Efficacy and safety of fluticasone furoate/vilanterol or tiotropium in subjects with COPD at cardiovascular risk

Henry Covelli¹
Bonavuth Pek²
Isabelle Schenkenberger³
Catherine Scott-Wilson⁴
Amanda Emmett⁵
Courtney Crim⁶

¹Kootenai Health, Coeur d’Alene, ID, USA; ²Clinique de Pneumologie et de Sommeil de Lanaudière, Quebec, Canada; ³Klinische Forschung, Berlin, Germany; ⁴GlaxoSmithKline Inc., Research Triangle Park, ⁵PAREXEL International, Durham, NC, USA

Background: Fluticasone furoate/vilanterol (FF/VI) is a novel, once-daily, inhaled corticosteroid/long-acting β₂-agonist combination approved for the treatment of COPD and asthma. We compared the safety and efficacy of FF/VI and tiotropium (TIO) in subjects with moderate-to-severe COPD with greater risk for comorbid cardiovascular disease (CVD).

Methods: This randomized, blinded, double-dummy, parallel-group study compared a once-daily morning dose of FF/VI 100/25 mcg delivered via ELLIPTA® with TIO 18 mcg via HandiHaler® for 12 weeks in subjects with diagnosed COPD, forced expiratory volume in 1 second (FEV₁) 30%–70% predicted, and CVD or CVD risk. The primary endpoint was change from baseline in 24-hour weighted mean FEV₁ on Day 84. Other efficacy endpoints included time to onset of bronchodilation, trough FEV₁, other spirometry measures, rescue medication use, symptoms, quality of life (St George’s Respiratory Questionnaire-COPD [SGRQ-C]), and health status (COPD Assessment Tests [CAT]) measures. Safety endpoints included cardiovascular monitoring, cortisol excretion, COPD exacerbations, and adverse events, including prespecified drug effects.

Results: Both FF/VI and TIO improved the 24-hour weighted mean FEV₁ from baseline after 12 weeks with no significant difference between treatments. Other endpoints favored FF/VI for time to onset of bronchodilation, rescue medication use, dyspnea, SGRQ-C and CAT scores, or favored TIO for change from baseline in forced vital capacity and inspiratory capacity. Pneumonia occurred more frequently in the FF/VI group, and two TIO-treated subjects died following cardiovascular events. Other safety measures were similar between groups, and cardiovascular monitoring did not reveal increased CVD risk.

Conclusion: Both FF/VI and TIO were efficacious in improving lung function in subjects with COPD and comorbid CVD or CVD risk factors, with minor differences in efficacy and safety profiles.

Keywords: fluticasone furoate/vilanterol, tiotropium, COPD, cardiovascular disease, ICS/LABA, anticholinergic

Introduction

Successful management of moderate-to-severe COPD can be accomplished with combination therapy that delivers a concurrent dose of inhaled corticosteroid and a long-acting β₂-agonist (ICS/LABA) or with tiotropium (TIO), an inhaled long-acting anticholinergic bronchodilator.¹ Both twice-daily ICS/LABA and once-daily TIO treatment for COPD have been demonstrated to improve lung function and health status as well as reduce COPD exacerbations.¹–⁶ Head-to-head comparisons between ICS/LABA drugs and TIO, however, are limited and the efficacy results inconclusive,⁷,⁸ leading researchers to call for further large, long-term randomized controlled trials to...
establish relative efficacy and safety of these treatments, as well as compare newly developed alternative ICS/LABAs with TIO. A newly introduced ICS/LABA treatment for COPD in a novel inhaler device combines fluticasone furoate (FF), an ICS with enhanced selectivity, retention, and affinity for the glucocorticoid receptor, and vilanterol (VI), a LABA with prolonged bronchodilating properties. FF/VI is the only once-daily ICS/LABA treatment and is well tolerated, improves lung function, and sustains bronchodilation after 24 weeks and 52 weeks of treatment for COPD. When once-daily treatment with FF/VI for 12 weeks was compared with twice-daily fluticasone propionate/salmeterol 250/50 (FP/SALM) for COPD, there was evidence of numerically greater effect on the forced expiratory volume in 1 second (FEV₁) and faster onset for FF/VI compared with FP/SALM with similar safety profiles for the two drugs.

COPD is not only a disease of the Airways and lung parenchyma, but there are several extrapulmonary comorbidities associated with COPD, of which cardiovascular disease (CVD) is arguably the most significant. CVD is a comorbidity of COPD, and the risk of cardiovascular (CV) events rises in subjects with COPD with a history of myocardial infarction. Furthermore, concerns regarding CV safety and COPD medications have been raised for both TIO and LABA drugs. Therefore, we conducted the study in a population of subjects with or at risk for CVD. The objective of this study was to compare the efficacy and safety of FF/VI and TIO, both delivered once daily, for COPD treatment over a 12-week period in a randomized, controlled trial. An additional objective was to investigate whether the efficacy and/or safety profile of FF/VI or TIO would differ in subjects with CVD or in subjects at risk for CVD.

Methods

The study (Protocol HZC115805, NCT01627327) was approved by an institutional review board or independent ethics committee at each site and conducted in accordance with good clinical practice guidelines and the Declaration of Helsinki. All subjects provided written informed consent. The study was conducted between April and December 2012 at 56 centers in six countries (Canada, Czech Republic, Germany, Poland, Romania, and the USA).

