ORIGINAL ARTICLE

Comparative safety of inhaled medications in patients with chronic obstructive pulmonary disease: systematic review and mixed treatment comparison meta-analysis of randomised controlled trials

Yaa-Hui Dong,¹ Hsien-Ho Lin,¹ Wen-Yi Shau,² Yun-Chun Wu,¹ Chia-Hsuin Chang,¹,³ Mei-Shu Lai¹

ABSTRACT

Background The active-treatment comparative safety information for all inhaled medications in patients with chronic obstructive pulmonary disease (COPD) is limited. We aimed to compare the risk of overall and cardiovascular death for inhaled medications in patients with COPD.

Methods Through systematic database searching, we identified randomised controlled trials of tiotropium Soft Mist Inhaler, tiotropium HandiHaler, long-acting β2 agonists (LABAs), inhaled corticosteroids (ICS), and LABA-ICS combination with at least a 6-month treatment duration. Direct comparison and mixed treatment comparison (MTC) meta-analyses were conducted to estimate the pooled ORs of death for each comparison.

Results 42 trials with 52,516 subjects were included. The MTC meta-analysis with the fixed effect model indicated tiotropium Soft Mist Inhaler was associated with an universally increased risk of overall death compared with placebo (OR 1.51; 95% CI 1.06 to 2.19), tiotropium HandiHaler (OR 1.65; 95% CI 1.13 to 2.43), LABA (OR 1.63; 95% CI 1.10 to 2.44) and LABA-ICS (OR 1.90; 95% CI 1.28 to 2.86). The risk was more evident for cardiovascular death, in patients with severe COPD, and at a higher daily dose. LABA-ICS was associated with the lowest risk of death among all treatments. No excess risk was noted for tiotropium HandiHaler or LABA. The results were similar for MTC and direct comparison meta-analyses, with less precision in the random effects model.

Conclusion Our study provided a comparative safety spectrum for each category of inhaled medications. Tiotropium Soft Mist Inhaler had a higher risk of mortality and should be used with caution.

INTRODUCTION

Chronic obstructive pulmonary disease (COPD) is a substantial disease burden worldwide.¹ ² The Global Initiative for Chronic Obstructive Lung Disease guidelines recommend inhaled long-acting anticholinergic and β2 agonists (LABAs) for maintenance therapy of COPD.² Add-on treatment with inhaled corticosteroids (ICS) is indicated for patients with repeated exacerbations.² However, an association of inhaled medications with cardiovascular complications has been noted, which is possibly related to their pharmacological effects.³ ⁴ Moreover, patients with COPD are susceptible to overall and cardiovascular death.⁵ ⁶ Accordingly, it is important to examine the safety profiles of both outcomes for these medications.

Tiotropium, the only marketed long-acting anticholinergic, is approved as dry powder delivered via a HandiHaler device and solution delivered via a Respimat Soft Mist Inhaler (Boehringer Ingelheim, Ingelheim am Rhein, Germany). Based on the results of a large placebo-controlled trial,⁷(please note, references with the prefix ‘w’ are listed in the online appendix), the US Food and Drug Administration (FDA) suggested that tiotropium HandiHaler has no excess risk of overall death and cardiovascular events.⁵ ⁸ ⁹ However, a non-significant increased risk of overall death for tiotropium Soft...
Mist Inhaler was noted in placebo-controlled trials, and a nearly 50% increased risk of overall death compared with placebo was reported in a meta-analysis. For LABAs, the US FDA has continually highlighted the increased risk of death in patients with asthma. However, safety concerns about LABAs remain inconclusive in patients with COPD. A nearly 50% increased risk of overall death compared with placebo was observed in several meta-analyses. 

We also recorded how adverse events were monitored. Any risk of mortality for inhaled medications in patients with COPD, left unspecified by the Medical Dictionary for Regulatory Activities (see the online appendix). First, none of the published trials simultaneously compared safety across various inhaled medications. Second, given the limited sample size, the risks of death for inhaled medications are generally undetermined by individual trials. Third, although traditional meta-analyses provide pooled risk estimates with better precision, the estimates are obtained only from direct comparison trials. Consequently, the information on comprehensive comparisons among different inhaled medications is insufficient.

