

Clinical use of aclidinium in patients with COPD

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Abstract: Chronic obstructive pulmonary disease (COPD) is the sixth-leading cause of death in the US. The Global Initiative for Chronic Obstructive Lung Disease (GOLD) guidelines provide evidence-based recommendations for the clinical management of chronic COPD. Long-acting inhaled bronchodilators continue to be the mainstay of current management. Aclidinium bromide (Tudorza™ Pressair™) joins tiotropium as a long-acting inhaled antimuscarinic bronchodilator approved by the US Food and Drug Administration for the maintenance treatment of COPD. Early studies demonstrated aclidinium's significant bronchodilatory effects supporting once-daily dosing; however, two Phase III studies, Aclidinium Clinical Trial Assessing Efficacy and Safety in Moderate to Severe COPD Patients (ACCLAIM/COPD) I and ACCLAIM/COPD II, in which patients were randomized to receive aclidinium 200 µg daily, failed to achieve the minimal clinically important difference in improvement of trough forced expiratory volume in 1 second (FEV₁), suggesting the need for higher doses or more frequent dosing. Additional studies – Aclidinium to Treat Airway Obstruction in COPD Patients (ATTAIN) and Aclidinium in Chronic Obstructive Respiratory Disease (ACCORD) I – were undertaken to compare 200 and 400 µg twice-daily dosing. The mean improvements from baseline in trough FEV₁ in the 400 µg groups were +129 mL over 24 weeks and +124 mL over 12 weeks in ATTAIN and ACCORD I, respectively. Aclidinium also had beneficial effects on health-related quality of life and other endpoints, such as rescue medication use and rates of exacerbations. Aclidinium bromide inhalation powder is generally well tolerated in patients with COPD, with headache, cough, diarrhea, and rhinosinusitis among the most commonly reported adverse events. Cardiovascular side effects were rarely reported. Patient satisfaction studies found that patients using the aclidinium delivery device had fewer errors affecting drug delivery than those using the tiotropium device and, overall, the aclidinium device was preferred to the tiotropium device. In conclusion, aclidinium bromide is approved for use in the US at a dose of 400 µg twice daily and is a promising alternative to tiotropium.

Keywords: chronic obstructive pulmonary disease, aclidinium, antimuscarinic, anticholinergic

Introduction

Chronic obstructive pulmonary disease (COPD) comprises a group of respiratory conditions that includes emphysema and chronic bronchitis.^{1,2} Most patients with COPD exhibit features of both chronic bronchitis and emphysema; differentiation is no longer considered relevant.² COPD is characterized by chronic inflammation that causes airway remodeling and parenchymal destruction, leading to persistent airflow limitation that is usually progressive.³ The resulting features are decline of expiratory flow and inadequate lung emptying on expiration. Exacerbations and comorbidities also contribute to the severity of the disease.²

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In 2008, COPD moved from the sixth- to the third-leading cause of death in the US.¹ This increase is thought to have been driven by the aging population, smoking epidemic, and reduced mortality from other common causes of death.⁴ Data analyzed from a 2011 Behavioral Risk Factor Surveillance System survey suggest that 6.3% of US adults (an estimated 15 million) have been diagnosed with the disease.¹

The Global Initiative for Chronic Obstructive Lung Disease (GOLD) guidelines provide evidence-based recommendations for the clinical management of chronic COPD. Smoking cessation is the most important intervention and all patients who smoke should be encouraged to quit. The goals of pharmacologic treatment are to reduce symptoms and exacerbations and improve exercise tolerance and health status. Long-acting inhaled bronchodilators, which include the long-acting beta-adrenoceptor agonists salmeterol, formoterol, arformoterol, and indacaterol, and the long-acting muscarinic receptor antagonist tiotropium continue to be the mainstay of the current management of COPD. The choice of agent is patient specific and should consider availability, cost, and delivery device, including the patient's ability to effectively use the device.² Acclidinium bromide (Tudorza™ Pressair™; Almirall, SA, Barcelona, Spain) is the second long-acting inhaled antimuscarinic bronchodilator to have been approved by the US Food and Drug Administration (FDA) for maintenance treatment of COPD.⁵ This article reviews the pharmacology, pharmacokinetics, efficacy, safety, and place in therapy of acclidinium bromide.

Data selection

A search of PubMed was performed using the search terms “chronic obstructive pulmonary disease”, “acclidinium”, “anticholinergic”, and “muscarinic receptor antagonist” to identify the relevant literature. References from each identified article were reviewed for applicable articles. US government websites, including the FDA's and the ClinicalTrials.gov database, and references from identified sources were also reviewed.

Pharmacology

The superfamily of G-protein-coupled receptors includes five subtypes of the muscarinic receptors, but only M₁, M₂, and M₃ are expressed in human lung tissue. M₃ receptors are located in the smooth muscle of the airways and are primarily responsible for the bronchoconstrictor response to cholinergic nerve stimulation. Antagonism of the muscarinic receptors, particularly M₃, mediates bronchodilation and

smooth-muscle relaxation. M₂ receptors are also expressed in the heart, the antagonism of which is theorized to cause bronchoconstriction and tachycardia.^{6,7} Acclidinium bromide is a potent inhibitor of all muscarinic receptors, exhibiting kinetic selectivity for the M₃ receptor (with a 29.2-hour residence half-life) comparable to that of tiotropium. Acclidinium is unique in that it dissociates from M₂ receptors (with a 4.7-hour residence half-life) more than three times faster than tiotropium, thereby minimizing the potential for cardiovascular side effects.⁷ In animal studies, acclidinium demonstrated an onset of action twice as fast as that of tiotropium and a long duration of action similar to that of tiotropium.^{7,8}