Subjects

Men and women aged ≥40 years with a clinical diagnosis of COPD were recruited. All subjects were current or former smokers (>10 pack years) and had a postbronchodilator FEV₁ ≥30% to ≤70% of predicted normal (NHANES III) and an FEV₁/forced vital capacity (FVC) ratio ≤70% at screening. Subjects also had a history of CVD/a CVD event or, in addition to being a current/former smoker, had at least one current CV risk factor (hypertension, hypercholesterolemia, or treated diabetes). Female subjects were either postmenopausal or using effective contraception.

Subjects were excluded if they had asthma or respiratory disorders other than COPD; recent (<12 months) lung resection; clinically significant abnormal chest X-ray, laboratory, Holter, or electrocardiogram (ECG) finding at screening; recent (<12 weeks) hospitalization for COPD; recent (<6 weeks) acute worsening of COPD requiring treatment with corticosteroids or antibiotics or lower respiratory tract infection requiring antibiotics; noncompliance, COPD exacerbation, or lower respiratory tract infection during the run-in period; cancer not in complete remission for ≥5 years; long-term or nocturnal oxygen therapy (>12 h/day); or other diseases or contraindications that would put the subject at risk.

Study design and treatment

This study was a 12-week, randomized (1:1), blinded, stratified, double-dummy, parallel-group, active comparator study. After providing written informed consent and prior to screening, subjects discontinued COPD medications (including maintenance medications; excluded medications included systemic/oral/parenteral corticosteroids [within 6 weeks prior to screening and thereafter], ICS or ICS/LABA combinations [within 4 weeks prior to screening and thereafter], anticholinergics [long acting, 1 week prior to screening and thereafter; short acting, 4 hours prior to screening and thereafter], oral phosphodiesterase type 4 inhibitors [PDE-4] inhibitors [1 week prior to screening and thereafter], and LABAs [within 48 hours prior to screening and thereafter]); exceptions were study-supplied rescue medication (albuterol) and mucolytics at a constant dosage. After the screening visit, eligible subjects entered a 2-week single-blind placebo run-in period to obtain baseline rescue medication use and COPD symptom (scores recorded on diary cards) and to evaluate subjects’ compliance and disease stability. At the end of the run-in period, eligible subjects were randomly assigned to once-daily morning treatment with either FF/VI 100/25 mcg (BREO™ [RELVAR™, REVINTY™], GlaxoSmithKline plc [GSK], London, UK) delivered via the ELLIPTA™ inhaler (GSK, London, UK) or TIO 18 mcg (Spiriva®, Boehringer Ingelheim, Ingelheim, Germany) delivered from blister capsules by the HandiHaler® (Spiriva, Boehringer Ingelheim). When used, the novel ELLIPTA dry powder inhaler simultaneously delivers...
one breath-actuated dose of both FF 100 mcg and VI 25 mcg from their individual blister strips.

A central randomization schedule was generated using a validated computerized system (RandAll; GSK) and communicated with a validated computerized voice response system, the Registration and Medication Ordering System (RAMOS; GSK), when the investigative site called to randomize a subject. To provide a double-dummy design, subjects were given two inhalers, one containing active drug and the other placebo, and they and their investigators were blinded to assigned treatment. TIO and placebo capsules were closely matched in color, but TIO capsules had trade markings that were not present on the placebo capsules. Whether subjects would notice and correctly or incorrectly interpret this difference is unclear. Capsule type was consistent for each subject for the duration of the study. Both the TIO and placebo blister packages were covered with opaque overlabels that hide the information on the TIO packaging. HandiHalers were covered with labels to mask identifying marks. Staff involved with safety and efficacy assessments were not present during dosing in the clinic so that they would not identify the capsules removed from the blisters.

Every morning, subjects in the FF/VI group inhaled once from the ELLIPTA inhaler containing active drug followed by two inhalations from the HandiHaler containing placebo. Subjects in the TIO group inhaled once from the ELLIPTA inhaler containing placebo followed by two inhalations from the HandiHaler containing TIO. Compliance with the study drug was assessed at each visit by reviewing the dose counter on the ELLIPTA or counting capsules from the HandiHaler. Subjects were also provided with albuterol (metered dose inhaler [MDI] and/or nebulus) to use as needed for symptomatic relief of COPD during the run-in and treatment periods, except for the 4-hour period prior to spirometry and during each study visit.

Subjects were stratified at randomization for COPD exacerbation history (either did or did not have ≥1 COPD exacerbation requiring oral corticosteroid and/or antibiotic treatment and/or hospitalization in the 3 years prior to screening) and for reversibility status (either reversible [defined as an increase in FEV₁ of ≥12% and ≥200 mL] or nonreversible [defined as an increase in FEV₁ <200 mL or ≥200 mL and <12% after albuterol administration at screening]). Subjects revisited the clinic on treatment days 28, 56, and 84. A follow-up safety telephone contact was performed 7 days after the last treatment visit. Subjects who experienced a COPD exacerbation or pneumonia were withdrawn; however, only exacerbations that resulted in hospitalization (ie, severe exacerbations) were captured as an adverse event (AE).

Study assessments

Standardized equipment (spirometry: microQuark, 12-lead ECG: Universal ECG, and 24-hour Holter monitor: Mortara Instrument), study-customized software, and training were provided, and data were transmitted and overread by independent centralized certified staff at CompleWare.