The mixed treatment comparison (MTC) meta-analysis has a unique strength to integrate data from direct and indirect comparisons and facilitate multiple head-to-head comparisons across various treatments. To comprehensively compare the risk of mortality for inhaled medications in patients with COPD, we conducted both direct comparison and MTC meta-analyses of randomised controlled trials.

METHODS

Data sources and searches
We searched MEDLINE, CINAHL, Cochrane Library and ClinicalTrials.gov from inception to July 2011 (see the online appendix for the detailed search strategy). We examined the bibliographies of eligible trials and systematic review articles for relevant trials. To identify unpublished trials, we searched the manufacturers’ clinical trials registers. If the outcomes of interest were not available from original articles or the above clinical trials registers, we contacted the authors or searched the US FDA website for additional information.

Study selection
The inclusion criteria were randomised, double-blind, active or placebo-controlled trials; patients with COPD of any severity; patients receiving predefined treatments, including tiotropium Soft Mist Inhaler, tiotropium HandiHaler, LABA, LABA-ICS and ICS; trials providing data about overall or cardiovascular death; and trials lasting for 6 months or more. We excluded trials if they included patients with asthma, involved non-predefined treatment arms, and were published only in protocols, in abstracts, or in non-English languages.

Outcome measures
The primary outcome was overall death. The secondary outcome was cardiovascular death based on the preferred terms across various treatments. To address the possible trial heterogeneities and misclassification of cardiovascular death, we performed additional analyses. Meta-regression was conducted to adjust for related demographic characteristics (see the online appendix). Sensitivity analyses were performed by excluding trials with the ICS withdrawal design and by restricting the analyses to trials with objective adjudication of cause of death.

STATA V.9.0 (StataCorp) and WinBUGS V.1.4.3 (MRC Biostatistics Unit, Cambridge, UK) was used for direct comparison and MTC meta-analyses, respectively. The WinBUGS code is shown in the online appendix. The predefined protocol is available from the authors on request.

RESULTS

Eligible trials
A total of 42 eligible trials reporting on overall death and 51 trials reporting on cardiovascular death were included in the meta-analysis (online appendix and figure 1). These 42 trials enrolled 52,516 subjects, with similar characteristics across trials with different treatments (64 years of age, 73% men, 44% current smokers, 1 year study duration, 44% of predicted value in FEV1). However, more subjects combined LABA or ICS use at baseline in tiotropium Soft Mist Inhaler and HandiHaler trials (table 1 and online tables S1 and S2).

Assessment of risk of bias of included studies
All 42 trials were randomised, double-blind, with 24 trials addressing adequate randomisation procedures. Forty-one trials stated the withdrawal rate, which varied across trials and treatment groups (with the lowest value of 17% in the tiotropium Soft Mist Inhaler group and the highest values of 33% in the ICS and placebo groups). Twenty-eight trials described the fraction of lost to follow-up, which was relatively disagreement was resolved by discussion and consensus (see the online appendix).

Statistical analysis
We used the intention-to-treat analysis. A two-sided value of 0.05 was defined for statistical significance. The Peto method was applied for the direct comparison meta-analysis of rare events. The Mantel-Haenszel method with the fixed effect model and different continuity correction factors was performed for the sensitivity analysis. For each pairwise comparison, we estimated the risks of overall and cardiovascular death with the pooled OR and 95% CI. Statistical heterogeneity was evaluated by the I² statistic, with a value of 50% or more indicating a substantial level of heterogeneity. Publication bias was assessed by the funnel plot, the Begg’s test and the Egger’s test (see the online appendix).

The Bayesian Markov chain Monte Carlo methods with fixed and random effects models were used for the MTC meta-analysis (see the online appendix). Results were presented as the OR and 95% credible interval (CrI). For each treatment, we also estimated the probability of overall and cardiovascular death and the probability of being ranked as the riskiest intervention.