Pharmacokinetics and pharmacodynamics

A Phase I study of 30 healthy individuals was conducted to assess the pharmacokinetics of acclidinium bromide 200 µg, 400 µg, and 800 µg twice daily (BID). Acclidinium bromide exhibited linear and time-independent pharmacokinetics. Steady state was achieved within the 7-day treatment period and was estimated to have been reached as early as the first day of treatment.⁹ The results from this study confirm findings from a previous study done with acclidinium at the same doses given once daily.¹⁰ In comparison, tiotropium requires 2–3 weeks to reach steady state.^{11,12} Acclidinium is rapidly converted to inactive alcohol and acid metabolites, resulting in low systemic exposure, which minimizes the potential for side effects.⁹ Although no formal drug interaction studies have been conducted, in vitro studies suggest low potential for drug–drug interactions. No relevant CYP450 involvement was noted in studies. Another small Phase I study of 24 participants with varying degrees of renal function demonstrated no clinically significant differences in pharmacokinetics.¹³ Acclidinium bromide has not been studied in patients with hepatic impairment.

Clinical trials

Multiple randomized controlled trials have assessed the efficacy of acclidinium in patients with COPD.^{11,14–24} Inclusion criteria for the majority of studies include patients diagnosed with moderate to severe COPD (80% > forced expiratory volume in 1 second [FEV₁] ≥30%), ≥40 years of age, with a history of cigarette smoking (≥10 pack-years). The majority of studies included similar exclusion criteria of other pulmonary disease, COPD exacerbation or respiratory-tract infection in the past 6 weeks (3 months if hospitalized), unstable

cardiac conditions, and contraindication to anticholinergic therapy. The studies allowed patients to continue on some concomitant COPD medications, including inhaled corticosteroids, systemic corticosteroids (prednisone ≤ 10 mg daily or ≤ 20 mg every other day), and methylxanthines. A summary of the efficacy trials can be found in Table 1.

An initial dose-ranging study in 17 adults compared acclidinium 100, 300, and 900 μg with placebo.¹⁴ Following acclidinium administration, investigators found statistically significant bronchodilatory effects of the drug as early as 15 minutes, peak effect by 2 hours, and mean change from baseline FEV_1 at 24 hours of +30 mL, +133 mL, +131 mL, and -28 mL for acclidinium 100, 300, 900 μg , and placebo, respectively. This 24-hour improvement compared with placebo suggested the appropriateness of once-daily dosing. A randomized, double-blind, Phase IIb study compared acclidinium at doses of 25, 50, 100, 200, and 400 μg with placebo and open-label tiotropium 18 μg daily over a 4-week period.¹⁵ The primary efficacy endpoint of trough FEV_1 of acclidinium 200 and 400 μg was shown to be statistically significantly superior to placebo at week 4 at +148 mL ($P=0.006$) and +128 mL ($P=0.018$), respectively, achieving similar results to tiotropium (+151 mL, $P=0.003$). Similar results were found in a double-blind, double-dummy, multicenter, crossover study comparing tiotropium 18 μg , placebo, and acclidinium 200 μg .¹⁶ In this study, similar increases in FEV_1 at 30 minutes and over 3 hours occurred between acclidinium and tiotropium, both of which were statistically superior to placebo. Results from these studies supported the initial dose of acclidinium 200 μg daily.

Acclidinium Clinical Trial Assessing Efficacy and Safety in Moderate to Severe COPD Patients (ACCLAIM/COPD) I and ACCLAIM/COPD II were two Phase III double-blind placebo-controlled trials performed over a 52-week period in which patients were randomized to acclidinium 200 μg or placebo daily in a patient population with an average age of 62 years and a mean baseline FEV_1 of 50%–55% predicted value.¹⁷ The primary efficacy endpoint, trough FEV_1 , was increased in the acclidinium bromide group compared with placebo by +61 mL and +63 mL at 12 weeks and +67 mL and +59 mL at 28 weeks in ACCLAIM/COPD I and ACCLAIM/COPD II, respectively ($P<0.001$ for all). This level of bronchodilation was maintained throughout the 52-week study, with both trials showing superiority over placebo at reported time points. However, these values do not meet the minimal clinically important difference (MCID) score of 100 mL for FEV_1 in COPD, suggesting suboptimal therapy.²⁵ The St George's Respiratory Questionnaire

(SGRQ), a self-administered health-status questionnaire, was also assessed and scores were significantly improved in the acclidinium groups compared with placebo in both studies. For ACCLAIM/COPD I, the mean improvement from baseline SGRQ scores was -4.63 versus -3.10 for acclidinium and placebo, respectively ($P=0.19$); similar treatment differences were noted for ACCLAIM/COPD II (-3.49 vs -1.28, $P=0.021$). These improvements in SGRQ scores were associated with statistically significantly more patients in both treatment arms achieving the MCID of -4 units on the SGRQ.²⁶ Further, statistically significantly more patients achieved the MCID for the Transitional Dyspnea Index (TDI) of 1 unit in both acclidinium groups compared with placebo.²⁷ The authors concluded that acclidinium is effective in patients with moderate to severe COPD.¹⁷ However, the low trough FEV_1 values in this study were below the MCID, and are lower than those found in studies of tiotropium, which suggests the need for higher doses or more frequent dosing.