At the screening visit, pre- and postbronchodilator spirometry, medical history, physical examination, 12-lead ECG, 24-hour Holter monitoring (subset of subjects), chest X-ray, vital signs, serum pregnancy tests, clinical laboratory tests, oropharyngeal examination, and 24-hour urine cortisol collection (subset of subjects) were conducted, and diary cards were dispensed for recording of medical problems, concomitant medications, rescue medication use, and symptoms (dyspnea, cough, sputum production). At all subsequent visits, eligible randomized subjects completed the St George’s Respiratory Questionnaire-COPD (SGRQ-C) and COPD Assessment Tests (CAT) prior to other assessments, and safety monitoring was conducted, including oropharyngeal examination, AE reporting, exacerbation and pneumonia assessment, and diary card review. At the randomization Visit 2, subjects also underwent vital sign measurement and blood draws for biomarkers (subset of subjects). At Visit 5, which was an overnight stay, 24-hour spirometry, ECGs, 24-hour Holter monitoring (subset of subjects), 24-hour urinary cortisol collection (subset of subjects), physical examinations, vital sign measurement, blood draws for clinical laboratory tests and biomarkers, and pregnancy tests were conducted. Each Holter was reviewed by a centralized overreader at CompleWare. Serum was collected for seven known inflammatory biomarkers at baseline and Week 12 and analyzed as exploratory endpoints: high-sensitivity C-reactive protein (hsCRP), pulmonary and activation-regulated chemokine (more currently referred to as chemokine [C-C motif] ligand 18 [CCL-18]), fibrinogen, interleukin-6 (IL-6), surfactant protein D (SP-D), Clara cell secretory protein-16 (CC-16), and brain natriuretic peptide (BNP).

Spirometry was performed after withholding rescue medication, study medication, and caffeinated drinks and after abstinence from smoking and physical exertion. Inspiratory capacity (IC) assessments were performed prior to FEV₁ assessments. At the randomization Visit 2, after predose spirometry and the first study dose, subjects underwent postdose serial spirometry at 5 minutes, 15 minutes, and 30 minutes and at 1 hour, 2 hours, and 4 hours. At Visit 5 on Day 84 after predose spirometry and after the last study dose, subjects underwent postdose serial spirometry at 5 minutes, 15 minutes, and 30 minutes and at 1 hour, 2 hours,
4 hours, 6 hours, 8 hours, 12 hours, 13 hours, 14 hours, 16 hours, 20 hours, and 24 hours.

**Statistical analysis**

Assuming a standard deviation of 215 mL and significance at the two-sided 5% level, a sample size of 271 subjects per treatment arm was required for 90% power to detect a 60 mL difference between FF/VI and TIO for 24-hour weighted-mean FEV$_1$, at Week 12, an estimate based on historical data. To allow for a 10% withdrawal rate, at least 300 subjects were to be randomized per treatment arm. Primary analyses were performed on the intent-to-treat population, which comprised all subjects randomized to treatment and who had received at least one dose of study medication.

The primary endpoint, change from baseline in 24-hour weighted mean FEV$_1$, at Week 12, was analyzed using analysis of covariance with covariates of baseline FEV$_1$, randomization stratum (COPD exacerbation history and reversibility status), smoking status at screening, and country. For calculation of 24-hour weighted mean FEV$_1$, missing serial spirometry FEV$_1$ values were linearly interpolated between two nonmissing values; however, the 0-hour and 24-hour values, as well as additional values between 0 hour and 6 hours and between 12 hours and 24 hours, must not have been missing.

The secondary endpoints (time to onset [increase of 100 mL above baseline in FEV$_1$] 0–4 hours postdose on Day 1 and change from baseline in trough FEV$_1$ on Day 84) were nested under the primary endpoint. To account for multiplicity across primary and secondary efficacy endpoints, a $P$-value <0.05 was required for the primary endpoint to allow statistical significance to be inferred for secondary endpoints. Time to onset was analyzed using a log-rank test, stratified by exacerbation history and reversibility stratum. Change from baseline in trough (predose) FEV$_1$ was analyzed using the same methodology as the primary endpoint.

Daily diary measures were averaged weekly and over the entire 12-week treatment period for analysis. Change from baseline in SGRQ-C and CAT scores was analyzed using mixed model repeated measures with covariates of baseline score, exacerbation history and reversibility stratum, smoking status, country, treatment group, day, day by baseline, and day by treatment group. A responder analysis was also performed for the SGRQ-C. Subjects were classified as a responder at a given visit if they had a change from baseline at that visit of $\geq$4 points improvement, the minimal clinically important difference (MCID). Otherwise, the subject was classified as a nonresponder. A logistic regression model with terms for treatment group, exacerbation history and reversibility stratum, and baseline score was used to compare the treatment groups. Other efficacy endpoints, as well as safety measures of pulse rate and log-transformed urinary cortisol, were analyzed using analysis of covariance.

**Results**

**Subject characteristics**

At screening, 890 subjects were evaluated for eligibility (Figure 1). Of those, 267 (30%) subjects were not randomized mainly because they failed the strict enrollment criteria, for example, FEV$_1$, 30%–70% predicted (n=112), and/or had an abnormal Holter (n=75) and/or ECG (n=55) finding. Following the run-in period, 310 and 313 subjects were randomized to treatment with FF/VI and TIO, respectively, and constituted the intent-to-treat population. Of those, 291 (94%) and 274 (88%) subjects, respectively, completed the study. More subjects were withdrawn from treatment with TIO, mainly due to AEs and lack of efficacy/COPD exacerbations, than with FF/VI. Three percent of TIO-treated and 1% of FF/VI-treated subjects withdrew from the study due to protocol deviations.