We performed subgroup analyses to address the risks of inhaled medications in trials with longer treatment durations (study duration ≥1 year) and trials enrolling patients with severe COPD (forced expiratory volume in 1 s (FEV1) <50% of predicted value). A potential dose–response relation of tiotropium Soft Mist Inhaler was examined by stratification analyses by individually comparing 5 µg/day and 10 µg/day of tiotropium Soft Mist Inhaler with other treatments.

To address the possible trial heterogeneities and misclassification of cardiovascular death, we performed additional analyses. Meta-regression was conducted to adjust for related demographic characteristics (see the online appendix). Sensitivity analyses were performed by excluding trials with the ICS withdrawal design and by restricting the analyses to trials with objective adjudication of cause of death.

Thorax first published as 10.1136/thoraxjnl-2012-201926 on 6 October 2012. Downloaded from http://thorax.bmj.com/ on August 1, 2023 by guest. Protected by copyright.
low and less variant across different treatment groups (Table 1 and online tables S2 and S3). Practice of adverse event monitoring was heterogeneous across trials, with six trials (25,533 subjects) describing objective adjudication of cause of death (online table S3).

Direct comparison meta-analysis

Figure 2 displays the network of each pairwise comparison. Statistical heterogeneity was minimal, with the exception of tiotropium HandiHaler versus LABA for overall death ($I^2 = 59.4\%$).

Table 1  Summary of trial characteristics and patient characteristics at baseline. Figures are means (ranges)

<table>
<thead>
<tr>
<th></th>
<th>Age (years)</th>
<th>Men (%)</th>
<th>Current smokers (%)</th>
<th>Study duration (years)</th>
<th>FEV$_1$ (% of predicted value)</th>
<th>Subjects with concomitant use of LABA (%)</th>
<th>Subjects with concomitant use of ICS (%)</th>
<th>Withdrawal rate (%)</th>
<th>Lost to follow-up (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total (N=42)</td>
<td>64.0</td>
<td>73.2</td>
<td>73.1</td>
<td>1.2 (0.5–4)</td>
<td>44.4 (34.4–86.6)</td>
<td>33.3 (3.0–60.1)</td>
<td>47.3 (14.0–83.2)</td>
<td>27.9 (0.8–59.0)</td>
<td>1.5 (0.0–8.6)</td>
</tr>
<tr>
<td>Stratified by treatment*</td>
<td></td>
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</tr>
<tr>
<td>TIO-SMI (n=3)</td>
<td>64.9</td>
<td>76.4</td>
<td>45.5</td>
<td>41.5 (29.7–53.4)</td>
<td>54.9 (53.7–56.1)</td>
<td>17.1 (16.0–20.4)</td>
<td>1.3 (1.1–1.7)</td>
<td></td>
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</tr>
<tr>
<td>TIO-HH (n=12)</td>
<td>64.4</td>
<td>77.0</td>
<td>45.3</td>
<td>42.5 (17.5–53.8)</td>
<td>59.1 (42.4–75.3)</td>
<td>24.3 (14.7–42.0)</td>
<td>1.1 (0.2–2.1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LABA (n=19)</td>
<td>63.7</td>
<td>72.0</td>
<td>43.9</td>
<td>34.3 (3.0–54.5)</td>
<td>37.3 (14.0–57.0)</td>
<td>25.2 (10.0–45.3)</td>
<td>1.1 (0.4–3.0)</td>
<td></td>
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</tr>
<tr>
<td>LABA-ICS (n=17)</td>
<td>64.9</td>
<td>72.2</td>
<td>36.5</td>
<td>28.6 (5.0–45.3)</td>
<td>34.6 (14.0–54.3)</td>
<td>26.8 (12.2–39.0)</td>
<td>1.9 (0.0–2.5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ICS (n=16)</td>
<td>63.3</td>
<td>69.5</td>
<td>44.0</td>
<td>29.1 (8.5–56.0)</td>
<td>44.0 (19.8–83.2)</td>
<td>33.5 (0.8–43.8)</td>
<td>2.0 (0.0–8.6)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>PL (n=34)</td>
<td>64.1</td>
<td>73.2</td>
<td>44.1</td>
<td>34.2 (8.5–60.1)</td>
<td>52.3 (19.8–83.2)</td>
<td>33.3 (0.8–59.0)</td>
<td>2.0 (0.0–6.8)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*N* and *n* represent the number of trials included in the meta-analysis and with each treatment, respectively. *The variables of age, proportion of men, percentage of current smokers, study duration, FEV$_1$, % of predicted value, and percentage of subjects with concomitant use of LABA or ICS were calculated by trials of individual treatments. The variables of withdrawal rate and lost to follow-up were calculated by treatment groups.

