Tiotropium in asthma: a systematic review

Introduction: The objective of this paper is to systematically review the existing evidence of the effectiveness and safety profile of a long-acting inhaled muscarinic antagonist as add-on therapy in patients with asthma that is uncontrolled despite inhaled corticosteroid (ICS) use.

Methods: With the assistance of two experienced research librarians, we searched Ovid MEDLINE/PubMed (1946 to September 12, 2013), the Cochrane Library review, and the TRIP database. The key search terms were “tiotropium and asthma.” The search was limited to human data published in English. Included in the systematic review were all randomized controlled trials that evaluated the efficacy of tiotropium in patients with asthma. The clinical trials had to be at least 4 weeks in duration and to provide adequate information on clinically appropriate end points in asthma care (eg, change in lung function, exacerbation rates, and/or ICS dosing). Data on patient characteristics, study design, outcome measures, concomitant asthma medication, and adverse events were extracted from the full text of each included individual study. Marked heterogeneity of study design precluded statistical pooling of results for a meta-analysis. Consequently, only descriptive summaries of outcomes are provided.

Results: Our database search retrieved 149 citations. We found five randomized controlled trials in humans that met our criteria for inclusion in the systematic review. We also found two open-label uncontrolled trials that were considered in the discussion. Each of the five included studies met the Consolidated Standards of Reporting Trials criteria for a well-designed randomized trial.

Discussion: The five clinical studies included in this systematic review focused on evaluating the efficacy of tiotropium as add-on therapy to ICS or ICS in combination with a long-acting inhaled β2-agonist (LABA) in patients with uncontrolled moderate to severe persistent asthma. Tiotropium maintained lung function when ICSs were tapered and when an LABA was discontinued. Tiotropium improved lung function when added to ICS alone or ICS–LABA combination therapy. In the only trial to have compared the addition of tiotropium with doubling the dose of ICS, tiotropium provided significantly superior results. In trials in which the addition of tiotropium was compared with salmeterol, the beneficial effects of these two bronchodilators were similar. No safety concerns were found with use of tiotropium as add-on therapy.

Conclusion: Tiotropium may have a beneficial role in moderate to severe persistent asthma despite use of an ICS or ICS and LABA. Use of tiotropium as add-on therapy poses no safety concerns.

Keywords: tiotropium, asthma, lung, inflammation, inhaled corticosteroid, LABA, LAMA

Introduction

Asthma is a chronic inflammatory airway disease. Airway inflammation in asthma is characterized by bronchial wall infiltration by eosinophils, activated mast cells, and T lymphocytes. Cytokines from T helper 2 cells and leukotrienes play key roles in mediating airway inflammation. Also seen are goblet cell hyperplasia, mucus gland hypertrophy, and airway mucus accumulation. Increased airway smooth
Muscle mass and reversible bronchoconstriction are other important features of asthma. Airway inflammation, mucus accumulation, and bronchospasm account for the symptoms of shortness of breath, wheezing, cough, and chest tightness in asthma. Understanding asthma pathophysiology is an important issue for health care practitioners in the US because asthma prevalence increased from 7.3% in 2001 to 8.4% in 2010. In 2010, an estimated 25.7 million Americans had asthma – 18.7 million adults aged 18 years and over and 7.0 million children aged 0–17 years.

The ultimate goal of asthma management is to minimize symptoms, prevent acute exacerbations, and avoid adverse medication effects. The pathophysiologic features of asthma explain why current US and international guidelines recommend the use of an inhaled corticosteroid (ICS) and a short-acting inhaled β₂-agonist (SABA) for treatment of persistent asthma. ICSs can control airway inflammation, and SABAs can manage acute symptoms related to bronchospasm. To reduce the risk of drug-related side effects, current asthma management guidelines recommend a stepwise treatment approach to achieving optimal asthma control, with more aggressive ICS dosing reserved for patients with more severe disease. However, prospective studies have confirmed that a significant proportion of asthma patients do not achieve asthma control, even with maximal approved doses of ICS. In patients who do not respond to ICS treatment alone, asthma guidelines recommend addition of another long-term controller, such as a long-acting inhaled β₂-agonist (LABA), a leukotriene-modifying agent (LMA), theophylline, and/or an anti-immunoglobulin E monoclonal antibody. Despite the addition of agents like LABAs and LMAs to ICSs, many asthma patients still remain both symptomatic and obstructed. There is clearly a need for alternative medications with different modes of action in asthma management. One intriguing option as an add-on therapy for asthma patients not responding to ICSs is an anticholinergic.