Subjects’ demographic and baseline characteristics were similar between treatment groups (Table 1), with the majority of subjects being white and male. In this population of subjects with a history of CVD/CV event or risk factors, the most common baseline conditions were hypertension and hypercholesterolemia. Slightly under half of the subjects had not had a COPD exacerbation in the last 3 years. Baseline lung function results indicated that the study population had moderate-to-severe airflow limitation (Table 1), and approximately one-third of the subjects demonstrated acute bronchodilator reversibility. Baseline health-related quality of life (QoL) and health status, indicated by SGRQ-C and CAT scores, respectively, suggested impairment and a high level of COPD symptoms.

**Efficacy**

Results of the primary efficacy measure, change from baseline trough in 24-hour weighted mean FEV$_1$ after 12 weeks’ treatment, were similar between the treatment groups (Figure 2), with a least squares (LS) mean change from baseline of 117 mL and 95 mL in the FF/VI and TIO groups, respectively, with a difference of 0.022 L (95% confidence interval [CI] −0.012 to 0.055) ($P$=0.201). In the reversible subgroup, however, the mean change was greater in the FF/VI group compared with the TIO group (181 mL and 110 mL, respectively), with a difference of 0.071 L.
Efficacy and safety of FF/VI or TIO in subjects with COPD at CV risk

Figure 1 Patient disposition.

Abbreviations: FF/VI, fluticasone furoate/vilanterol (100/25 mcg); TIO, tiotropium (18 mcg).

Table 1 Summary of demographic and baseline characteristics

<table>
<thead>
<tr>
<th></th>
<th>FF/VI (n=310)</th>
<th>TIO (n=313)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age, years</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>62.9 (8.1)</td>
<td>62.3 (8.0)</td>
</tr>
<tr>
<td>Min–max</td>
<td>42–85</td>
<td>40–86</td>
</tr>
<tr>
<td><strong>Sex, n (%), female/male</strong></td>
<td>117/193 (38/62)</td>
<td>104/209 (33/67)</td>
</tr>
<tr>
<td><strong>Race and racial combinations, n (%)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>White – White/Caucasian/European heritage</td>
<td>302 (97)</td>
<td>305 (97)</td>
</tr>
<tr>
<td>African American/African heritage</td>
<td>8 (3)</td>
<td>7 (2)</td>
</tr>
<tr>
<td>Asian – East Asian heritage</td>
<td>0</td>
<td>1 (&lt;1)</td>
</tr>
<tr>
<td><strong>Body mass index, kg/m²</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>28.4 (6.0)</td>
<td>28.6 (5.5)</td>
</tr>
<tr>
<td>Min–max</td>
<td>16–49</td>
<td>16–62</td>
</tr>
<tr>
<td><strong>Smoking history</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Current/former smoker, n (%)</td>
<td>161/149 (52/48)</td>
<td>161/152 (51/49)</td>
</tr>
<tr>
<td>Pack years, mean (SD)</td>
<td>43.2 (24.5)</td>
<td>45.6 (25.2)</td>
</tr>
<tr>
<td><strong>Cardiovascular history or risk factors, n (%)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hypertension</td>
<td>251 (81)</td>
<td>271 (87)</td>
</tr>
<tr>
<td>Hypercholesterolemia</td>
<td>134 (43)</td>
<td>143 (46)</td>
</tr>
<tr>
<td>Coronary artery disease</td>
<td>84 (27)</td>
<td>81 (26)</td>
</tr>
<tr>
<td>Diabetes</td>
<td>54 (17)</td>
<td>71 (23)</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td>32 (10)</td>
<td>24 (8)</td>
</tr>
<tr>
<td>Arrhythmia</td>
<td>16 (5)</td>
<td>27 (9)</td>
</tr>
<tr>
<td>Congestive heart failure</td>
<td>14 (5)</td>
<td>15 (5)</td>
</tr>
<tr>
<td>Cerebrovascular accident</td>
<td>12 (4)</td>
<td>4 (1)</td>
</tr>
<tr>
<td><strong>COPD exacerbation history, n (%)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No exacerbation in past 3 years</td>
<td>145 (47)</td>
<td>136 (43)</td>
</tr>
<tr>
<td>≥1 exacerbations in past 3 years</td>
<td>165 (53)</td>
<td>177 (57)</td>
</tr>
</tbody>
</table>

(Continued)
Table 1 (Continued)  

<table>
<thead>
<tr>
<th>FF/VI (n=310)</th>
<th>TIO (n=313)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline lung function, mean (SD)</td>
<td></td>
</tr>
<tr>
<td>Prebronchodilator FEV₁ (L), mean (SD)</td>
<td>1.35 (0.47)</td>
</tr>
<tr>
<td>Prebronchodilator FVC (L), mean (SD)</td>
<td>2.67 (0.78)</td>
</tr>
<tr>
<td>Inspiratory capacity (L), mean (SD)</td>
<td>2.06 (0.60)</td>
</tr>
<tr>
<td>Reversibility, n (%), reversible/non-reversible</td>
<td>101/202 (33/67)</td>
</tr>
<tr>
<td>Postbronchodilator FEV₁ (L), mean (SD)</td>
<td>1.46 (0.45)</td>
</tr>
<tr>
<td>Postbronchodilator % predicted normal FEV₁, mean (SD)</td>
<td>49.4 (10.5)</td>
</tr>
<tr>
<td>Postbronchodilator FVC (L), mean (SD)</td>
<td>2.90 (0.80)</td>
</tr>
<tr>
<td>Postbronchodilator FEV₁/FVC</td>
<td>0.51 (0.10)</td>
</tr>
<tr>
<td>Baseline health outcome scores, mean (SD)</td>
<td></td>
</tr>
<tr>
<td>SGRQ-C</td>
<td>48.84 (16.70)</td>
</tr>
<tr>
<td>CAT</td>
<td>18.1 (6.9)</td>
</tr>
</tbody>
</table>

Abbreviations: CAT, COPD Assessment Test; FEV₁, forced expiratory volume in 1 second; FF/VI, fluticasone furoate/vilanterol (100/25 mcg); FVC, forced vital capacity; SD, standard deviation; SGRQ-C, St George’s Respiratory Questionnaire-COPD; TIO, tiotropium (18 mcg).