FEV$_1$, forced expiratory volume in 1 s; ICS, inhaled corticosteroid; LABA, long-acting $\beta_2$ agonist; PL, placebo; TIO-HH, tiotropium dry powder delivered via HandiHaler; TIO-SMI, tiotropium solution delivered via Respinmat Soft Mist Inhaler.
tiotropium Soft Mist Inhaler had universally increased risks compared with those receiving placebo (OR 1.51; 95% CrI 1.06 to 2.19) or those using tiotropium HandiHaler (OR 1.65; 95% CrI 1.13 to 2.43), LABA (OR 1.63; 95% CrI 1.10 to 2.44) and LABA-ICS (OR 1.90; 95% CrI 1.28 to 2.86). In contrast, LABA-ICS demonstrated a beneficial profile versus placebo (OR 0.80; 95% CrI 0.67 to 0.94) or ICS (OR 0.77; 95% CrI 0.64 to 0.93). For cardiovascular death, tiotropium Soft Mist Inhaler had a more apparent risk compared with placebo (OR 2.07; 95% CrI 1.09 to 4.16), tiotropium HandiHaler (OR 2.38; 95% CrI 1.20 to 4.99), LABA (OR 3.04; 95% CrI 1.48 to 6.55), LABA-ICS (OR 2.79; 95% CrI 1.87 to 6.02) and ICS (OR 2.39; 95% CrI 1.18 to 5.12). In contrast, LABA had a decreased risk versus placebo (OR 0.68; 95% CrI 0.50 to 0.95). In the random effects model, tiotropium Soft Mist Inhaler consistently demonstrated an increased risk of overall death versus any comparators, and showed an increased risk of cardiovascular death versus LABA-ICS. Between direct comparison and MTC meta-analyses, the direction in OR was identical for each comparison. The difference in OR was also minimal (within 10%), with the exception of tiotropium HandiHaler versus LABA or LABA-ICS (over 30%).

Among all the treatments, tiotropium Soft Mist Inhaler had the highest probability of overall and cardiovascular death (8% and 5.5%, respectively), with an approximate probability of 95% of being ranked as the riskiest treatment. In contrast, LABA-ICS had the lowest probability of overall death (4.5%), with a probability of 0% of being ranked as the riskiest treatment (table 5).

The analyses restricted to trials with longer treatment duration and trials enrolling patients with severe COPD showed similar results to the main analysis, although the risk of cardiovascular death associated with tiotropium Soft Mist Inhaler was slightly higher for patients with severe COPD using the fixed effect model (table 4). Three tiotropium Soft Mist Inhaler trials included a group treated with 5 μg/day and two included a group treated with 10 μg/day. Use of 10 μg/day tiotropium Soft Mist Inhaler tended to be associated with a higher risk of overall death against all comparators, although the risks of cardiovascular death were irrespective of the dose of tiotropium (figure 3).

The MTC meta-regression adjusting for demographic characteristics did not substantially change the increased risk for tiotropium Soft Mist Inhaler versus other treatments (online tables S5 and S6). The sensitivity analyses which excluded trials with the ICS withdrawal design and restricted trials with objective adjudication of cause of death yielded similar results to the main analysis (online tables S7 and S8).