Tiotropium bromide is a long-acting inhaled muscarinic antagonist (LAMA). It has high potency as a selective antagonist at the muscarinic acetylcholine (Ach) receptors. Tiotropium rapidly dissociates from M2 Ach receptors but slowly dissociates from M3 Ach receptors. Consequently, tiotropium provides 24-hour bronchodilation as maintenance therapy in moderate to severe chronic obstructive pulmonary disease (COPD). Tiotropium has also been shown to reduce goblet cell metaplasia and airway smooth muscle mass in allergen-challenged mice and guinea pigs. Early evidence of clinical benefit with LAMA therapy in asthma was identified in COPD patients with airway hyper-responsiveness and concomitant asthma.

The objective of this paper is to systematically review the existing evidence of the use of tiotropium in patients with uncontrolled asthma and to provide recommendations for its potential role in the treatment of asthma. We sought to answer the question what is the existing evidence of the effectiveness and safety profile of an LAMA as add-on therapy in patients with asthma that is uncontrolled despite ICS use?

Materials and methods

Search strategy and study selection

We adhered to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) criteria when performing this study. With the assistance of two experienced research librarians, we searched Ovid MEDLINE/PubMed (1946 to September 12, 2013), the Cochrane Library review, and the TRIP database. The key search terms were “tiotropium and asthma.” The search was limited to human data published in English.

Abstracts of articles identified in the search were reviewed independently by two of the authors. Articles meeting pre-identified inclusion and exclusion criteria were selected for evaluation in this systematic review. Disagreements about inclusion of individual articles were resolved by consensus or review by a third author. Included in the systematic review were all randomized controlled trials that evaluated the efficacy of tiotropium in patients with asthma. The clinical trials had to be at least 4 weeks in duration and to provide adequate information on clinically appropriate end points in asthma care (eg, change in lung function, exacerbation rates, and/or ICS dosing). Excluded were unpublished and ongoing research, animal studies, reviews, case reports, case series, editorials, and studies in COPD patients (with or without asthma). The references of published articles identified in the search were examined for additional studies appropriate for inclusion in the review.

Data extraction and quality assessment

Data on patient characteristics, study design, outcome measures, concomitant asthma medication, and adverse events were extracted from the full text of each included individual study. The methodologic quality of the trials was evaluated using criteria reported in the Consolidated Standards of Reporting Trials (CONSORT) Statement. Marked heterogeneity of study design precluded statistical pooling of results for a meta-analysis. Consequently, only descriptive summaries of outcomes are provided.
Results

Our database search retrieved 149 citations of which 93 were excluded because they did not meet the inclusion criterion of being specific to the effect of tiotropium in asthma (Figure 1). An additional 37 citations were excluded because they did not describe results of clinical trials in humans. Twelve other citations were not included. Two were single-patient case reports, one was in Japanese, six had duplicate information, and three studies did not make a distinction between asthma and COPD.

We found five randomized controlled trials in humans that met our criteria for inclusion in the systematic review. One open-label study without a control was identified in the database search. This study was not included in the systematic review but was considered in the discussion. Following review of the references of the selected papers, an additional open-label trial without a control group was identified. This study was not included in the systematic review but was considered in the discussion. Summary descriptions of the five publications that met inclusion criteria are provided in Tables 1 and 2. Detailed descriptions of the five selected studies are provided as follows.