(95% CI 0.012–0.129). Other stratification subgroups (ie, nonreversible, ≥1 COPD exacerbation in last 3 years, no COPD exacerbation in last 3 years) demonstrated small or no treatment group differences.

The secondary endpoints were nested under the primary endpoint. Because the primary endpoint difference did not reach statistical significance, all differences reported for the secondary efficacy endpoints are presented descriptively only as statistical inference cannot be made.

Table 2 shows that 79% and 82% of subjects receiving FF/VI or TIO, respectively, achieved an increase in FEV₁ of ≥100 mL from baseline within 4 hours postdose on treatment Day 1, with a somewhat faster onset in the FF/VI group, with the median time to onset of 17.0 minutes compared with the median time of 20.5 minutes in the TIO group. More subjects in the FF/VI group than the TIO group demonstrated an onset of effect within the first 5 minutes of dosing (Table 2).

Other efficacy endpoints are summarized in Table 3. FF/VI and TIO improved trough FEV₁ on Day 84 to a similar degree. Both change from baseline in trough IC and change from baseline in trough FVC favored TIO treatment.

Subjects reported their COPD symptoms daily on diary cards. Changes from baseline in the individual symptoms of cough and sputum production were similar between groups (Table 3). However, reductions in dyspnea over the 12-week period favored FF/VI.

Treatment with FF/VI reduced subjects’ use of rescue medication during Weeks 1–12, with an LS mean change difference of −0.37 inhalations (95% CI −0.55 to −0.19) and during each of the week-long treatment periods (Figure 3A). Likewise, the percentage of rescue-free 24-hour periods was increased by FF/VI treatment compared with TIO during Weeks 1–12, with an LS mean change difference of 9.1% (95% CI 4.0–14.2) and during each week (Figure 3B).

Health-related QoL was measured at baseline and at each posttreatment visit. While SGRQ-C baseline scores were similar between treatment groups (Table 1) and both groups improved with treatment (Figure 4A), FF/VI demonstrated a consistently greater numerical improvement in total score than TIO treatment, and only FF/VI treatment reached the clinically

Table 2 Time to onset of action (postdose change from baseline of ≥100 mL) on treatment day 1

<table>
<thead>
<tr>
<th>Time to onset</th>
<th>FF/VI (n (%) of subjects)</th>
<th>TIO (n (%) of subjects)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>304</td>
<td>309</td>
</tr>
<tr>
<td>5 minutes</td>
<td>110 (36)</td>
<td>72 (23)</td>
</tr>
<tr>
<td>15 minutes</td>
<td>37 (12)</td>
<td>66 (21)</td>
</tr>
<tr>
<td>30 minutes</td>
<td>25 (8)</td>
<td>38 (12)</td>
</tr>
<tr>
<td>60 minutes</td>
<td>23 (8)</td>
<td>30 (10)</td>
</tr>
<tr>
<td>120 minutes</td>
<td>23 (8)</td>
<td>32 (10)</td>
</tr>
<tr>
<td>240 minutes</td>
<td>21 (7)</td>
<td>16 (5)</td>
</tr>
<tr>
<td>Increase of ≥100 mL not met</td>
<td>65 (21)</td>
<td>55 (18)</td>
</tr>
</tbody>
</table>

Abbreviations: FF/VI, fluticasone furoate/vilanterol (100/25 mcg); TIO, tiotropium (18 mcg).
Table 3 Other efficacy endpoints

<table>
<thead>
<tr>
<th>Endpoint Description</th>
<th>FF/VI (LS mean change difference (95% CI))</th>
<th>TIO (LS mean change difference (95% CI))</th>
<th>LS mean change difference (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trough (predose) FEV(_1), n</td>
<td>268 (0.98)</td>
<td>209 (0.83)</td>
<td>-0.09 (0.02 to 0.03)</td>
</tr>
<tr>
<td>Trough (predose) IC, n</td>
<td>267 (0.09)</td>
<td>163 (0.07)</td>
<td>-0.005 (0.002 to 0.003)</td>
</tr>
<tr>
<td>Trough (predose) FVC, n</td>
<td>268 (0.05)</td>
<td>209 (0.04)</td>
<td>-0.08 (0.002 to 0.003)</td>
</tr>
<tr>
<td>Dyspnea symptom score, a</td>
<td>301 (2.3)</td>
<td>298 (3.0)</td>
<td>2.9 (0.002 to 3.4)</td>
</tr>
<tr>
<td>Cough symptom score, b</td>
<td>299 (1.32)</td>
<td>296 (1.32)</td>
<td>1.5 (0.03 to 1.4)</td>
</tr>
<tr>
<td>Sputum production symptom score, b</td>
<td>297 (2.3)</td>
<td>296 (2.3)</td>
<td>1.5 (0.3 to 1.4)</td>
</tr>
</tbody>
</table>

Notes: aScored from 0 (not breathless at rest or upon exertion) to 4 (breathless at rest); bScored from 0 (none) to 3 (severe); 24h period.

