cough, wheeze, shortness of breath, and phlegm (5-point scale: 0= “no symptoms”; 4= “very severe symptoms”), as well as limitation of morning activities (5-point scale: 1= “not at all”; 5= “a very great deal”). Since this study, these questionnaires have been developed and evaluated further.15,16

Safety and tolerability
Treatment-emergent adverse events (TEAEs) were recorded throughout the study.

Statistical analyses
Efficacy data are reported for the intent-to-treat population, defined as all randomized patients who received at least one dose of study medication and who had at least one baseline and post-baseline FEV₁ value. Endpoints were assessed using an analysis of covariance model with treatment and sex as factors, and age and baseline values as covariates. Between-group least squares mean differences and 95% confidence intervals were calculated for all treatment group comparisons.

Results
Patients
In all, 414 patients were randomized to treatment in the overall study (2:2:1 ratio), of which 277 were defined as symptomatic (E-RS baseline score ≥10 units) and included in this post hoc subgroup analysis (placebo: n=60; aclidinium 400 μg: n=116; tiotropium 18 μg: n=101) (Figure 1). The percentages of patients in each treatment arm of this post hoc analysis were similar to those in the primary study (placebo, 21.7% vs 20.5%; aclidinium 400 μg, 41.9% vs 41.3%; tiotropium 18 μg, 36.5% vs 38.2%, respectively).

Demographics and baseline characteristics in the subgroup of symptomatic patients were similar to those in the overall study population (symptomatic patients: mean age 62.1 years, 65.0% male, 54.5% current smokers, post-bronchodilator FEV₁ 111% of predicted). Patient demographics and baseline characteristics for symptomatic patients were also similar across treatment arms, with the exception of a higher proportion of male patients in the active treatment groups compared with placebo, and a higher proportion of patients with severe COPD in the tiotropium group (Table 1). Mean post-bronchodilator percent predicted FEV₁ and COPD symptoms scores at baseline were similar across treatment arms (Table 1).

Efficacy
Lung function
Lung function endpoints in the subgroup of symptomatic patients were similar to those in the overall population.
Aclidinium 400 μg BID and tiotropium 18 μg QD both improved FEV₁ over 24 hours (AUC₀₋₂₄ h) from baseline at week 6 compared with placebo (aclidinium, 140 mL; tiotropium, 106 mL; both P<0.01). Furthermore, treatment with aclidinium 400 μg BID and tiotropium 18 μg QD improved FEV₁ from baseline at week 6 at all time points over 24 hours, compared with placebo (Figure 2). During the nighttime period (AUC₁₂₋₂₄ h), improvements from baseline compared with placebo were greater with aclidinium 400 μg BID than tiotropium 18 μg QD on day 1 (157 vs 67 mL for aclidinium and tiotropium, respectively; P<0.001) and week 6 (153 vs 90 mL for aclidinium and tiotropium, respectively; P<0.05).

Aclidinium 400 μg BID also demonstrated improvements in trough FEV₁ from baseline versus placebo and tiotropium at day 1 (136 vs 68 mL for aclidinium and tiotropium, respectively; P<0.05) and week 6 (137 vs 71 mL for aclidinium and tiotropium, respectively; P<0.05) in symptomatic patients (Figure 3).

COPD symptoms in symptomatic patients
In this subgroup of symptomatic patients, the improvement from baseline in E-RS Total score was greater with aclidinium compared with placebo (P<0.001) and tiotropium (P<0.05) over 6 weeks (Figure 4A): −2.8 units with aclidinium versus −0.7 units with placebo and −1.6 units with tiotropium. For each of the E-RS domains, greater improvements from baseline in E-RS score in symptomatic patients were also observed for aclidinium over 6 weeks of treatment (RS-Breathlessness and RS-Cough and Sputum: P<0.05 vs tiotropium and P<0.01 vs placebo; RS-Chest Symptoms: P<0.05 vs placebo) (Figure 4A). A higher percentage of patients in the aclidinium 400 μg treatment arm were E-RS responders (52.6%) compared with placebo (28.3%; P<0.01) and tiotropium 18 μg (37.6%; P<0.05) (Figure 4B) over 6 weeks.

Overall early-morning symptom severity was reduced in the subgroup of symptomatic patients over 6 weeks with aclidinium treatment versus placebo (P<0.01) and tiotropium (P<0.05; Figure 5A). Aclidinium also demonstrated improvements in individual early-morning symptom domains; shortness of breath and cough symptom scores improved in symptomatic patients treated with aclidinium compared with placebo over 6 weeks (both P<0.05; Figure 5A). A reduction in overall nighttime symptom severity from baseline was observed over 6 weeks with aclidinium versus placebo and tiotropium in symptomatic patients (both P<0.05; Figure 5B). Numerical improvements in early-morning or nighttime symptom severity were observed for tiotropium versus placebo in this subgroup.
In symptomatic patients, limitation of early-morning activity caused by COPD symptoms was reduced from baseline over 6 weeks with aclidinium versus placebo (P<0.01) and tiotropium (P=0.05), but not with tiotropium versus placebo (Figure 5C).