Based on the FEV_1 results of the Phase III studies, new dose-ranging studies were completed. A Phase IIa study compared acclidinium 400 μg BID with placebo and tiotropium.¹¹ The primary efficacy endpoint of mean change in FEV_1 area under the curve ($\text{AUC}_{0-12/12\text{h}}$) on day 15 was significantly higher in the acclidinium and tiotropium groups compared with the placebo (+221 mL and +244 mL, respectively, $P<0.0001$). Similar improvements were noted in trough FEV_1 on day 15 for acclidinium and tiotropium groups versus placebo (+186 mL, +150 mL, $P<0.0001$) and in reduction of rescue medication use. On day 15, acclidinium had significantly greater improvements in $\text{FEV}_1 \text{AUC}_{12-24/24\text{h}}$ ($P<0.05$) and forced vital capacity (FVC) $\text{AUC}_{12-24/24\text{h}}$ ($P<0.05$) compared with tiotropium, suggesting better evening bronchodilation associated with the second dose of acclidinium. Supporting this finding was the significant improvements in evening symptom scores for the acclidinium group versus the placebo group, unlike for the tiotropium group versus placebo group. Results of this trial were supported by a Phase IIb study comparing acclidinium 100, 200, and 400 μg BID with placebo and formoterol 12 μg BID in patients with moderate to severe COPD over a 7-day period.¹⁸ The primary efficacy endpoint of $\text{FEV}_1 \text{AUC}_{0-12}$ on day 7 compared with placebo for acclidinium 100, 200, 400 μg , and formoterol 12 μg BID was +154 mL, +176 mL, +208 mL, and +210 mL, respectively ($P<0.0001$ for all), with acclidinium 400 μg producing statistically significant improvements compared with 100 μg ($P<0.01$). All active control medications produced significantly greater improvements in $\text{FEV}_1 \text{AUC}_{12-24}$ compared with placebo, with formoterol

Table 1 Summary of acclidinium efficacy trials

Trial	Study design	Study duration	Intervention	Outcomes
Joos et al ¹⁴	n=17 CO, DB, MC, PC, R Phase IIa	Single dose	Acclidinium 100 µg, 300 µg, 900 µg vs placebo	Significant improvements in $AUC_{0-24/24}$ for acclidinium versus placebo ($P<0.001$)
Chanez et al ¹⁵	n=464 (441 completed) AC, DB, MC, MN, PC, R Phase IIb	4 weeks	Acclidinium 25 µg, 50 µg, 100 µg, 200 µg, 400 µg daily vs placebo daily and tiotropium 18 µg daily (open label)	Significant increases in trough FEV_1 at week 4 for acclidinium 200/400 µg and tiotropium compared with placebo
Vestbo et al ¹⁶	n=115 (107 completed) CO, DB, MC, MN, R Phase II	Single dose	Acclidinium 200 µg vs placebo and tiotropium 18 µg	Significant improvements in percentage of patients with FEV_1 increase $\geq 10\%$ at 30 minutes compared with placebo ($P<0.0001$ for both medications)
Jones et al ¹⁷ ACCLAIM/COPD I	n=843 (707 completed) R, DB, PC, MC, MN Phase III	52 weeks	Acclidinium 200 µg daily vs placebo	Significant improvements in mean change in FEV_1 at 30 minutes: +122 mL acclidinium, +118 mL tiotropium, +27 mL placebo ($P<0.0001$ for both compared with placebo) Significant improvements in FVC, IC, and perception of dyspnea Significant increase in trough FEV_1 at 12/28 weeks Range +37–67 mL throughout the study Significantly more acclidinium patients achieved the MCID for SGRQ at week 52; mean difference in SGRQ at 52 weeks –4.63 acclidinium, –3.1 placebo ($P=0.19$) Significantly more acclidinium patients achieved the MCID for TDI at week 52; mean difference in TDI at 52 weeks +1.83 acclidinium, +0.61 placebo ($P<0.0001$) Significant increase in trough FEV_1 at 12/28 weeks Range +51–78 mL throughout the study Significantly more acclidinium patients achieved the MCID for SGRQ at week 52; mean difference in SGRQ at 52 weeks –3.49 acclidinium, –1.28 placebo ($P=0.021$) Nonsignificantly more acclidinium patients achieved the MCID for TDI at week 52; mean difference in TDI at 52 weeks +1.61 acclidinium, +1.09 placebo ($P=0.12$) Fewer patients in acclidinium group compared with placebo group experienced a moderate to severe COPD exacerbation (33.2% vs 39.8%, $P=0.0046$)
Jones et al ¹⁷ ACCLAIM/COPD II	n=804 (564 completed) R, DB, PC, MC, MN Phase III	52 weeks	Acclidinium 200 µg daily vs placebo daily	Mean change in FEV_1 $AUC_{0-12/12h}$ was significantly higher in the acclidinium and tiotropium groups than placebo ($P<0.0001$) Significant differences between acclidinium and tiotropium existed for FEV_1 $AUC_{12-24/24h}$ and FVC $AUC_{12-24/24h}$ COPD symptoms significantly improved with acclidinium but not with tiotropium compared with placebo Significant improvements in FEV_1 AUC_{0-12} in both active medication groups compared with placebo ($P<0.0001$)
Fuhr et al ¹¹	n=30 (27 completed) AC, CO, DB, MC, PC, R Phase IIa	15 days	Acclidinium 400 µg BID vs placebo and tiotropium 18 µg daily	
Singh et al ¹⁸	n=79 (68 completed) AC, CO, DB, MC, MN, PC, R Phase IIb	7 days	Acclidinium 100 µg, 200 µg, 400 µg BID vs placebo and formoterol 12 µg BID	