**DISCUSSION**

This study demonstrated that tiotropium Soft Mist Inhaler was associated with higher risks of overall and cardiovascular death compared with placebo and other inhaled medications, with a potential dose–response effect on overall death. The risk of cardiovascular death associated with tiotropium Soft Mist Inhaler was also higher for patients with severe COPD, although this may simply be because more severe comorbidities impair cardiovascular systems. Instead, LABA-ICS was associated with the lowest risk of overall death. No excess risk was noted for tiotropium HandiHaler or LABA. In contrast with the Global Initiative for Chronic Obstructive Lung Disease guidelines recommending tiotropium or LABA as first-line maintenance therapy, our study highlighted the potential harm of tiotropium Soft Mist Inhaler. This finding should be weighed against other risks and benefits for individual treatments and allows a revision of the recommendations for management of COPD.
Data suggested that the elevated risks of tiotropium Soft Mist Inhaler had an increased risk of overall death (risk ratio 1.52; 95% CI 1.06 to 2.16) and cardiovascular death (risk ratio 2.05; 95% CI 1.06 to 3.99) versus placebo.11 Our study further highlighted safety concerns of tiotropium Soft Mist Inhaler versus other active treatments. Some possible mechanisms have been proposed. One is that use of tiotropium Soft Mist Inhaler could yield higher systematic drug exposure compared with tiotropium HandiHaler,29 although this was not observed in Japanese research.30 Evidence also suggested that patients receiving a higher dose of tiotropium Soft Mist Inhaler had higher peak concentrations.29 Another mechanism is the potential effect of underlying rhythm disorders on mortality. Data suggested that the elevated risks of tiotropium Soft Mist Inhaler could be attributed to the higher peak concentrations which increase the risk of overall death and cardiovascular death.

### Table 2: Risk of overall death and cardiovascular death for each pairwise comparison from the direct comparison and MTC meta-analyses

<table>
<thead>
<tr>
<th>Comparison</th>
<th>Overall death (N = 42)</th>
<th>Cardiovascular death (N = 31)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Direct comparison Peto OR (95% CI)</td>
<td>MTC fixed effect OR (95% CI)</td>
</tr>
<tr>
<td>TIO-SMI vs</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TIO-HH</td>
<td>1.65 (1.13 to 2.43)</td>
<td>1.66 (1.04 to 2.75)</td>
</tr>
<tr>
<td>LABA</td>
<td>1.63 (1.10 to 2.44)</td>
<td>1.61 (1.002 to 2.68)</td>
</tr>
<tr>
<td>LABA-ICS</td>
<td>1.90 (1.28 to 2.96)</td>
<td>1.93 (1.20 to 3.24)</td>
</tr>
<tr>
<td>ICS</td>
<td>1.47 (0.99 to 2.21)</td>
<td>1.55 (0.96 to 2.65)</td>
</tr>
<tr>
<td>PL</td>
<td>1.49 (1.05 to 2.11)</td>
<td>1.51 (1.06 to 2.19)</td>
</tr>
<tr>
<td>TIO-HH vs LABA</td>
<td>0.76 (0.55 to 1.06)</td>
<td>0.96 (0.63 to 1.49)</td>
</tr>
<tr>
<td>TIO-HH vs LABA-ICS</td>
<td>1.81 (1.07 to 3.05)</td>
<td>1.16 (0.95 to 1.41)</td>
</tr>
<tr>
<td>TIO-HH vs ICS</td>
<td>0.89 (0.73 to 1.09)</td>
<td>0.93 (0.71 to 1.31)</td>
</tr>
<tr>
<td>TIO-HH vs PL</td>
<td>0.93 (0.81 to 1.07)</td>
<td>0.92 (0.81 to 1.05)</td>
</tr>
<tr>
<td>TIO-SMI vs LABA</td>
<td>1.10 (0.91 to 1.32)</td>
<td>1.17 (0.98 to 1.39)</td>
</tr>
<tr>
<td>TIO-SMI vs ICS</td>
<td>0.86 (0.71 to 1.04)</td>
<td>0.90 (0.76 to 1.07)</td>
</tr>
<tr>
<td>TIO-SMI vs PL</td>
<td>0.90 (0.75 to 1.08)</td>
<td>0.93 (0.79 to 1.09)</td>
</tr>
<tr>
<td>LABA-ICS vs ICS</td>
<td>0.78 (0.64 to 0.94)</td>
<td>0.77 (0.64 to 0.93)</td>
</tr>
<tr>
<td>LABA-ICS vs PL</td>
<td>0.81 (0.67 to 0.98)</td>
<td>0.80 (0.67 to 0.94)</td>
</tr>
<tr>
<td>LABA vs PL</td>
<td>1.01 (0.86 to 1.20)</td>
<td>1.03 (0.88 to 1.21)</td>
</tr>
</tbody>
</table>