Fardon et al evaluated the ICS sparing ability of tiotropium in patients with severe persistent asthma.
<table>
<thead>
<tr>
<th>Study</th>
<th>Inclusion criteria</th>
<th>Design</th>
<th>Duration</th>
<th>Size</th>
<th>Intervention</th>
<th>Overall conclusion</th>
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<tr>
<td>Bateman et al</td>
<td>Inadequately controlled moderate persistent asthma: B16-Arg/Arg patients FEV&lt;sub&gt;1&lt;/sub&gt; &lt; 90% for patients on ICS-LABA or &lt; 80% for patients on ICS Reversibility (&gt; 12% and 200 mL ≤ 10 pack-year smoking)</td>
<td>Randomized, double-blind, double-dummy, placebo-controlled, parallel-group</td>
<td>16 weeks</td>
<td>388</td>
<td>Initial 4 weeks: usual ICS dose (400–1,000 μg daily) and 50 μg salmeterol twice daily (run-in) Subsequent 16 weeks: 5 μg tiotropium daily in evening 50 μg salmeterol twice a day</td>
<td>Tiotropium superior to placebo and noninferior to salmeterol in moderate persistent asthma with B16-Arg/Arg genotype</td>
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<tr>
<td>Fardon et al</td>
<td>Severe persistent asthma: FEV&lt;sub&gt;1&lt;/sub&gt; &lt; 65% predicted FVC &lt; 80% predicted Reversibility &gt; 5% Nonsmokers</td>
<td>Randomized, double-blind, placebo-controlled, crossover</td>
<td>4 weeks</td>
<td>18</td>
<td>Initial 4 weeks: fluticasone 1,000 mcg daily (run-in) Subsequent 4-week crossover periods: Fluticasone 500 μg all patients Salmeterol and placebo Salmeterol and tiotropium</td>
<td>Tiotropium may offer small benefit when added to salmeterol in stepping down ICS with no effect on the quality of life or subjective symptoms</td>
</tr>
<tr>
<td>Kerstjens et al</td>
<td>Uncontrolled severe persistent asthma: FEV&lt;sub&gt;1&lt;/sub&gt; &lt; 80% predicted despite high-dose ICS and LABA ACQ ≥ 1.5 &lt; 10 pack-year smoking</td>
<td>Randomized, double-blind, placebo-controlled, crossover</td>
<td>8 weeks</td>
<td>107</td>
<td>Initial 2 weeks: usual dose of ICS and LABA (screen) Subsequent 8-week crossover periods: Tiotropium 5 mcg in morning Tiotropium 10 mcg in morning Placebo</td>
<td>Tiotropium significantly improved lung function when added to ICS - LABA</td>
</tr>
<tr>
<td>Kerstjens et al</td>
<td>Inadequately controlled severe persistent asthma: FEV&lt;sub&gt;1&lt;/sub&gt; ≤ 80% ACQ ≥ 1.5 despite high-dose ICS and LABA &lt; 10 pack-year smoking One exacerbation requiring oral CS in previous year</td>
<td>Randomized, double-blind, placebo-controlled, parallel-group</td>
<td>48 weeks</td>
<td>912</td>
<td>Initial 4 weeks: usual ICS (≤ 800 μg) and LABA (run-in) Subsequent 48 weeks: Tiotropium 5 μg daily in morning Placebo group</td>
<td>Tiotropium improved lung function and reduced exacerbations when added to ICS-LABA</td>
</tr>
<tr>
<td>Peters et al</td>
<td>Inadequately controlled severe persistent asthma: FEV&lt;sub&gt;1&lt;/sub&gt; &lt; 70% or symptoms &gt; 6 days/week or nocturnal symptoms &gt; twice/week &lt; 10 pack-year smoking</td>
<td>Randomized, three-way double-blind, triple-dummy, crossover</td>
<td>14 weeks</td>
<td>210</td>
<td>Initial 4 weeks: ICS 160 μg daily (run-in) Subsequent 14-week crossover periods: Tiotropium 18 μg in morning Salmeterol 50 μg twice daily ICS 320 μg daily (double-dose ICS)</td>
<td>Tiotropium noninferior to salmeterol and superior to double-dose ICS regarding symptoms and lung function</td>
</tr>
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</table>

Abbreviations: ACQ, Asthma Control Questionnaire; CS, corticosteroid; FEV<sub>1</sub>, forced expiratory volume in 1 second; FVC, forced vital capacity; ICS, inhaled corticosteroid; LABA, long-acting inhaled β<sub>2</sub>-agonist.
Table 2 Results summary of the reviewed articles