Jones et al ¹⁹ ATTAIN	n=828 (737 completed) DB, MC, MN, PC, R Phase III	24 weeks	Acclidinium 200 µg, 400 µg BID vs placebo	Significant improvements in trough FEV ₁ compared with placebo at week 24; +99 mL acclidinium 200 µg, +128 mL acclidinium 400 µg ($P<0.0001$ for both) Significant improvements in SGRQ compared with placebo at week 24; -3.8 units acclidinium 200 µg ($P<0.001$), -4.6 units acclidinium 400 µg ($P<0.0001$) Significant improvements in TDI at week 24 compared with placebo: +0.6 units acclidinium 200 µg ($P<0.05$), +1 unit acclidinium 400 µg ($P<0.001$) Statistically significant improvements in trough FEV ₁ at week 12 for acclidinium 200 µg (+86 mL, $P<0.0001$). Clinically and statistically significant improvements in FEV ₁ at week 12 for acclidinium 400 µg (+124 mL, $P<0.0001$) Significant improvements in TDI, SGRQ, and symptom scores at week 12 Clinically and statistically significant improvements in SGRQ from baseline in all groups Improvements in trough and peak FEV ₁ Statistically but not clinically significant improvements in FEV ₁ at week 12: +51 mL acclidinium 200 µg, +72 mL acclidinium 400 µg Numerically higher results in SGRQ from baseline in both groups compared with placebo Statistically significant improvements in TDI from baseline Statistically significant improvements in FEV ₁ , AUC ₀₋₂₄ at week 6 in both active treatments compared with placebo Statistically significant improvements in E-RS compared with placebo in both active treatment arms Numerical improvements in trough FEV ₁ compared with baseline Clinically significant improvements in SGRQ compared with baseline
Kerwin et al ²⁰ ACCORD I	n=561 (467 completed) DB, MC, MN, PC, R Phase III	12 weeks	Acclidinium 200 µg, 400 µg BID vs placebo	
D'Urzo et al ²¹	n=291 (199 completed) DB, MC, MN, R Phase III extension	52 weeks	Acclidinium 200 µg, 400 µg BID	
Rennard et al ²² ACCORD II	n=544 (454 completed) DB, PC, MC, MN, R Phase III	12 week	Acclidinium 200 µg, 400 µg BID vs placebo	
Beier et al ²³	n=414 (400 completed) Phase IIb AC, DB, MC, MN, PC, R	6 weeks	Acclidinium 400 µg BID vs placebo and Tiotropium 18 µg daily	
Gelb et al ²⁴	n=605 (341 completed) DB, R	52 weeks	Acclidinium 200 µg, 400 µg BID	

Abbreviations: AC, active comparator; ACCCLAIM/COPD, Acclidinium Clinical Trial Assessing Efficacy and Safety in Moderate to Severe COPD Patients; ACCORD, Acclidinium in Chronic Obstructive Respiratory Disease; ATTAIN, Acclidinium to Treat Airway Obstruction in COPD Patients; BID, twice daily; CO, crossover; DB, double blind; E-RS, EXACT (exacerbations of chronic pulmonary disease tool)-Respiratory Symptoms; FEV₁, forced expiratory volume in 1 second; FVC, forced vital capacity; IC, inspiratory capacity; MC, multicenter; MCI, minimal clinically important difference; MN, multinational; O, open label; PC, placebo controlled; R, randomized; SGRQ, St George's Respiratory Questionnaire; TDI, Transitional Dyspnea Index.

+56 mL compared with acclidinium 400 µg BID ($P<0.01$). Trough FEV₁ on day 7 was significantly greater for all active controller medications compared with placebo, with improvements of +106 mL, +114 mL, +154 mL, and +148 mL for acclidinium 100, 200, 400, and formoterol 12 µg BID, respectively. Results of these two studies suggested that doses of 200 and 400 µg BID were appropriate for Phase III evaluation.^{11,18}

Based on the Phase IIa/b data, a Phase III randomized, double-blind, placebo-controlled trial comparing acclidinium 200 and 400 µg BID compared with placebo in moderate to severe COPD was undertaken.¹⁹ The Acclidinium to Treat Airway Obstruction in COPD Patients (ATTAIN) study consisted of primarily male subjects with an average age of 62.4±8 years, an average smoking history of 40.2±19.8 pack-years, 52.8% of whom were current smokers, and an average baseline FEV₁% predicted of 52.5±14.1%. Patients were well matched at baseline overall, with a SGRQ score approximately 46.3±16.8 units and a baseline dyspnea index (BDI) of 6.8±2.1 units. In terms of the primary efficacy endpoint, both acclidinium bromide 200 and 400 µg significantly improved trough FEV₁ compared with baseline at 24 weeks versus placebo (+99±22 mL and +128±22 mL, respectively, $P<0.001$ for both). Of the two doses, only one met the MCID for this efficacy outcome, suggesting clinically significant improvement for only acclidinium 400 µg. This improvement in FEV₁ with acclidinium 400 µg was maintained throughout the study time points, with a range of +105 to 140 mL, maintaining clinical significance. Peak FEV₁ was significantly improved at all time points for both doses, with similar FEV₁ results at weeks 12 and 24. Both doses produced statistically significant improvements in FVC and inspiratory capacity (IC) compared with placebo at all prespecified time points. Both doses of acclidinium provided statistically significant improvements in health status over the 24-week period, with a mean change of -3.8±1.1 versus -4.6±1.1 units in SGRQ scores for acclidinium 200 and 400 µg, respectively, which met the MCID. A lower proportion of patients met the MCID in the placebo compared with the acclidinium 200 and 400 µg groups (41%, 56%, and 57.3%, respectively, $P<0.001$ for acclidinium compared with placebo). Similar to the results for the SGRQ, both acclidinium 200 and 400 µg improved TDI scores by +0.6±0.3 units ($P<0.05$) and +1±0.3 units ($P<0.001$), respectively, with the 400 µg dose meeting MCID for clinical significance. This increase in health status was paralleled by a significant reduction in the use of rescue medication for both doses of acclidinium compared with placebo. Further, reductions in the rate ratio of COPD exacerbations