Abbreviations: CI, confidence interval; FEV\(_1\), forced expiratory volume in 1 second; FF/VI, fluticasone furoate/vilanterol (100/25 mcg); FVC, forced vital capacity; h, hours; h, hours; IC, inspiratory capacity; LS, least squares; SD, standard deviation; SE, standard error; TIO, tiotropium (18 mcg).

Figure 3 Weekly rescue medication use.

Notes: (A) Mean weekly use of albuterol/salmeterol rescue medication in inhalations/day over the 12-week course of the study and (B) mean weekly percentage of rescue-free days over the 12-week course of the study. During Weeks 1–12, the LS mean change difference between groups was (A) -0.37 inhalations (95% CI -0.55 to -0.19) and (B) 9.1% (95% CI 4.0–14.2). Albuterol use was measured as occasions used/day.

Abbreviations: CI, confidence interval; FF/VI, fluticasone furoate/vilanterol (100/25 mcg); LS, least squares; TIO, tiotropium (18 mcg).
significant MCID change score of −4 (Week 8); achievement of the total score MCID was not persistent at Week 12. With respect to individual SGRQ-C components (symptoms, impacts, activity), FF/VI consistently reduced each component score more than TIO, and only the FF/VI group achieved the MCID for symptoms across all time points. A responder analysis for total SGRQ-C score revealed that a higher percentage of FF/VI-treated than TIO-treated subjects, respectively, achieved the MCID after 4 weeks (46% vs 34%), 8 weeks (49% vs 36%), and 12 weeks (48% vs 42%) of treatment.

CAT scores at baseline were high (≥10) and similar between groups (Table 1). The FF/VI group consistently demonstrated a numerically greater improvement in CAT score over the treatment period (Figure 4B).

Biomarkers

Seven biomarkers (hsCRP, CCL-18, SP-D, fibrinogen, IL-6, CC-16, and BNP) were measured in a subset of subjects (165 FF/VI, 169 TIO) at baseline and Week 12. There were no significant changes from baseline in BNP, fibrinogen, or IL-6 with either FF/VI or TIO. For both the treatment groups, there were decreases in CCL-18 and CC-16 and an increase in SP-D; hsCRP only increased at Week 12 in the TIO group. However, for all seven biomarkers, the change from baseline was not significantly different between the two treatment groups (data not shown).

Safety

Over one-third of the subjects experienced an AE during treatment (Table 4), the most frequent of which were headache, nasopharyngitis, back pain, and oral candidiasis. More TIO-treated than FF/VI-treated subjects were withdrawn from the study due to AEs. Two TIO-treated subjects died (one due to cardiopulmonary arrest and the other due to cardiorespiratory arrest and cardiac failure). Both subjects had hypertension; one smoked and had a family history of CVD. More FF/VI than TIO-treated subjects experienced an AE considered drug related by the investigator.

Table 4 Adverse events

<table>
<thead>
<tr>
<th>Category</th>
<th>FF/VI (n=310)</th>
<th>TIO (n=313)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AEs during treatment, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any AE during treatment</td>
<td>113 (36)</td>
<td>99 (32)</td>
</tr>
<tr>
<td>Drug-related AE</td>
<td>21 (7)</td>
<td>12 (4)</td>
</tr>
<tr>
<td>AE leading to withdrawal^b</td>
<td>6 (2)</td>
<td>14 (4)</td>
</tr>
<tr>
<td>Serious AEs</td>
<td>10 (3)</td>
<td>10 (3)</td>
</tr>
<tr>
<td>Fatal AEs</td>
<td>0</td>
<td>2 (&lt;1)</td>
</tr>
<tr>
<td>Most frequent^c AEs, n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headache</td>
<td>18 (6)</td>
<td>23 (7)</td>
</tr>
<tr>
<td>Nasopharyngitis</td>
<td>16 (5)</td>
<td>13 (4)</td>
</tr>
<tr>
<td>Back pain</td>
<td>9 (3)</td>
<td>9 (3)</td>
</tr>
<tr>
<td>Oral candidiasial</td>
<td>9 (3)</td>
<td>5 (2)</td>
</tr>
<tr>
<td>AEs of special interest,^d n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cardiovascular effects</td>
<td>13 (4)</td>
<td>15 (5)</td>
</tr>
<tr>
<td>Local steroid effects/candidiasis</td>
<td>17 (5)</td>
<td>11 (4)</td>
</tr>
<tr>
<td>Hypersensitivity</td>
<td>5 (2)</td>
<td>4 (1)</td>
</tr>
<tr>
<td>LRTI excluding pneumonia</td>
<td>3 (&lt;1)</td>
<td>4 (1)</td>
</tr>
<tr>
<td>Bone disorders/fractures</td>
<td>3 (&lt;1)</td>
<td>1 (&lt;1)</td>
</tr>
<tr>
<td>Pneumonia</td>
<td>3 (&lt;1)</td>
<td>0</td>
</tr>
<tr>
<td>Ocular effects/glaucoma</td>
<td>0</td>
<td>1 (&lt;1)</td>
</tr>
</tbody>
</table>

Notes: *Number of subjects reporting an event (not number of events); ^any AE leading to permanent withdrawal from the study or withdrawal of study drug; ^AEs reported in ≥3% of subjects in either treatment group; ^prespecified AEs of special interest with corticosteroid and LABA treatment.

Abbreviations: AE, adverse event; FF/VI, fluticasone furoate/vilanterol (100/25 mcg); LRTI, lower respiratory tract infection; TIO, tiotropium (18 mcg).

