Navafenterol (AZD8871) in patients with COPD: a randomized, double-blind, phase I study evaluating safety and pharmacodynamics of single doses of this novel, inhaled, long-acting, dual-pharmacology bronchodilator

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Abstract

Background: Navafenterol (AZD8871) is a dual-pharmacology muscarinic antagonist β2-agonist (MABA) molecule in development for the treatment of chronic obstructive pulmonary disease (COPD). The pharmacodynamics, safety and tolerability of single doses of navafenterol were investigated in patients with moderate to severe COPD.

Methods: This was a randomized, five-way complete cross-over study. Patients received single doses of navafenterol 400 µg, navafenterol 1800 µg and placebo (all double-blind) and indacaterol 150 µg and tiotropium 18 µg (both open-label active comparators). The primary pharmacodynamic endpoint was change from baseline in trough forced expiratory volume in 1 s (FEV1) on day 2. Safety and tolerability were monitored throughout.

Results: Thirty-eight patients were randomized and 28 (73.7%) completed the study. Navafenterol 400 µg and 1800 µg demonstrated statistically significant improvements vs placebo in change from baseline in trough FEV1 (least squares mean [95% confidence interval]: 0.111 [0.059, 0.163] L and 0.210 [0.156, 0.264] L, respectively, both P < .0001). The changes were significantly greater with navafenterol 1800 µg vs the active comparators (least squares mean treatment difference: 0.065–0.069 L, both P < .05). The frequency of treatment-emergent adverse events was similar for placebo and the active comparators (range 34.4–37.5%), slightly higher for navafenterol 400 µg (52.9%), and lowest for navafenterol 1800 µg (22.6%).

Conclusions: Both doses of navafenterol demonstrated sustained bronchodilation over 24 h. Navafenterol was well tolerated and no safety concerns were raised.

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Keywords: Bronchodilator, COPD, MABA, Dual-pharmacology muscarinic receptor antagonist β2-adrenoceptor agonist, Safety, Pharmacokinetics

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Background

Inhaled long-acting bronchodilators are integral in the pharmacological management of chronic obstructive pulmonary disease (COPD). Whilst bronchodilator monotherapy is recommended for treatment initiation in most patients with COPD [1], there is considerable evidence that the combination of a long-acting muscarinic receptor antagonist (LAMA) with a long-acting β₂-agonist (LABA) offers additional benefits over monotherapy [2, 3]. The Global Initiative for Chronic Obstructive Lung Disease (GOLD) report recommends a combination of a LAMA and LABA for treatment escalation, or initiation, in patients with greater symptom burden or exacerbation risk [1].

Co-formulation of fixed-dose combinations (FDCs), including LAMA/LABA combinations, is technically challenging [4]. The development of dual-pharmacology muscarinic antagonist β₂-agonist (MABA) molecules offers a new approach to the treatment of COPD [4]. These molecules combine two mechanisms of action within a single entity, offering potential advantages such as the delivery of a fixed ratio of LAMA/LABA activity into each lung region, simplification of technical and clinical development, and the potential for additive or synergistic bronchodilation over either entity alone [4, 5]. The use of MABAs also creates a platform for the inclusion of another drug, such as an inhaled corticosteroid (ICS) or another anti-inflammatory agent [4]. Navafenterol (AZD8871, formerly LAS191351) is one of the few MABAs in clinical development for the treatment of COPD [1]. Its pharmacological profile has been extensively studied in preclinical investigations in vitro and in vivo, and these investigations have confirmed its dual action at β₂- and muscarinic receptors [6].

A study of navafenterol was conducted in two parts: a first-in-human single ascending-dose study in patients with mild, persistent asthma (Part 1) and a five-way crossover, single-dose study in patients with moderate to severe COPD (Part 2). In Part 1, single ascending doses of navafenterol 50, 200, 400, 900, 1800, and 2100 μg were well tolerated and doses ≥200 μg produced clinically meaningful, sustained bronchodilation [7]. Here, we report data from Part 2 of this study; the primary objective was to assess the pharmacodynamics, safety, and tolerability of single doses of navafenterol in patients with moderate to severe COPD, with exploratory comparisons vs placebo and the active comparators indacaterol (a LABA) and tiotropium (a LAMA).

Methods

Study design

This was a randomized, double-blind, five-way complete crossover, placebo and active-controlled single-dose study (NCT02573155). A subset of patients participated in a pharmacokinetic (PK) sub-study, the methods and results of which are reported in Additional file 1. The study was conducted at a single site in the UK (Medicines Evaluation Unit, Manchester). The study protocol was approved by an Independent Ethics Committee (NRES Committee – Cambridgeshire and Hertfordshire, Health Research Authority, Nottingham, UK; Reference No 15/EE/0329; see Additional file 1) and the UK Medicines and Healthcare Products Regulatory Agency. The informed consent form was also reviewed by the Independent Ethics Committee. The study was performed in accordance with the Declaration of Helsinki and the International Conference on Harmonization/Good Clinical Practice guidelines. All patients provided written, informed consent before enrolment in the study. The first patient was randomized on April 25, 2016; the last patient visit was August 22, 2016.