Safety and tolerability

In the subgroup of symptomatic patients, the incidence of TEAEs was comparable in the placebo (26.7%), aclidinium (28.4%), and tiotropium (32.7%) groups. Similar to the overall study population, the most commonly reported TEAEs in symptomatic patients were headache (5.8%) and nasopharyngitis (5.1%). Other common TEAEs (≥2% of patients overall) were COPD exacerbation (2.5%), back pain (2.5%), and cough (2.2%). The majority of TEAEs were mild or moderate in intensity. There were few serious TEAEs (1.4% overall) and no deaths in the subgroup of symptomatic patients. In total, five patients (1.8%) discontinued due to TEAEs and one patient (0.4%) discontinued due to a serious TEAE, with COPD exacerbation being the most common cause (1.4%).

Discussion

Assessment of treatment efficacy in symptomatic patients has clinical significance, as treatment guidelines recommend that such patients are treated in order to improve lung function and reduce symptoms. This post hoc analysis was performed to evaluate the 24-hour effect of treatment with aclidinium bromide 400 µg BID, tiotropium 18 µg QD, or placebo, in 277 symptomatic patients with moderate to severe
COPD participating in a Phase III study. This subgroup of symptomatic patients constituted a substantial proportion of the overall patients (277/414; 45%).

In the subgroup of symptomatic patients, 6 weeks of treatment with aclidinium BID showed improvements from baseline in FEV₁ at all time points over 24 hours (compared with placebo), and FEV₁ was higher than tiotropium QD at most 12- to 24-hour time points. Aclidinium treatment also led to greater improvements in trough FEV₁ compared with tiotropium. Furthermore, during the nighttime period at both day 1 and week 6, improvements from baseline in FEV₁ (compared with placebo) were greater with aclidinium than with tiotropium.

Patient symptoms also improved following treatment. After 6 weeks, the improvement in E-RS score was greater with aclidinium compared with both placebo and tiotropium, as indicated by changes in RS-Breathlessness, RS-Cough and Sputum, and E-RS Total scores. In addition, the percentage of patients defined as E-RS responders increased with aclidinium compared with tiotropium or placebo. Improvement over placebo and tiotropium was also observed in early-morning symptom severity, nighttime symptom severity, individual early-morning symptoms (shortness of breath and cough), and limitation of early-morning activity caused by symptoms. Safety and tolerability in symptomatic patients appeared to be similar to that in the overall study population, with the most commonly reported TEAEs being headache and nasopharyngitis.

The overall study population included both symptomatic and asymptomatic patients. Results have been previously reported and indicated that aclidinium provided significant 24-hour bronchodilation versus placebo from day 1, with comparable efficacy to tiotropium after 6 weeks. In this post hoc analysis, some notable differences were observed in the symptomatic patient group. Improvements in bronchodilation during the nighttime period were greater with aclidinium than with tiotropium in symptomatic patients, whereas in the overall population, no differences were observed between the two treatments. Furthermore, symptomatic patients experienced a greater reduction in nighttime symptom severity from baseline to 6 weeks with aclidinium,
Efficacy of aclidinium bromide in symptomatic patients with COPD

Figure 4 Symptomatic patients: change from baseline in E-RS Total and domain symptom scores versus placebo (A) and percentage of E-RS responders (B) at week 6.

Notes: Percentage reduction is shown in brackets. Placebo N=60, aclidinium 400 μg N=116, tiotropium 18 μg N=101. *P<0.05, **P<0.01, ***P<0.001 versus placebo; P<0.05 versus tiotropium. Scores: 0–40 for Total, 0–17 for Breathlessness, 0–12 for Chest symptoms, 0–11 for Cough and sputum.

Abbreviations: E-RS, Evaluating Respiratory Symptoms; ls, least squares; OR, odds ratio.

Figure 5 Symptomatic patients: change from baseline in symptom severity in the early morning (A) and nighttime (B), and limitation of early-morning activity (C) over 6 weeks.

Notes: Percentage reduction is shown in brackets. Placebo N=60, aclidinium 400 μg N=116, tiotropium 18 μg N=101. *P<0.05, **P<0.01 versus placebo; *P<0.05 versus tiotropium. Specific symptoms (A): 1= mild, 2= moderate, 3= severe, 4= very severe; any symptom: 1= no symptoms, 2= mild, 3= moderate, 4= severe, 5= very severe. For patients with no symptom, a zero value was assigned. Nighttime symptom severity (B): 1= mild, 2= moderate, 3= severe, 4= very severe. Limitation of activity (C): 1= not at all, 2= a little, 3= moderately, 4= a good deal, 5= a very good deal.

Abbreviation: ls, least squares.
compared with tiotropium. In the overall population, although nighttime symptom severity was significantly reduced with aclidinium versus placebo, the difference between the two comparators was not statistically significant. For both the overall population and the symptomatic patient group, no differences in nighttime symptom severity were observed for tiotropium versus placebo.