Figure 4 Least squares mean change from baseline in (A) total SGRQ-C score and (B) total CAT score at Weeks 4, 8, and 12.

Notes: The LS mean change difference between groups for Week 4 was (A) −2.25 (95% CI −4.00 to −0.51) and (B) −0.9 (95% CI −1.8 to 0.0), for Week 8 was (A) −2.84 (95% CI −4.70 to −0.99) and (B) −0.7 (95% CI −1.6 to 0.2), and for Week 12 was (A) −1.38 (95% CI −3.38 to 0.62) and (B) −0.4 (95% CI −1.3 to 0.5). The MCID of −4 is indicated by a dotted line.

Abbreviations: CAT, COPD Assessment Test; CI, confidence interval; FF/VI, fluticasone furoate/vilanterol (100/25 mcg); LS, least squares; MCID, minimal clinically important difference; SGRQ-C, St George’s Respiratory Questionnaire-COPD; TIO, tiotropium (18 mcg).
Pneumonia was reported in three subjects during treatment with FF/VI, all of which were radiologically confirmed and treated with antibiotics. Two of the three subjects were hospitalized. The third subject experienced a COPD exacerbation 4 days after developing pneumonia. All three subjects were withdrawn from the study. One FF/VI-treated subject developed pneumonia 19 days after study medication (FF/VI) was finished. None of the pneumonias were fatal.

AEs of special interest based on the known effects of corticosteroid and LABA treatment were prespecified. The most commonly reported were CV effects and local steroid effects (Table 4). The number of other events was low and similar between groups.

Fewer subjects in the FF/VI group (7/310; 2%) experienced a COPD exacerbation than in the TIO group (11/313; 4%), and all were withdrawn from the study. Although the protocol required subjects to be withdrawn upon experiencing an exacerbation, one subject in the FF/VI group was not withdrawn until after having a second event. Of those withdrawn, five FF/VI-treated and nine TIO-treated subjects had a history of COPD exacerbation. None of the COPD exacerbations were fatal. Three of eight exacerbations in the FF/VI group required hospitalization compared with one of eleven in the TIO group; per protocol, only the hospitalized events were captured as adverse events. Compared with the FF/VI group, more subjects in the TIO group who had an exacerbation were treated with antibiotics alone or in conjunction with oral corticosteroids.

There were no clinically significant differences between groups in mean changes from baseline for pulse rate, heart rate, or QTc intervals. At Week 12, few subjects (2% in each group) had a QTc(F) >450, and few (≤1% in each group) had a change from baseline in QTc(F) ≥60 ms.

The percentage of subjects with ECG findings defined a priori as abnormalities of potential clinical importance was similar at baseline (9% FF/VI group, 13% TIO group) and at the end of treatment (8% FF/VI group, 7% TIO group). Most abnormalities were categorized as depolarization abnormalities or prior myocardial infarction.

Holter monitoring was conducted in a subset of subjects (158 FF/VI, 153 TIO), with no significant difference between FF/VI and TIO groups in abnormalities of potential clinical significance, including mean heart rate, cardiac rhythm abnormalities (ventricular ectopic events), or other prespecified abnormalities of potential clinical importance. Most of the abnormalities were nonsustained ventricular tachycardias (defined as >100 beats/min, 3–30 beats) or right bundle branch blocks. There was a higher incidence of nonsustained ventricular tachycardia in the FF/VI group (9%) than the TIO group (4%) and of right bundle branch block in the TIO group (5%) than the FF/VI group (2%). There were no reports of sustained ventricular tachycardia or sustained supraventricular tachycardia (defined as ≥100 beats/min, >30 beats) at any time postrandomization in either group. Abnormalities of potential clinical importance were reported in more subjects at the end of treatment (14% FF/VI, 12% TIO) than at screening (4% FF/VI, 5% TIO). More subjects in both groups had ventricular ectopics with ≥1 ventricular run at the end of treatment (11% FF/VI, 6% TIO) than at screening (<1% FF/VI, 2% TIO).

Clinical chemistry and hematology results were generally similar between groups, and mean changes from baseline were minimal. No subject had laboratory abnormalities of potential clinical concern. Cortisol excretion was measured by 24-hour urine collection in a subset of subjects (104 on FF/VI, 103 on TIO). There were no clinically significant differences in LS mean urinary cortisol ratio of Week 12 to baseline between groups (FF/VI [1.06] vs TIO [1.02]) ratio of 1.038; 95% CI 0.86–1.26).

**Discussion**

Our results in this head-to-head comparator trial between FF/VI and TIO in subjects with COPD who had or were at risk of CVD showed that both treatments resulted in lung function improvement >12 weeks and were consistent with previous reports of FF/VI treatment effects from randomized controlled trials.\(^{11-13,22-24}\) The change from baseline in 24-hour weighted mean FEV\(_1\) after 12-week treatment (117 mL for FF/VI and 95 mL for TIO) exceeded a proposed MCID for weighted mean FEV\(_1\);\(^{23}\) and was within the putative MCID range of 100–140 mL for trough FF/VI;\(^{26}\) however, the treatment difference between the two therapies was not statistically significant.