<table>
<thead>
<tr>
<th>Study</th>
<th>Primary outcome</th>
<th>Secondary outcome</th>
<th>Mode of delivery</th>
<th>Safety</th>
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<tbody>
<tr>
<td>Bateman et al</td>
<td>Change in mean weekly predose PEF: Tiotropium: −3.9±4.87 L/min Salmeterol: −3.2±4.64 L/min Placebo: −24.6±4.84 L/min.</td>
<td>Mean weekly PEF and FEV₁ worse in the placebo group; similar between salmeterol and tiotropium (P&lt;0.01). Rescue medication use, limitation of activities, and daytime asthma symptoms maintained in active treatment groups but worsened in the placebo group. Reduction of rescue medication use statistically significant only in the salmeterol group. Number of asthma-free days slightly increased in the tiotropium and salmeterol groups, more than placebo. Salmeterol slightly superior regarding asthma-free days than tiotropium (1.84 vs 1.34, respectively, 95% CI: −0.955 to −0.069).</td>
<td>Respimat</td>
<td>Adverse events reported by 41.3%, 39.8%, and 41.8% of patients in the placebo, tiotropium, and salmeterol groups, respectively. Individual adverse event reporting generally similar for the three treatment groups. Asthma exacerbations reported in similar rates for the three treatment groups. No cardiovascular events were reported.</td>
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<tr>
<td>Kerstjens et al</td>
<td>Peak FEV₁ within 3 hours after dosing at end of 8-week treatment 5 μg tiotropium: peak FEV₁ significantly higher than placebo (139 mL difference, 95% CI: 96–181 mL). 10 μg tiotropium: peak FEV₁ significantly higher than placebo (170 mL difference, 95% CI: 128–213 mL). There were no significant differences between active treatments.</td>
<td>Significant differences between 5 μg tiotropium and placebo for trough FEV₁ (difference of 86 mL, 95% CI: 41–132 mL) and between 10 μg tiotropium and placebo (difference 113 mL, 95% CI: 67–159 mL). Both treatment doses of tiotropium were superior to placebo in peak FVC, trough FVC, and area under the curve for FVC during the first 3 hours, and weekly AM and PM PEF. The 10 μg dose of tiotropium resulted in significantly better AM and PM PEF than the lower dose. No significant differences in asthma symptoms, use of rescue medications, and change in Mini AQLQ with either dose of tiotropium or placebo.</td>
<td>Respimat</td>
<td>Adverse events in 40%, 42%, and 50% of patients receiving placebo, 5 μg tiotropium, and 10 μg tiotropium, respectively. Dry mouth reported by more patients in the 10 μg tiotropium dose group. Severe adverse events in 5 patients (all nonlife-threatening and nondrug related). No blood pressure, pulse rate, ECG, or laboratory test abnormalities could be attributed to the study drug or dose.</td>
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Table 2 (Continued)

<table>
<thead>
<tr>
<th>Study</th>
<th>Primary outcome</th>
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<th>Mode of delivery</th>
<th>Safety</th>
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<tr>
<td>Peters et al.</td>
<td>AM PEF: tiotropium was superior to double dose of ICS (mean difference 25.8 L/min, P&lt;0.001); Tiotropium was noninferior to salmeterol (difference of 6.4 L/min, P=0.26).</td>
<td>Tiotropium was superior to double dose of ICS for PM PEF (difference of 35.3 L/min, P&lt;0.001), FEV1 (difference of 0.10 L, P=0.004), and daily symptom scores (difference of -9.11 points, P&lt;0.001). Tiotropium was noninferior to salmeterol in other lung function and symptom assessments except for prebronchodilator FEV1 (tiotropium superior with difference of 0.11 L, P=0.003).</td>
<td>HandiHaler</td>
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Abbreviations: ACQ, Asthma Control Questionnaire; AM, morning; AQLQ, Asthma Quality of Life Questionnaire; CI, confidence interval; ECG, electrocardiogram; FEV1, forced expiratory volume in 1 second; FVC, forced vital capacity; ICS, inhaled corticosteroid; NO, nitric oxide; PEF, peak expiratory flow; PM, evening; Rv, residual volume; TLC, total lung capacity.

Inclusion criteria for this clinical trial required that study participants be adults with a clinically established diagnosis of asthma and with ≥15% reversibility after an SABA or a short-acting inhaled muscarinic antagonist (SAMA). Inclusion criteria also required a forced expiratory volume in 1 second (FEV1) of ≤65%. Patients who had had a recent exacerbation or had smoked at any time during their life. The study was a double-blind, randomised, placebo-controlled crossover study. Patients were changed from their usual asthma medications to 1,000 μg of fluticasone daily for a 4-week run-in period. Their dose of fluticasone was then halved to 500 μg/day and an LABA of formoterol was added (12.5 μg per puff and two puffs twice daily). At that point, patients entered a crossover phase in which they were randomized first in the addition of either a metered dose inhaler (MDI) or a matched placebo before, secondly, receiving bronide (18 μg administered once daily from a metered dose inhaler) or a matched placebo, before, secondly, receiving Handihaler (1 μg administered twice daily). A specific primary outcome measure was not identified in the run-in phase, but only 18 patients completed the study.
response to tiotropium. Safety information was not provided in this study.