While noting that the BID dosing regimen of aclidinium may have contributed to the observed improvement in nighttime efficacy versus tiotropium QD in symptomatic patients, improvement of nighttime lung function is of particular importance for patients with COPD as nighttime symptoms and poor quality sleep are common. To date, relatively few studies have demonstrated significant improvements in nighttime lung function and/or sleep quality following bronchodilator therapy.

The improvements in E-RS score observed in symptomatic patients receiving treatment with aclidinium are likely to be clinically significant since E-RS has been shown to be a valid and reliable tool for the assessment of respiratory symptoms of COPD in clinical trials. The minimal clinically important differences for the different aspects of the E-RS tool have recently been proposed: RS-Total ≥−2 units; RS-Breathlessness ≥−1 unit; RS-Cough and Sputum ≥−0.7 units; and RS-Chest Symptoms ≥−0.7 units, and the ability of the E-RS to capture treatment effects has recently been evaluated. In this study, change in E-RS Total score was −2.8 units with aclidinium versus −0.7 units with placebo and −1.6 units with tiotropium. With aclidinium, changes from baseline in individual domain scores at week 6 were −1.3 for Breathlessness, −0.6 for Chest Symptoms, and −0.8 for Cough and Sputum.

There are some potential limitations of this post hoc analysis. There was found to be a significantly higher proportion of patients with baseline bronchial reversibility in the aclidinium group compared with the tiotropium group, which may potentially account for the significant difference in efficacy observed. The apparent higher proportion of patients with severe COPD in the tiotropium group is, however, unlikely to have influenced the FEV, response in these patients, since the differences between groups were not found to be significant. Also, the 6-week study period may not be long enough to reflect a patient’s symptom burden. In addition, the E-RS threshold used to distinguish symptomatic from asymptomatic patients has not been formally validated and requires further investigation. One must also consider that unvalidated, early versions of the early-morning (Early Morning Symptoms of COPD Instrument) and nighttime symptoms (Nighttime Symptoms of COPD Instrument) questionnaires were used in this study. Both questionnaires have subsequently been developed and evaluated further, and there are published data indicating that these are valid tools for measuring COPD symptoms in large randomized trials. The use of a patient questionnaire, rather than clinical assessments, to evaluate limitation of early-morning activity could be considered a potential constraint of this study; however, the benefit of this method is that it can assess if patients are restricted in their usual morning activities, such as getting washed and dressed.

One further consideration of this post hoc analysis is that although symptomatic patients constituted 45% of the overall study population, the total sample size remains relatively small (n=414). Furthermore, it should be noted that it is possible that a different group of symptomatic patients may have been identified if the COPD Assessment Test or modified Medical Research Council criteria outlined in the GOLD report were applied.

**Conclusion**

Results from this post hoc analysis of a symptomatic patient group with moderate to severe COPD showed that aclidinium 400 μg BID provided additional improvements compared with tiotropium 18 μg QD in: 1) bronchodilation, particularly during the nighttime, 2) E-RS responder status, 3) early-morning, daytime, and nighttime symptoms, and 4) early-morning limitation of activity. These results suggest that symptomatic patients may achieve greater benefits during the nighttime with aclidinium treatment than patients with fewer symptoms.

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Disclosure

JB has received consulting fees, speaker’s fees, and travel expenses from AstraZeneca and has also received compensation for organizing or participating in advisory boards for Cytos, Boehringer Ingelheim, Almirall, AstraZeneca, Novartis, and Revotar Biopharmaceuticals. The institution where JB is currently employed has received compensation for the design, performance, or participation in single or multicenter clinical trials in the past 5 years from several companies including Almirall, Altana, AstraZeneca, Boehringer Ingelheim, Cytos, GSK, Meda Pharmaceuticals, Merck Sharp & Dohme, Mundipharma, Novartis, Pfizer, and Revotar Biopharmaceuticals. RM has received consulting fees, speaker’s fees, and travel expenses from Boehringer Ingelheim and has also received compensation for participating in advisory boards for Boehringer Ingelheim, Almirall, AstraZeneca, and Novartis. Furthermore, RM has received compensation for participation in multicenter clinical trials in the past 5 years from several companies including Almirall, AstraZeneca, Boehringer Ingelheim, GSK, Merck Sharp & Dohme, Mundipharma, Novartis, Pearl, Roche, and Takeda. A-MK is a current employee of Pulmonary Research Institute at LungenClinic Grosshansdorf; the institution received compensation for the design of and/or participation in clinical trials from Almirall, AstraZeneca, Boehringer Ingelheim, GSK, Novartis, Pfizer, Infinity Pharmaceuticals, TEVA, Stema Biologicals, Chiesi, Bayer, and Takeda. Furthermore, A-MK has received consulting fees, and speaker’s fees from AstraZeneca, Boehringer Ingelheim, and Roche. FC and EGG are employees of AstraZeneca PLC, Barcelona, Spain, and former employees of Almirall S.A., Barcelona, Spain.

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