In our study, secondary endpoints were nested under the primary endpoint; since the primary endpoint difference did not reach statistical significance, all differences reported for the secondary efficacy endpoints must be regarded as descriptive only. Nonetheless, numerical differences were observed between the treatment groups in some other lung function endpoints, symptoms, use of rescue medication, and QoL. Rescue medication use and dyspnea were consistently reduced over the 12 weeks of treatment with FF/VI compared with TIO. QoL (measured by mean SGRQ-C total and component scores and by the percentage of SGRQ-C responders) and health status (measured by mean CAT scores) improved with both treatments, but the improvement was greater with FF/VI treatment compared with TIO. The total SGRQ-C
improvement with FF/VI was −5.0 at 8 weeks and −3.9 at 12 weeks, similar to the result of −4.3 at 12 weeks in another study. A previous comparison of another ICS/LABA with TIO provided similar findings of a superior effect of ICS/LABA on QoL.7 In our study, mean SGRQ-C improvements reached or approached the MCID after treatment with FF/VI, but not TIO. In contrast, CAT improvement scores were more modest after FF/VI treatment and did not approach the MCID. Although the most reliable estimate of MCID may be two points,27 the CAT only allows integer scores, and the bar may be set too high for population-based comparisons.28

As COPD progresses, patients experience increasing air trapping and hyperinflation reflected by IC declines.29 In our study, 12 weeks’ TIO treatment compared with FF/VI provided improvements in trough IC and FVC. Improvements in IC have been associated with decreased exertional but not resting dyspnea.30,31 In this regard, it is noteworthy that although treatment with TIO was associated with a greater improvement in IC, dyspnea improved more so with FF/VI.

The safety profiles of FF/VI and TIO in this study were similar, with a couple of exceptions. Pneumonia was reported more frequently in FF/VI-treated subjects, all radiographically confirmed, consistent with previous reports of a higher incidence with FF/VI treatment13 or other ICS treatment alone or in combination with a LABA.32,33

More subjects failed to complete the study while receiving TIO (12% vs 6% for FF/VI), including those who experienced COPD exacerbations (4% vs 2%, respectively). In 1-year studies of FF/VI versus VI,13 treatment with FF/VI reduced exacerbation rates compared with treatment with VI alone. Likewise, in a 2-year study comparing TIO with FP/SALM, more TIO-treated subjects withdrew, but the rates of COPD exacerbation did not differ between groups.7

In our population of patients with comorbid CVD or increased risk of CVD, there were no apparent effects of FF/VI or TIO treatment on the CV safety profile or differences between treatments, with the exception of two deaths from CV events in the TIO-treated subjects. We paid particular attention to measuring CV safety, employing 24-hour Holter monitoring in a subset of subjects, performing frequent ECGs, and monitoring CV AEs because this was an at-risk population and due to previous reports of TIO and LABA CV safety. While some earlier analyses suggested that the risk of CVD or CV mortality was increased in COPD patients using the TIO Respimat® (Boehringer Ingelheim) inhaler,5,34,35 this was not confirmed in a subsequent trial,35 and the TIO HandiHaler has not been associated with an increased risk compared with placebo in recent studies.6,16–39

In our study, the rate of ventricular tachycardias was somewhat higher than anticipated. However, the definition can vary from that used here (>100 beats/min) to that of >120 beats/min.40–41 Also, subjects in both groups experienced a greater incidence of ventricular ectopic beats at the end of the study than at screening, but the circumstances of those Holter measurements differed, as the end-of-study visit was conducted with a 24-hour in-patient stay.

The CV risk profile of FF/VI versus TIO has also been examined in a recent clinical study of FF/VI effects on arterial stiffness, a marker that is elevated in COPD and is a predictor of CV events and mortality.42,43 Arterial stiffness, as measured by aortic pulse wave velocity, was lowered by both FF/VI and TIO in subjects with moderate-to-severe COPD and baseline aortic pulse wave velocity >11.0 m/s.44 These subjects also had a high incidence of baseline CVD, but the number of CV events over the 12-week treatment period was low and similar between treatments.44 Results of the ongoing SUMMIT trial in 16,000 subjects with comorbid CVD risk and COPD will further determine the impact of FF/VI on CVD and mortality.45

The major limitations of our study were its lack of placebo arm, relatively small sample size, and 12-week duration, making the study incapable of establishing effects on pneumonia incidence, CVD, or mortality.

However, this randomized study had several strengths. First, it provided for prospective collection of COPD exacerbation and pneumonia data. Second, it applied rigorous CV monitoring and compared ICS/LABA and TIO treatment in a population with comorbid COPD. Most importantly, the study is the first to compare the efficacy of a once-daily ICS/LABA with TIO. Patients with COPD demonstrate low adherence to study medication,46 which is significantly associated with increased risk of death and COPD exacerbations.47 Once-daily medications such as TIO have been shown to improve adherence in medication use over medications with more frequent dosing regimens,48–50 and it is expected that FF/VI will have similar benefits, especially in certain patient populations who prefer once-daily medications.51 Adherence to study medication in the current study was high, with no difference between the FF/VI and TIO groups.

Conclusion

In conclusion, treatment with FF/VI or TIO for 12 weeks improved lung function in COPD subjects who have or are at risk for comorbid CVD, but there was no significant difference between the treatment groups. Reduction in rescue medication use and dyspnea and improving SGRQ-C-indicated
QoL favored FF/VI, whereas improvements in FVC and IC favored TIO. Other than more reports of pneumonia in FF/VI-treated subjects and CV AEs in TIO-treated subjects, the safety profiles of the treatments were similar. The risk of pneumonia should be weighed against the benefits of FF/VI treatment, such as the reduction in COPD exacerbations, increased QoL, and medication adherence, when clinicians consider the potential use of FF/VI for patients with COPD. The results of an ongoing larger FF/VI trial in subjects with COPD with CVD (SUMMIT)45 are anticipated to provide additional evidence regarding efficacy and safety in this population.

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