Peters et al. compared the benefit of tiotropium as add-on therapy to an ICS with that obtained by either adding an LABA or doubling the ICS dose in patients with uncontrolled persistent asthma. Patients were included if they were nonsmokers (less than 10 pack-years per lifetime), were aged 18 years or older, had an FEV₁ >40% of the predicted value, and had a diagnosis of asthma confirmed by bronchodilator reversibility or bronchial hyperresponsiveness. The study was a three-way, double-blind, triple-dummy, crossover trial. Patients entered a 4-week run-in period during which they were treated with 160 μg of beclomethasone daily. They were entered into the trial if they had either an FEV₁ ≤70% or daytime symptoms for 6 or more days per week or night-time symptoms 2 or more nights per week during the final 2 weeks of the run-in period. The three crossover periods involved adding tiotropium (18 μg every AM from a dry powder inhaler), adding salmeterol (50 μg twice daily by dry powder inhaler), or doubling the ICS to 320 μg daily. Each treatment period lasted 14 weeks with a 2-week washout phase between treatment periods. The identified primary outcome measure in this study was the AM PEF, and the primary hypothesis was that the addition of tiotropium to ICS would provide greater benefit than doubling the dose of ICS. Two hundred and ten severe asthma patients with a mean FEV₁ of 2.31 L (71.5% predicted) met the entry criteria and were randomized. Thirty-six participants dropped out during the treatment phase due to withdrawal of consent, loss to follow-up, drug-related adverse events, or miscellaneous reasons. The use of tiotropium resulted in a superior primary outcome compared with doubling the dose of an ICS. There was a greater improvement in the primary outcome measure AM PEF with tiotropium compared with a double dose of ICS, with a mean difference of 25.8 L/min (P<0.001). Tiotropium also showed superiority in most secondary outcomes. PM PEF was larger, with a difference of 35.5 L/min (P<0.001); the proportion of asthma control days was greater, with a difference of 0.079 (P=0.01); the FEV₁ before bronchodilation was larger, with a difference of 0.10 L (P=0.004); and daily asthma symptom scores were lower, with a difference of −0.11 points (P<0.001). In the secondary outcome analysis, addition of tiotropium to an ICS was found to be noninferior to salmeterol, as measured by the AM PEF. Although the prebronchodilator and postbronchodilator FEV₁ responses favored tiotropium, tiotropium was noninferior to salmeterol in other results, including PM PEF, asthma daily symptom scores, and sputum eosinophils. Interestingly, a small FEV₁ decrease in patients receiving albuterol in addition to the standing dose of salmeterol suggested a tachyphylaxis effect for β-agonists that was not found in the tiotropium group. Not increasing the dose of ICS by more than a factor of two was proposed as a potential weakness of the trial, together with the short study duration (14 weeks) and the small sample size. Although long-term safety issues were not assessed in this trial, numerically more asthma exacerbations occurred during the double-dose ICS treatment period than during the tiotropium treatment period (n=16 versus [vs] n=9), with the fewest occurring with salmeterol treatment (n=5).

Kerstjens et al. sought to compare the efficacy and safety of different doses of tiotropium in the Respimat® inhaler (Boehringer Ingelheim Pharma GmbH and Co, KG, Ingelheim, Germany) against placebo as add-on therapy in patients with uncontrolled severe persistent asthma. Patients included in this trial were aged 18 years and older and had at least a 5-year history of asthma. They were required to have a postbronchodilator FEV₁ ≥80% and to have an Asthma Control Questionnaire (ACQ) score of ≥1.5 despite treatment with high-dose ICS and an LABA. Interestingly, patients were allowed to remain on a stable dose of theophylline, LMA, and oral corticosteroid throughout the study. Patients were excluded if they had a smoking history of ≥10 pack-years or a diagnosis of COPD. The authors conducted a randomized, double-blind, crossover study with three 8-week treatment periods. After a 2-week run-in period, eligible patients were randomized to receive each of three treatments in a random sequence for 8 weeks in a crossover design (5 μg or 10 μg of tiotropium or matching placebo administered as two actuations once daily in the AM from the Respimat inhaler). There was no washout phase between treatment periods. Throughout the 24-week treatment period, participants recorded PEF and FEV₁ values twice daily at home using the Asthma Monitor AM2+ and responded to daily questions in the electronic diary component of this device. The primary end point was peak FEV₁ (obtained within 3 hours of dosing) at the end of each treatment period. Of the 107 randomized patients, 100 completed all treatment periods. Peak FEV₁ was significantly higher with 5 μg of tiotropium (difference 139 mL; 95% CI: 96–181 mL) and 10 μg (difference 170 mL; 95% CI: 128–213 mL) than with placebo (both P<0.0001). Trough peak FEV₁, at the end of the dosing interval was also higher with both doses of tiotropium than with placebo (difference with 5 μg of 86 mL [95% CI: 41–132]; difference with 10 μg of 113 mL [95% CI: 67–159 mL], both P<0.0004). Although there were numerically greater FEV₁ responses with the 10 μg dose of tiotropium than with the 5 μg dose compared with placebo.
for both peak and trough FEV₁, the differences in outcome measures between the two active doses were not significant. Across groups, no significant difference was noted in asthma-related health status, rescue medication use, or symptoms measured in the electronic diary. There was a trend for more adverse events to be reported with the 10 μg tiotropium treatment group than with the placebo group. Patients receiving the highest dose of tiotropium specifically reported more complaints about dry mouth than the placebo group (6.8% vs 1.0%). Serious adverse events and asthma exacerbations were few and balanced among the treatment groups.