of any severity was significantly lower with acclidinium 200 (rate ratio =0.72, 95% confidence interval [CI] =0.52–0.99) and 400 µg (rate ratio =0.67, 95% CI =0.48–0.94) compared with placebo, despite the study not being powered specifically for exacerbations.

Similar to the ATTAIN study, the Acclidinium in Chronic Obstructive Respiratory Disease (ACCORD) I study, a Phase III, randomized, double-blind, placebo-controlled trial over 12 weeks, compared acclidinium 200 and 400 µg BID with placebo in patients with moderate to severe COPD.²⁰ The subjects had an average age of 64.2±9.4 years, 53% were male, and 44.8% were current smokers with a smoking history of 54.3±26.8 pack-years. Baseline FEV₁ was 47.2%±14.1% predicted, an average baseline SGRQ of 46.5±17.1 units and BDI of 6.4±2.1 units. For the primary efficacy outcomes, acclidinium 200 µg statistically significantly increased trough FEV₁ at 12 weeks (+86 mL, $P<0.0001$), while acclidinium 400 µg both clinically and statistically significantly increased trough FEV₁ (+124 mL, $P<0.0001$). In addition, both doses were associated with significant improvements in peak FEV₁ at 12 weeks (+146 mL, +192 mL, $P<0.0001$ for both). Both IC and mean FVC AUC_{0-3h} statistically significantly improved from baseline compared with placebo at the end of 12 weeks. In the first week, maximal bronchodilation was achieved and maintained throughout the study. Mean improvements in SGRQ scores compared with placebo at 12 weeks were -2.7 units in acclidinium 200 µg ($P=0.013$) and -2.5 units in acclidinium 400 µg ($P=0.019$), meeting statistical but not clinical significance. Similarly, statistically significant improvements were found for both doses at week 12 in the TDI score, with acclidinium 400 µg achieving clinical significance. Non-validated questionnaires developed by the sponsor specifically for this study were utilized to assess changes in symptoms and rescue medication use. Compared to placebo, acclidinium 200 and 400 µg produced significant improvements in daily symptoms at 12 weeks, with acclidinium 400 µg numerically higher than 200 µg. Acclidinium 200 µg did not meet statistical significance in improvement of nighttime breathing and nighttime sputum production, where the 400 µg dose also did not meet statistical significance in improvement in sputum production. Both doses of acclidinium were associated with significant reductions in rescue medication use at the end of the 12-week period. Both doses of acclidinium were associated with significant reductions in rescue medication use at the end of the 12-week period. Similar to the ATTAIN study, a reduction in rates of COPD exacerbations of any kind with acclidinium 200 and 400 µg compared with placebo was demonstrated, with 0.55, 0.41, and 0.79 exacerbations,

respectively, resultant per patient year. Only the comparison of the 400 µg dose with placebo demonstrated a significant reduction in rates of exacerbations of any kind (rate ratio =0.52, $P=0.009$).

In a 1-year extension of the ACCORD I study, the patients assigned originally to placebo were randomized (1:1) to either acclidinium 200 or 400 µg and active medication patients continued their current dose of acclidinium for 52 weeks.²¹ At the end of the treatment period, patients on continuous acclidinium 200 and 400 µg had a trough FEV₁ change from baseline of +69 mL and +56 mL, respectively. For patients re-randomized from placebo to acclidinium, improvements were seen in FEV₁ similar to those seen in the initial ACCORD I study, although results were not reported. Clinically and statistically significant improvements in SGRQ scores occurred at 52 weeks in the continuous acclidinium group, with a mean improvement of -7.0 units in 200 µg and -7.9 units in the 400 µg group. In patients changed from placebo to acclidinium, clinically and statistically significant improvements in SGRQ scores also occurred, with improvements of -4.9 units in the 200 µg group and -5.7 units in the 400 µg group. All groups also experienced reduced use of rescue medication for the treatment of COPD symptoms.