**Strengths of this study**

This study compared different formulations of tiotropium and different categories of inhaled medications. Given that inhaled pharmacological treatment is the cornerstone of management of COPD, the question for physicians is not whether one drug should be prescribed but rather which one to choose. Our approach accordingly addressed the practical challenge and provided information for making treatment decisions. In addition, compared with previous meta-analyses,4–16 21 26–28 45 we included more trials, including a large comparative trial of inhaled long-acting bronchodilators20. This amplified the sample size of the study and enabled us to make more precise estimates. Furthermore, our subgroup and stratification analyses facilitated special populations or scenarios to be identified that require more caution when using tiotropium Soft Mist Inhaler.

**Safety of tiotropium Soft Mist Inhaler**

A recent meta-analysis reported that tiotropium Soft Mist Inhaler had an increased risk of overall death (risk ratio 1.52; 95% CI 1.06 to 2.16) and cardiovascular death (risk ratio 2.05; 95% CI 1.06 to 3.99) versus placebo.11 Our study further highlighted safety concerns of tiotropium Soft Mist Inhaler versus other active treatments. Some possible mechanisms have been proposed. One is that use of tiotropium Soft Mist Inhaler could yield higher systematic drug exposure compared with tiotropium HandiHaler,29 although this was not observed in Japanese research.30 Evidence also suggested that patients receiving a higher dose of tiotropium Soft Mist Inhaler had higher peak concentrations.29 Another mechanism is the potential effect of underlying rhythm disorders on mortality. Data suggested that the elevated risks of tiotropium Soft Mist...
Table 4  Subgroup analysis for risk of overall death and cardiovascular death for each pairwise comparison from the MTC meta-analysis