Bateman et al²² compared the efficacy and safety of tiotropium as add-on therapy to patients with moderate persistent asthma not adequately controlled by ICS therapy alone against salmeterol and placebo. As a novel twist of this study, the authors focused their study on a specific group of asthma patients with the B16-Arg/Arg genotype. Patients with a history of asthma with the B16-Arg/Arg genotype and receiving maintenance therapy with ICS of 400–1,000 μg/day and receiving maintenance therapy with ICS of 400–1,000 μg/day or budesonide or equivalent were eligible for enrollment. The main inclusion criteria were reversible airway obstruction (improvement in FEV₁ of at least 12% and 200 mL after sequential inhalation of 80 μg of ipratropium bromide and 400 μg of salbutamol) and prebronchodilator FEV₁ of 90% of predicted value or less for patients previously receiving ICSs only. The main exclusion criteria were smoking history (>10 pack-years), COPD, serious concomitant illnesses, and concomitant respiratory medications. They conducted a double-blind, double-dummy, placebo-controlled, parallel-group trial. There was a 4-week run-in period during which patients received 50 μg of twice-daily salmeterol administered through a metered dose inhaler along with their ICS. After the run-in period, patients were randomized to 16 weeks of treatment with tiotropium (two puffs of 2.5 μg daily in the PM through the Respimat inhaler), salmeterol metered dose inhaler (two puffs of 25 μg twice daily plus tiotropium-matching placebo), or placebo (with both a tiotropium and salmeterol matching placebo). The primary end point was change in weekly AM (predose) PEF from the last week of run-in period to the last week of treatment. A total of 530 patients were eligible for the trial and 388 were randomized. At baseline almost all patients were receiving ICSs. The majority of patients in each treatment group were also receiving a LABA. The prebronchodilator FEV₁ was between 68% and 70% for the three treatment groups. There was, on average, between a 24% and 28% increase in FEV₁ after ipratropium and albuterol. The AM PEF was maintained with tiotropium (−3.9±4.87 L/min change from run-in) and salmeterol (−3.2±4.64 L/min change from run-in), but the AM PEF decreased with placebo (−24.62±4.84 L/min from run-in). Both active treatments provided significantly superior effects on AM PEF compared with placebo. The tiotropium effect was noninferior to salmeterol. Weekly FEV₁ measures showed the same treatment effects as the AM PEF. Surprisingly, little evidence of treatment effect with tiotropium and salmeterol was found in asthma symptoms, rescue medication use, and the Mini AQLQ. In general, adverse event reporting was comparable across treatment groups. There were numerically more asthma exacerbations with salmeterol treatment (n=4) than with either placebo (n=1) or tiotropium (n=0).

In two large replicate studies, Kerstjens et al²³ compared the effect of adding 5 μg of tiotropium each AM to placebo on lung function and exacerbations in poorly controlled asthma patients already receiving ICS and LABA. Included patients were aged between 18 years and 75 years and had a 5-year or longer asthma history. Other inclusion criteria consisted of an ACQ score of 1.5 or higher, a postbronchodilator FEV₁ ≤80% predicted, and a postbronchodilator FVC ≤70% predicted despite daily therapy with high-dose ICS and LABA. Included patients also had at least one asthma exacerbation in the preceding year that required treatment with systemic corticosteroids. Main exclusion criteria consisted of a pre-existent diagnosis of COPD, serious coexisting illness, and concurrent use of anticholinergic bronchodilators. Patients with a ≥10 pack-years smoking history and who had smoked within the past year were also excluded. The study was a randomized, double-blind, placebo-controlled, parallel-group design. After a 4-week screening period, patients were randomized to treatment with either tiotropium (5 μg daily) or placebo, both administered by the Respimat Soft Mist Inhaler in the AM for 48 weeks, which made it the longest trial carried out on this subject. The primary outcomes were the peak FEV₁ (measured within 3 hours of treatment) and trough FEV₁ at week 24 expressed as change from baseline. In the two studies, 912 eligible patients were randomized. At 24 weeks, the mean change in the peak FEV₁ from baseline was greater with tiotropium than with placebo in the two trials: a difference of 86±34 mL in trial 1 (P=0.01) and 154±32 mL in trial 2 (P<0.001). Similar significant differences in favor of tiotropium treatment were found with the trough FEV₁. The effects of tiotropium continued throughout the entire 48 weeks of the study. In addition, the addition of tiotropium increased the time to the first severe
exacerbation (282 days vs 226 days) with an overall reduction of 21% in the risk of a severe exacerbation (hazard ratio 0.79; \( P = 0.03 \)). Consistent differences between tiotropium and placebo treatment on the ACQ, the AQLQ symptom-free days, and rescue medication use were not found, however. Adverse event reporting was slightly lower with tiotropium than with placebo. No differences were found between the groups in cardiac adverse events and serious adverse events. Treatment with tiotropium was associated with a slightly higher complaint rate of dry mouth.

