the TONADO® studies did not show a significant difference in the hazard ratios for time to exacerbation end points. These findings are partially attributable to a higher number of severe exacerbations with T/O 5/5 µg in TONADO® 1. TONADO® was not designed for formal comparison of exacerbations with T/O versus T; however, a study powered to assess this is ongoing.

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Please refer to page A273 for declarations of interest in relation to abstract P296.

**Abstract P297**

**LUNG-FUNCTION PROFILE BEFORE AND AFTER THE FIRST MODERATE TO SEVERE EXACERBATION DURING THE WISDOM STUDY**

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**Rationale** The WISDOM study (NCT00975195) showed no increased risk of exacerbation when inhaled corticosteroid (ICS) was withdrawn stepwise in patients with severe COPD on LAMA+LABA maintenance therapy versus continued LAMA+LABA+ICS.1 Daily home spirometry measured the time course of lung-function changes throughout the study. The aim of this post hoc analysis was to address the lung-function profile leading up to, during and following the first moderate-to-severe exacerbation.

**Methods** WISDOM was a multinational, randomised, double-blind study.1 Patients with severe to very severe COPD entered a 6-week run-in with LAMA+LABA+ICS (tiotropium 18 µg once daily; salmeterol 50 µg twice daily; fluticasone propionate 500 µg twice daily), and were randomised to continue LAMA+LABA+ICS or salmeterol/tiotropium for 52 weeks while discontinuing ICS in a stepwise manner over 12 weeks. On-treatment daily forced expiratory volume in 1 second (FEV₁) change from baseline was calculated before and after the first moderate-to-severe exacerbation. In this post hoc analysis, we included patients who experienced a moderate-to-severe exacerbation after the ICS-withdrawal visit, did not have an exacerbation in the 8 weeks before or after the exacerbation, and had daily home-measured FEV₁ data available for every week analysed.

**Results** Of 2488 patients, 262 experienced a moderate-to-severe exacerbation after the ICS-withdrawal visit and had lung-function data for every week. For all patients combined (ICS and ICS withdrawal), change in FEV₁ remained relatively stable 56–14 days before the first moderate-to-severe exacerbation (mean FEV₁ change from baseline values: 0.04 to 0.07 L) (Figure). There was a decline in lung function starting 2–3 weeks before exacerbation (FEV₁ change value of 0.12 L from baseline), followed by a moderate improvement over ~14 days. Post-exacerbation lung function did not reach pre-exacerbation levels.

**Conclusions** Lung function was relatively stable in both treatment groups. Home spirometry measurements showed a marked decline in FEV₁ prior to moderate-to-severe exacerbation with improvements seen post-exacerbation, although not to pre-exacerbation levels. These findings support the usefulness of home

**Abstract P297 Figure 1** Mean change from baseline in on-treatment daily FEV₁ before and after first moderate to severe exacerbation for both treatment combined.

**Plot includes patients whose first moderate to severe exacerbation after the ICS-withdrawal visit was neither preceded by an exacerbation of any severity in 8 weeks prior nor followed by a further exacerbation of any severity within 8 weeks (56 days) after Note that Day 0 is the day of the exacerbation**
spiroscopy to predict exacerbations and to indicate subsequent worsening of lung function resulting from a previous COPD exacerbation.

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Please refer to page A273 for declarations of interest in relation to abstract P297.

REFERENCE

Rationale The combination of tiotropium (T), a long-acting muscarinic antagonist (LAMA), plus olodaterol (O), a long-acting β₂-agonist (LABA), is approved for once-daily maintenance treatment of COPD. The randomised, double-blind, Phase IIIb OTEMTO® 1 and 2 studies (NCT01431274; NCT01431287) showed improvements in quality of life and lung function after 12 weeks' treatment with T/O compared to T alone or placebo in patients with moderate to severe COPD. This post hoc analysis investigated whether previous maintenance treatment with a long-acting bronchodilator or inhaled corticosteroid (ICS) influenced symptomatic benefits of T/O.

Methods Patients aged ≥40 years received T/O 2.5/5 μg, T/O 5/5 μg, T 5 μg or placebo once daily for 12 weeks via Respimat® inhaler. St George’s Respiratory Questionnaire (SGRQ) total score was a primary end point, alongside lung function (FEV₁ area under the curve from 0–3 hours and trough FEV₁ responses). Secondary end points included Mahler Transition Dyspnoea Index (BDI) focal score. Salbutamol/albuterol was provided as rescue medication and use was recorded in an e-diary. We report comparisons between T/O 5/5 µg, T 5 µg and placebo.

Results Of the 1621 patients evaluated, 943 had received prior maintenance treatment (66.7% LABA; 59.4% LAMA; 64.5% ICS) and 678 had not. Similar improvements in mean SGRQ total score were observed with T/O compared to T and placebo, respectively, in patients receiving prior maintenance treatment (–2.02 and –4.39 units) and those without (–2.20 and –4.78 units) (Table). TDI focal scores improved with T/O compared to T and placebo, respectively, in patients receiving prior maintenance treatment (0.60 and 1.87 units) and those without (0.60 and 1.33 units) (Table). Patients with and without prior maintenance treatment demonstrated similar improvements in daytime and night-time rescue medication use and lung-function improvements with T/O compared to T and placebo.

Conclusions T/O provides symptomatic benefits as demonstrated by improvements in SGRQ score, TDI focal score and decreased rescue medication use compared to placebo and T, independent of previous maintenance treatment. These findings suggest T/O is beneficial over monotherapy when used as first COPD maintenance treatment.

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Please refer to page A273 for declarations of interest in relation to abstract P298.