A third Phase III randomized, double-blind, placebo-controlled trial in patients with moderate to severe COPD comparing acclidinium 200 and 400 µg with placebo over a 12-week period, ACCORD II, was undertaken to confirm results of previous trials.²² This study was performed in patients with an average age of 62.8±8.9 years and 53.1% of subjects were male. Approximately 53.3% of patients were current smokers, with a 53.8±29.1 pack-year history. Baseline pulmonary function showed an FEV₁ of 52.5%±13.4% predicted, a baseline SGRQ score of 49±17.6 units, and TDI of 6.2±2.1 units. Two patients in the placebo group had mild COPD and one patient in the acclidinium 200 µg group had very severe COPD discovered after randomization. Additionally, an uneven dispersion of patients with moderate and severe COPD existed between the placebo and both acclidinium groups, with more moderate COPD favored in the placebo group and more severe COPD in the acclidinium 400 µg group. The primary efficacy endpoint of trough FEV₁ at 12 weeks was statistically but not clinically significantly superior to placebo, with an improvement of +51 mL and +72 mL for acclidinium 200 and 400 µg BID, respectively ($P<0.05$ for both). Patients also showed improvements in peak FEV₁, IC, and FVC from baseline in both acclidinium groups compared with placebo. Improvements in SGRQ scores were seen in all three groups, with

a score of -5.4, -6, and -4.3 units for acclidinium 400 µg, 200 µg, and placebo, respectively. TDI improved with acclidinium 200 and 400 µg by +1.0 and +1.3 units from baseline compared with +0.3 units with placebo ($P<0.05$ compared with placebo for both acclidinium groups). This minimal improvement in outcomes may be attributed to the uneven patient dispersion among COPD severity groups favoring the placebo group and thereby minimizing the treatment effect. Numerically, less use of rescue medication occurred in the acclidinium groups compared with in the placebo group.

Patients with moderate to severe COPD were assessed in a Phase IIIb trial comparing acclidinium 400 µg BID, tiotropium 18 µg daily, and placebo.²³ This study included patients with an average age of ~62 years and post-bronchodilator FEV₁ of 55%–56% predicted, with ~53% current smokers with a 39.5–45.0 pack-year history of smoking depending on group. The primary efficacy outcome of FEV₁ AUC_{0–24} at week 6 in both acclidinium and tiotropium groups significantly improved compared with placebo (+150 mL, +140 mL, respectively, $P<0.0001$ for both). At week 6, numerically greater change in FEV₁ AUC_{12–24} in acclidinium (+160 mL) compared with tiotropium (+123 mL), similar to previous Phase II studies demonstrating superior bronchodilatory activity of acclidinium in the evening was observed.¹¹ Tiotropium had numerically greater FEV₁ AUC_{0–12} compared with acclidinium, although this did not meet statistical significance.²³ Symptoms were assessed using the Exacerbations of Chronic Pulmonary Disease Tool (EXACT), a 14-item instrument. Responses for 11 of the 14 questions were used to calculate another score called the “EXACT-Respiratory Symptoms” (E-RS) score, in which higher numbers are associated with increased symptoms on a scale from 0 to 40. At week 6, a significant improvement in E-RS score occurred with both tiotropium and acclidinium compared with placebo, $P<0.05$ and $P<0.0001$, respectively. Similar to previous Phase II studies, and in accordance with the FEV₁ AUC_{12–24}, acclidinium, but not tiotropium, led to a significant reduction in nighttime symptom severity compared with placebo.¹¹ However, differences in symptom scores between active treatment groups were not significant.²³

A 52-week study in patients with moderate to severe COPD compared acclidinium 200 and 400 µg BID in patients with a mean age of 63.6 years and mean post-bronchodilator FEV₁ of 52.3% predicted.²⁴ In this study, mean improvements in trough FEV₁ at 52 weeks of +34 mL and +72 mL in the 200 and 400 µg acclidinium groups, respectively, were observed. At week 52, mean improvements in SGRQ scores were clinically significant, with -5.3 units and -5.2 units

in the 200 and 400 µg groups, respectively, compared with baseline. Although no clinically significant improvements in FEV₁ were reached at week 52, the numerical improvement along with the results from the previous Phase III trials support the FDA-approved dose of acclidinium 400 µg BID.

Safety

Acclidinium studies with doses of up to 6,000 µg have shown that acclidinium is safe.¹⁰ Further studies of up to 900 µg inhaled have demonstrated that there are minimal amounts of the medication in the blood after administration.¹⁴ After doses of 100, 300, and 900 µg, the amount of active medication and alcohol-based metabolite were both below the lower limit of quantification of <0.1 ng/mL in the blood at all time points.¹⁴ The inactive acid metabolite was detected at a few time points after the 300 µg dose, and, after the 900 µg dose, the maximum concentration was detected at 3 hours after dosing with a concentration between 3 and 6 ng/mL.¹⁴ This promotes a reduction in rates of both adverse events and the potential for drug interactions. As a long-acting muscarinic antagonist, acclidinium has the propensity for side effects, including dry mouth, constipation, and increased risk of urinary-tract infections. Of the clinical studies available, few of the reported anticholinergic adverse events have been directly associated with acclidinium bromide. The most common side effects of acclidinium include headache, nasopharyngitis, cough, and diarrhea.⁵

Muscarinic antagonist activity at the M₂ receptor has been associated with the potential for cardiovascular adverse events associated with muscarinic antagonists. Results of a meta-analysis performed by Singh et al raised concerns for adverse cardiovascular events associated with tiotropium use.²⁸ Data from the Understanding Potential Long-term Impacts on Function with Tiotropium (UPLIFT) trial, which compared tiotropium 18 µg daily with placebo over a 4-year treatment period, found no significant difference in mortality between tiotropium and placebo in the intention-to-treat analysis (hazard ratio [HR] =0.89, 95% CI =0.79–1.02).²⁹ In 67 patients compared with 85 patients in the tiotropium and placebo groups, respectively, myocardial infarction occurred (relative risk [RR] =0.71, 95% CI 0.52–0.99). Similarly, stroke developed in 80 placebo patients and 82 tiotropium patients (RR =0.95; 95% CI =0.70–1.29).²⁹ Based on the currently available information, the FDA does not presently support an increased risk of stroke, myocardial infarction, or death associated with tiotropium delivered by the HandiHaler® (Spiriva® HandiHaler,