We performed an assessment of the methodologic quality of the five studies, and we assessed the consistency and adherence of the five trials to the CONSORT. We concluded that each study met the CONSORT criteria for a well-designed randomized trial.\(^{18}\)

**Discussion**

Tiotropium is an inhaled long-acting anticholinergic currently approved in the US for once-daily maintenance treatment of bronchospasm associated with COPD.\(^{26}\) It has been proven to be an effective bronchodilator and to reduce the frequency of COPD exacerbations.\(^{10,11}\) Despite the well-studied pharmacology of tiotropium, its role in asthma remains unproven. The drug is currently not approved for non-COPD obstructive airway disease. In this systematic review we found that adding tiotropium to pre-existing therapy with only an ICS or an ICS–LABA combination resulted in significant improvements in lung function. There was also evidence from one study suggesting a reduction in exacerbations in patients with uncontrolled severe asthma. These observations are supported by findings in open-label, uncontrolled studies. The findings are consistent with the known effect of a muscarinic antagonist and its mechanism of action in the context of asthma pathophysiology.

The five clinical studies included in this systematic review focused on evaluating the efficacy of tiotropium as add-on therapy to ICS or ICS in combination with an LABA in patients with uncontrolled moderate to severe persistent asthma. Tiotropium maintained lung function when ICSs were tapered and when an LABA was discontinued.\(^{19,22}\) Tiotropium improved lung function when added to an ICS alone or an ICS–LABA combination therapy.\(^{20–23}\) In the only trial to have compared the addition of tiotropium with doubling the dose of ICS, tiotropium provided significantly superior results.\(^{20}\) In trials in which the addition of tiotropium was compared with salmeterol, the beneficial effects of these two bronchodilators was similar.\(^{20,22}\) Although the improvements in lung function appeared to be clinically meaningful, these clinical trials failed to show that tiotropium provided consistent and clinically relevant improvements in respiratory symptoms, rescue medication use, or quality of life. Perhaps the failure to show symptomatic improvements was a function of study size. In the one report powered sufficiently to assess any change in the symptomatic burden of asthma, the addition of tiotropium resulted in an overall reduction of exacerbations and an increase in the time between them.\(^{23}\) Overall, the consistent improvement in lung function with the addition of tiotropium should probably lead to US Food and Drug Administration approval of tiotropium as add-on therapy for patients with uncontrolled moderate to severe persistent asthma already on ICSs. Unfortunately, the available information does not enable health care providers to determine whether an individual patient might respond more to tiotropium or an LABA as add-on therapy.\(^{27}\)

Although excluded from the systematic review, the two open-label trials we found on this topic support the findings of the other five studies. Iwamoto et al\(^ {24}\) added tiotropium to 17 patients with severe persistent asthma who were receiving high-dose ICSs. Many of these patients were also taking an LABA and/or oral corticosteroids. The study was an open-label, uncontrolled design. They found that use of tiotropium for 4 weeks significantly improved FEV\(_1\), whereas the neutrophil proportion was directly correlated with an increase in FEV\(_1\). This suggested that the noneosinophilic sputum profile may separate responders vs nonresponders to tiotropium therapy. A similar interest in distinguishing responders from nonresponders was reflected in the study conducted by Park et al.\(^ {25}\) They studied 138 patients with severe persistent asthma who had reduced lung function despite high-dose ICSs and LABA. Tiotropium, administered as 18 \( \mu \)g daily from a dry powder inhaler, was added to this regimen. Lung function tests were evaluated every 4 weeks for a minimum of 12 weeks. Responders were defined as having a > 15% (or > 200 mL) improvement in FEV\(_1\), with a sustained effect lasting at least 8 successive weeks. Forty-six of the 138 patients (33.3%) responded to tiotropium treatment. Logistic regression analysis (controlled for age, sex, and smoking status) showed that Arg16Gly in \( ADRB2 \) \( (P = 0.003, \) odds ratio \( 0.21, \) 95% CI: \( 0.07–0.59) \) in a minor allele-dominant model was significantly associated with the response to tiotropium.