<table>
<thead>
<tr>
<th>Study duration ≥ 1 year</th>
<th>FEV&lt;sub&gt;1&lt;/sub&gt; &lt; 50% predicted value</th>
<th>Cardiovascular death (N=31)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Study duration ≥ 1 year (N=18)</td>
<td>FEV&lt;sub&gt;1&lt;/sub&gt; &lt; 50% predicted value (N=22)</td>
</tr>
<tr>
<td></td>
<td>Fixed effect OR (95% CrI)</td>
<td>Random effects OR (95% CrI)</td>
</tr>
<tr>
<td>TIO-SMI vs TIO-HH</td>
<td>1.65 (1.12 to 2.43)</td>
<td>1.65 (0.999 to 2.78)</td>
</tr>
<tr>
<td></td>
<td>1.66 (1.11 to 2.48)</td>
<td>1.66 (1.01 to 2.80)</td>
</tr>
<tr>
<td>LABA vs LABA-ICS</td>
<td>1.94 (1.30 to 2.92)</td>
<td>2.02 (1.22 to 3.47)</td>
</tr>
<tr>
<td></td>
<td>1.49 (0.999 to 2.22)</td>
<td>1.58 (0.99 to 2.79)</td>
</tr>
<tr>
<td>ICS vs PL</td>
<td>1.52 (1.06 to 2.19)</td>
<td>1.55 (1.02 to 2.47)</td>
</tr>
<tr>
<td>TIO-HH vs LABA</td>
<td>1.01 (0.84 to 1.21)</td>
<td>1.00 (0.75 to 1.35)</td>
</tr>
<tr>
<td></td>
<td>1.18 (0.97 to 1.44)</td>
<td>1.21 (0.90 to 1.70)</td>
</tr>
<tr>
<td></td>
<td>0.90 (0.74 to 1.10)</td>
<td>0.95 (0.71 to 1.42)</td>
</tr>
<tr>
<td></td>
<td>0.92 (0.81 to 1.05)</td>
<td>0.93 (0.74 to 1.24)</td>
</tr>
<tr>
<td>LABA vs LABA-ICS</td>
<td>1.17 (0.98 to 1.40)</td>
<td>1.21 (0.94 to 1.61)</td>
</tr>
<tr>
<td></td>
<td>0.90 (0.75 to 1.07)</td>
<td>0.95 (0.74 to 1.35)</td>
</tr>
<tr>
<td></td>
<td>0.92 (0.78 to 1.07)</td>
<td>0.93 (0.73 to 1.23)</td>
</tr>
<tr>
<td>LABA-ICS vs ICS</td>
<td>0.77 (0.64 to 0.92)</td>
<td>0.79 (0.60 to 1.10)</td>
</tr>
<tr>
<td></td>
<td>0.78 (0.66 to 0.93)</td>
<td>0.77 (0.59 to 1.01)</td>
</tr>
<tr>
<td>PL vs ICS</td>
<td>1.02 (0.87 to 1.20)</td>
<td>0.98 (0.73 to 1.23)</td>
</tr>
</tbody>
</table>

*N* represents the number of trials reporting on each outcome, with a study duration ≥ 1 year, or enrolling patients with mean FEV<sub>1</sub> < 50% of predicted value at baseline.

CrI, credible interval; FEV<sub>1</sub>, forced expiratory volume in 1 s; ICS, inhaled corticosteroid; LABA, long-acting β<sub>2</sub> agonist; MTC, mixed treatment comparison; PL, placebo; TIO-HH, tiotropium dry powder delivered via HandiHaler; TIO-SMI, tiotropium solution delivered via Resipmat Soft Mist Inhaler.

Inhaler compared with placebo were up to 3.2-fold for overall death and 8.6-fold for cardiovascular death in patients with cardiac rhythm disorders. However, this substantial risk was not detected in patients without cardiac dysrhythmias. Further studies, such as well designed randomised controlled trials with a head-to-head comparison of different formulations and doses of tiotropium are warranted to clarify these hypotheses.

Safety of tiotropium HandiHaler, LABA and LABA-ICS

Inhaler compared with placebo were up to 3.2-fold for overall death and 8.6-fold for cardiovascular death in patients with cardiac rhythm disorders. However, this substantial risk was not detected in patients without cardiac dysrhythmias. Further studies, such as well designed randomised controlled trials with a head-to-head comparison of different formulations and doses of tiotropium are warranted to clarify these hypotheses.

Figure 3 Stratification analysis for risk of overall death and cardiovascular death for each pairwise comparison from the MTC meta-analysis, stratified by the daily dose of tiotropium Soft Mist Inhaler. *The unit of daily dose is micrograms. CrI, credible interval; ICS, inhaled corticosteroid; LABA, long-acting β2 agonist; MTC, mixed treatment comparison; PL, placebo; TIO-HH, tiotropium dry powder delivered via HandiHaler; TIO-SMI, tiotropium solution delivered via Resipmat Soft Mist Inhaler.