Boehringer Ingelheim Pharmaceuticals, Inc., Ridgefield, CT, USA) device.³⁰ As a result of these concerns, acclidinium studies have included cardiovascular safety assessments, although the drug's reduced residence time at M₂ receptors and more rapid plasma hydrolysis compared with tiotropium is thought to contribute to its positive cardiovascular safety profile.⁷ This is supported by the lack of significant electrocardiography changes in clinical trials and a specific safety study focusing on the effect of acclidinium on corrected QT interval (QTc). This study was a Phase I trial completed in 272 healthy patients who were randomized to placebo, acclidinium 200 µg, or acclidinium 800 µg once daily for 3 days. Patients were also given open-label moxifloxacin 400 mg as an active control for the treatment period. All patient groups had similar values for QT interval corrected for individual heart rate (QTcI) at baseline, with no significant changes between either acclidinium dose or placebo noted at any time period. Moxifloxacin, however, demonstrated significant increase in QTcI compared with placebo at all time points.³¹ Overall, cardiac-related adverse events were rare in the studies and often unrelated to the active medication.

For example, in the ATTAIN study, two incidents of cardiovascular adverse events – a myocardial infarction in a patient on 200 µg acclidinium and an acute cardiac failure in a patient on acclidinium 400 µg – occurred, both of which were not thought to be treatment related.¹⁹ Few adverse events occurred in the ACCORD COPD II study, five of which were cardiovascular mediated. Of these five events, four occurred in patients on placebo therapy, and one incident of tachycardia occurred in a patient on acclidinium 400 µg BID.²²

In the ACCORD COPD I extension study, patients were re-randomized in a 1:1 ratio to placebo: acclidinium 200 or 400 µg BID, or maintained on their original dose of acclidinium 200 or 400 µg BID for a 52-week period. Treatment-emergent adverse events (TEAEs) were similar across the treatment arms, with 77.4% in the acclidinium 200 µg and 73.7% in the acclidinium 400 µg group experiencing at least one TEAE, the most common of which were acute exacerbations of adverse events. The TEAE was determined by investigators to be attributable to acclidinium 200 µg and 400 µg 10.2% and 12.5% of the time, respectively. Anticholinergic side effects were uncommon, with urinary-tract infections occurring in 3.6% and 5.9% of patients taking acclidinium 200 µg and 400 µg, respectively. Similarly, constipation occurred in 2.9% and 1.3% in acclidinium 200 and 400 µg BID, respectively. Dry mouth associated with acclidinium only occurred in one patient in the 400 µg group. Of the patients who experienced cardiac adverse events, only three of these

events were considered treatment related: one patient in the 200 µg group experienced acute coronary syndrome and atrial fibrillation – prior to study initiation, this patient had a cardiovascular history significant for coronary artery bypass and coronary angioplasty; one patient on acclidinium 400 µg with a medical history significant for cardiac disorders had congestive heart failure; and a third patient experienced first-degree atrial-ventricular blockade in the acclidinium 400 µg group.²¹ Results of this study are similar to those of another 52-week study comparing acclidinium 200 and 400 µg BID as well as those of other Phase III trials.

Patient satisfaction

Rates of medication nonadherence have approached 50% for chronic disease states in developed countries.³² This has important implications for COPD patients, as medication nonadherence has been associated with increased mortality, health-care cost, and hospitalizations.^{33,34} Various factors have been associated with patient nonadherence, including patient-based, societal, and treatment-related factors, which can affect adherence and thus the underlying disease state. Treatment-related factors such as method of administration, ease of use, side-effect profile, and accessibility can affect patient satisfaction and, therefore, patient adherence and the proper treatment of COPD. Acclidinium bromide has been formulated in a unique breath-actuated, multi-dose dry-powder inhaler (Tudorza™ Pressair™) with multiple patient feedback mechanisms to promote proper use of the inhaler. These feedback mechanisms include both sound and color feedback. When activating the inhaler, a color window at the front of the inhaler turns from red to green indicating the inhaler is ready to use; once a dose has been properly inhaled, the color turns back to red. Similarly, during inhalation, when a dose is properly taken, the patient should hear a click to indicate appropriate use.⁵

Beier et al sought to assess inhaler preference in 414 patients through a double-blind, double-dummy, randomized, multicenter, placebo- and active-controlled study comparing acclidinium 400 µg BID to tiotropium 18 µg daily over a 6-week period.²³ At the end of the 6-week period, overall willingness to continue on either the acclidinium Genuair™ (Almirall SA) inhaler or tiotropium HandiHaler® inhaler was assessed based on a 0–100 scale, with higher numbers being associated with increased willingness to continue on the inhaler; overall inhaler preference was also assessed. Additionally, patient preference for either the Genuair or HandiHaler, based on specific attributes such as ease of use, ease of learning to use, operation, holding, dose

preparation, and inhaler feedback indicating proper use, was analyzed. More patients at the end of the 6-week period preferred Genuair to HandiHaler, with few patients having no preference for either inhaler (80.1%, 10.7%, and 9.2%, respectively, $P<0.0001$). When assessing inhaler-based specific factors between Genuair and HandiHaler, Genuair was preferred in each category, as can be seen in Table 2. When assessing patient willingness to continue with a device, more patients were willing to continue with the Genuair than the HandiHaler, as indicated by a higher mean rating in the preference scale at week 6 (88.8 vs 45.4, $P<0.0001$).²³