Improved lung function (increased FEV\(_1\) and PEF) in asthma with tiotropium can be explained by its mechanism of action in the context of asthma pathophysiology. The major bronchi (the site of bronchoconstriction in asthma) have the
densest population of cholinergic nerves in the respiratory tract. Bronchoconstriction is mainly mediated through cholinergic mechanisms in humans. Stimulation of cholinergic fibers leads to activation of M3 Ach receptors in the airway smooth muscle, which, in turn, causes bronchoconstriction and mucus secretion. Tiotropium is an effective M3 Ach receptor antagonist. Cholinergic stimulation may also have influences in addition to controlling airway smooth muscle contraction. Cholinergic pathways influence mucus secretion and possibly airway inflammation. It has been speculated that tiotropium might be able to affect airway remodeling in asthma through these mechanisms.  

Only four of the clinical trials reported complete information on adverse events related to tiotropium use. In general, similar total and individual reporting of adverse events was described for tiotropium and salmeterol compared with placebo. The only exception was that patients receiving higher doses of tiotropium (eg, 10 μg) tended to report dry mouth more frequently. The cardiovascular safety profile of tiotropium is of particular interest. In COPD patients, Verhamme et al found that tiotropium administered 5 μg daily by the Respimat Soft Mist Inhaler was associated with about a 30% increase in mortality compared with a comparable dose of tiotropium administered by a dry powder inhaler. The mortality association was strongest for cardiovascular and cerebrovascular death. However, it remains unclear whether this association is causal or due to residual confounding by COPD severity. A more recent study, also performed in COPD patients, found that tiotropium administered by the Respimat Soft Mist Inhaler at a dose of 5 μg or 2.5 μg had a safety profile similar to that of tiotropium given by dry powder inhaler at a dose of 18 μg. The 48-week studies reported by Kerstjens et al described no deaths and few cardiovascular safety events in asthma patients using tiotropium administered by the Respimat Soft Mist Inhaler.

There are strengths and limitations to this systematic review. Although the number of studies included was small, the studies were of high quality. They were randomized and controlled. All studies met the CONSORT criteria for reporting of randomized trials. There were limitations, though, in the five studies. One study was small, with only 18 completed patients. Two studies were short, only 4 and 8 weeks. Consequently, information available regarding sustainability of benefit is somewhat limited. The trials were a mixture of crossover and parallel-group design, making integration of efficacy results impractical. There may still be unresolved issues regarding the appropriate dosing of tiotropium in the treatment of asthma. Different inhalation devices (with different doses) were used to administer tiotropium, metered dose inhaler, dry powder inhaler, and the Respimat Soft Mist Inhaler. One study compared the effect of 5 μg of tiotropium from the Respimat vs 10 μg. The effects on lung function of the 10 μg tiotropium dose tended to be greater, but there was also a higher reporting rate of dry mouth with the larger dose. It is unclear whether the 10 μg tiotropium dose might provide additional benefits with long-term use, such as a further reduction in exacerbation rates. Although two of the studies using tiotropium in the Respimat had patients self-administer their dose in the AM, one of the studies required PM dosing. It is difficult to determine from these studies whether there might be greater benefits of PM vs AM dosing.

There was limited information available from these studies regarding the anti-inflammatory benefits of tiotropium in uncontrolled asthma. Fardon et al found that addition of tiotropium was associated with a significant reduction in exhaled nitric oxide levels, but Peters et al found that immunoglobulin E levels, sputum eosinophil counts, and exhaled nitric oxide levels were not predictive of a clinical response to tiotropium. Similarly, reversibility testing to either an SABA or an SAMA was not helpful in identifying patients who would have a beneficial tiotropium response. Because all studies required SABA reversibility as an inclusion criterion, it is unclear whether an uncontrolled asthma patient with moderate to severe persistent disease but without reversibility would have a beneficial response to tiotropium.

In summary, there may be a beneficial role for tiotropium in moderate to severe persistent asthma patients who are uncontrolled despite use of an ICS or an ICS and an LABA. However, it remains to be determined whether the spirometric changes will ultimately reciprocate in long-term symptom improvement. The currently available literature lays a sound foundation for further studies to be designed with particular focus on the asthma population. An important issue will be to determine whether patterns of genotypes, physiology, or inflammatory markers can be identified that would identify patients with the maximal potential benefit from treatment with tiotropium in addition to currently approved therapy.

Disclosure
Drs Befekadu and Onofrei report no conflicts of interest in this work. Dr Colice is a consultant/advisory board member/speaker for Teva, MedImmune, Pearl, Dey, Mylan, Forest, and Alitair, which are pharmaceutical companies developing treatments for asthma and COPD.
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