Safety of tiotropium HandiHaler, LABA and LABA-ICS

Despite the potential harm of LABA among patients with asthma, we found no excess risk of LABA in patients with COPD. LABA tended to have a non-significant increased risk compared with LABA-ICS; however, the estimate was imprecise, with a likely inflated type I error due to multiple statistical tests. Meanwhile, we observed that tiotropium HandiHaler posed a nearly 80% increased risk of overall death when directly compared with LABA-ICS, although the result was based on one trial and became non-significant in the MTC analysis. A large population-based cohort study reported that tiotropium HandiHaler was associated with an excess risk of overall death versus LABA (HR 1.14; 95% CI 1.09 to 1.19). However, it was noted that more patients in the LABA group combined ICS use during the study period. Considering all the evidence, LABA-ICS may
be associated with the lowest risk profile. Further studies are needed to identify optimal therapeutic combinations and regimes for treating patients with COPD.

Limitations
Our study has limitations. First, the validity of our MTC meta-analysis relies on the assumptions of similarity of demographic characteristics across trials and homogeneity of each relative treatment effect. In our study, despite some variation, demographic characteristics were generally similar and the I² statistic was minimal. To manage possible trial heterogeneities in the MTC meta-analysis, we used the random effects model and the meta-regression and yielded similar findings. Risk estimates were also consistent between direct comparison and MTC meta-analyses in terms of directions and magnitudes. All these approaches enhanced the validity of our study. However, we cannot rule out the influences from unmeasured covariates. Given rare events and few trials for some comparisons, statistical heterogeneity or publication bias is likely underpowered. We also note that the difference in OR between direct comparison and MTC meta-analyses was particularly apparent in the comparisons of tiotropium HandiHaler versus LABA or LABA-ICS, which may be associated with the small number of trials involved in these comparisons. Second, trials were prevalent-user designs or designed for investigating ICS withdrawal effects rather than treatment effects. Notably, a high percentage of subjects combined LABA or ICS use in tiotropium trials. Our sensitivity analysis and meta-regression adjustment for percentage of concomitant use of LABA or ICS did not change the main findings. However, further studies are necessary to clarify the standalone or add-on effects of these drugs taking into account these complicated elements. Third, all included trials excluded patients with significant diseases and half of them excluded patients with specific cardiovascular morbidities. This may limit the generalisability of our findings to frailler populations in real practice. Fourth, few tiotropium Soft Mist Inhaler trials were involved in our study and precluded further exploration, such as the dose–response effect. Fifth, cardiovascular death was a sparse, non-predefined outcome and without a homogeneous definition across trials. This may result in imprecise estimates and outcome misclassification is possible. However, we constructed this composite endpoint according to the internationally recognised Medical Dictionary for Regulatory Activities definition. The analysis restricted to trials with objective adjudication of cause of death also provided similar results to the main analysis. Finally, in our included trials, the withdrawal rates were variable across treatment groups, with the lowest value in the tiotropium Soft Mist Inhaler group. This may raise concerns of overestimating the relative risk of tiotropium Soft Mist Inhaler due to underestimating the mortality of placebo and other active treatments. However, the proportions of lost to follow-up were low across treatment groups. Moreover, vital status information was ascertained in all placebo-controlled trials of tiotropium Soft Mist Inhaler, even for patients who withdrew early. Therefore, the unfavourable bias for tiotropium Soft Mist Inhaler should be limited.

Clinical implications and conclusions
In view of the safety spectrum for each inhaled medication, our results provide substantial implications for physicians. Tiotropium Soft Mist Inhaler was associated with a 50–90% increased risk of overall death and a twofold to threefold increased risk of cardiovascular death versus placebo and other inhaled medications. Until more evidence is available, physicians should use Tiotropium Soft Mist Inhaler with caution and the dose should not exceed the recommended daily dose. Alternative treatments may be considered in patients with severe COPD or with cardiac dysrhythmias.

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Contributors
Conception and design: YHD, HHL, WYS, CHC, MSL. Analysis of the data: YHD, YCW. Interpretation of the data: YHD, HHL, WYS, CHC, MSL. Drafting of the article: YHD, HHL, WYS, CHC, MSL. Statistical expertise: HHL, WYS, CHC, MSL. Obtaining funding: MSL. All the authors have read the manuscript, revised it critically for important intellectual content, and approved the final version.

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