Beier et al's results are supported by an open-label, randomized, crossover study by van der Palen et al comparing patient preference and satisfaction between the acclidinium Genuair/Pressair and tiotropium HandiHaler in patients with moderate to severe COPD.³⁵ In this 2-week, placebo-only trial, 105 patients were assessed on their satisfaction with the inhaler in terms of eight inhaler-related attributes, willingness to continue with a particular inhaler, and the number of errors related to inhaler use. Overall, patients demonstrated a higher satisfaction with Genuair than with HandiHaler (79.1% vs 20.9%, $P<0.0001$). In terms of the specific inhaler attributes, significantly more patients were satisfied with and preferred Genuair to HandiHaler, as can be seen in Table 2. Although still significantly preferred compared with HandiHaler, fewer patients demonstrated satisfaction with the feedback to indicate correct dose inhalation of Genuair compared with satisfaction in other categories. When comparing the number of errors between the two inhaler types, seven patients experienced an error with Genuair compared with 19 patients with HandiHaler (6.7% vs 18.1%, $P<0.01$). When comparing patients who had at least one “critical” error, defined as an error affecting drug delivery, three patients had at least one with Genuair compared with 20 patients with HandiHaler (2.9% vs 19%, $P<0.0001$), excluding those patients who made errors with both inhaler types. The majority of errors with Genuair occurred during dose inhalation, with common errors being failure to hold the inhaler upright and to hold their breath for several seconds after inhalation. In terms of willingness to continue using an inhaler, as assessed on a 0–100 scale on which higher numbers indicated willingness to continue, the mean scores for Genuair were higher (84.0 vs 62.5, $P<0.0001$).

An advantage of van der Palen et al's study was the use of placebo instead of active medication, which thereby limited the effect of perceived effectiveness of therapy on satisfaction results.³⁵ This is especially important in a study comparing tiotropium and acclidinium due to the longer time

Table 2 Patient satisfaction with Pressair™/Genuair™ versus HandiHaler devices

Parameter	Beier et al, ²³ n=414 Patient preference (%)		Van der Palen et al, ³⁵ n=105 Patient preference (%)	
	Pressair/Genuair	HandiHaler	Pressair/Genuair	HandiHaler
Ease of use	86.7*	9.0	83.9*	16.1
Convenience	87.4*	7.8	79.8*	20.2
Ease of learning	79.1*	8.0	82.4*	17.6
Ease of holding	79.9*	8.5	85.4*	14.6
Ease of operation	81.1*	9.2	84.5*	15.5
Ease of dose preparation	86.4*	7.8	87.6*	12.4
User feedback	76.2*	9.5	69.2**	30.8

Notes: * $P < 0.0001$; ** $P < 0.01$ compared with HandiHaler.

to steady state for tiotropium therapy. The authors concluded that the Genuair inhaler is preferred to HandiHaler by patients with moderate to severe COPD. The limitations of this trial included its small population; short duration; and exclusion of patients who would have difficulty with proper inhaler use, including patients with body malformation, unable to read, and/or with disease states that affect coordination. These exclusion criteria greatly limit the applicability of the results to patients who would have more difficulty in using inhalers properly at baseline; as such, further studies in this area are needed.³⁵

Similar results were found in a Phase IIb trial comparing aclidinium 100, 200, and 400 µg BID to placebo and formoterol 12 µg BID. When patients' preference for the Genuair versus Foradil Aerolizer® (Novartis AG, Basel, Switzerland) devices was assessed, more patients demonstrated preference for the Genuair (75.6% vs 10.3%) and found the inhaler easier to use.¹⁸

Conclusion

GOLD guidelines recommend long-acting inhaled bronchodilators for the first-line treatment of COPD. Until recently, tiotropium was the only long-acting muscarinic receptor antagonist available for use in the US. Although limited head-to-head trials comparing the efficacy and safety of tiotropium and aclidinium have been conducted, placebo-controlled trials show they are comparable in terms of improving lung function and health status in patients with moderate to severe COPD. Clinical studies suggest a good safety profile and low incidence of both anticholinergic side effects and medication-related cardiovascular events. The adverse events most commonly reported with aclidinium use were headache, cough, diarrhea, and rhinosinusitis. Cardiovascular side effects were rarely reported in two extension trials and one dedicated safety study (all of 52 weeks' duration). Post-marketing studies will be conducted to further determine the risk of potential cardiovascular side effects.

The choice of agent should also consider patient preference for dosing schedule and administration. Once-daily dosing would be expected to be the preferred option, however, studies that examined patient satisfaction found the aclidinium delivery device was preferred to the tiotropium device. Aclidinium's feedback mechanisms appear to reinforce proper administration with fewer administration errors compared with the tiotropium inhaler.

Aclidinium bromide, a long-acting muscarinic antagonist, is approved for use in the US at a dose of 400 µg BID for the maintenance treatment of COPD. It is a promising alternative to tiotropium, having demonstrated similar efficacy and safety in several clinical trials. Future studies will likely include products that combine muscarinic receptor antagonists and long-acting beta-adrenoceptor agonists in one device.

Disclosure

The authors report no conflicts of interest in this work.

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