Following a screening evaluation (Fig. 1) and run-in period of 14 to 28 days, patients were assigned to one of 10 treatment sequences, each with a 1:1:1:1:1 randomization ratio, according to a William’s design for crossover studies (Fig. 2). There were five treatment periods, with patients remaining on-site for 36 h post-dose in each treatment period. The treatments administered were navafenterol 400 μg, navafenterol 1800 μg, and placebo (all double blind, each administered via a variant of the Genuair™/Pressair™ dry-powder inhaler [DPI] adapted internally to deliver a single dose of inhalation powder) and indacaterol 150 μg and tiotropium 18 μg (both open-label, delivered by Onbrez Breezhaler® and HandiHaler® DPIs, respectively). A single dose was administered at the same time (±1 h) on day 1 of each treatment period, between 7 am and 10 am. There was a washout period of 10 to 21 days following navafenterol or placebo treatment and 7 to 21 days following indacaterol or tiotropium treatment.

Patients were withdrawn from their usual COPD therapy after signing the informed consent form but maintained on their usual ICS dose; those receiving an ICS/LABA FDC were switched to ICS monotherapy. Salbutamol was provided as reliever medication for the duration of the study.

A follow-up visit was performed 7 (±2) days after the last dose or after discontinuation. Patients were contacted by telephone 14 (±2) days after the last dose, to record any adverse events (AEs).

Patients

Male and female (of non-child-bearing potential) patients aged ≥40 years with a clinical diagnosis of stable moderate to severe COPD according to GOLD 2015 criteria [8] were eligible for inclusion. Other inclusion criteria included: postbronchodilator forced expiratory volume in 1 s (FEV₁) < 80% and ≥30% predicted normal
and FEV₁/forced vital capacity (FVC) ratio < 70%; body mass index < 40 kg/m², current or ex-smoker with a smoking history of ≥10 pack-years, no evidence of clinically significant respiratory and/or cardiovascular conditions or laboratory abnormalities and no history of thoracic surgery. Other inclusion and exclusion criteria and study restrictions are reported in Additional file 1.

Assessments
Pharmacodynamics
The primary pharmacodynamic endpoint was change from baseline in trough FEV₁. Trough FEV₁ was the mean of the FEV₁ values obtained at 23 h and 24 h after study drug administration. Secondary endpoints included change from baseline in normalized FEV₁ area under the curve from 0 to 6 h (AUC₀–₆), 0–12 h (AUC₀–₁₂), 12–24 h (AUC₁₂–₂₄), and 0–24 h (AUC₀–₂₄) post-dose, change from baseline in FEV₁ at each scheduled timepoint post-dose, number and percentage of patients achieving ≥100 mL change from baseline in FEV₁ (minimal clinically important difference) during the 6 h post-dose, change from baseline in and time to peak FEV₁ on day 1, and change from baseline in trough FVC on day 2.

Details of the timing and measurement standards for FEV₁ and FVC assessments are provided in Additional file 1.

Fig. 1 Patient disposition and flow. AE = adverse event; PK = pharmacokinetic. *Five patients discontinued due to treatment emergent AEs leading to study drug discontinuation, 1 patient withdrew due to a treatment emergent AE during the washout period, and 1 withdrew due to a non-treatment-emergent AE; † patient had a positive drug screen test for cocaine; ‡ patient did not meet stability/variability criteria.

Fig. 2 Study design. TP = treatment period. *The figure shows an example of the washout timings for one patient. The washout period was 10 to 21 days following navafenterol or placebo, and 7 to 21 days following the active comparators indacaterol and tiotropium. Therefore the length of each washout period depended on the patient’s randomization sequence. One exception per patient of up to 28 days was considered acceptable; † days following administration of last dose.
### Safety and tolerability

AEs were collected from consent until the telephone follow-up. Treatment-emergent AEs (TEAEs) were defined as AEs that appeared or worsened after the first dose of study drug and within 14 days following the last dose. Other safety assessments included physical examination, blood pressure, clinical laboratory assessments (blood chemistry, hematology, urinalysis, and blood potassium and glucose [both measured via i-STAT]), and 12-lead electrocardiography (see Additional file 1).

### Statistical analysis

There was no formal sample size calculation. It was considered that a sample size of 30 patients would be sufficient to meet the objectives of the study. Approximately 40 patients were randomized to account for an approximate 25% dropout rate.

Pharmacodynamic variables were analyzed in the per protocol population, defined as all randomized patients who satisfied the main inclusion/exclusion criteria, completed at least one treatment period, and had no major protocol violations. All primary and secondary pharmacodynamic variables, with the exception of time to peak FEV₁, were analyzed by an analysis of covariance model for crossover designs. All statistical comparisons used 2-sided hypothesis tests, and the significance level was set at .05 without multiplicity adjustment. Further details are in Additional file 1.

Safety outcomes were analyzed descriptively in the safety population (all randomized patients who received at least one dose of the study drug).

SAS version 9.2 or later (SAS Institute, Inc., Cary, NC) was used for all statistical analyses.

### Results

#### Patient demographics and baseline characteristics

A total of 72 patients were screened; of these, 38 were randomized and 28 (73.7%) completed the study (Fig. 1). Patient demographics and baseline characteristics for the safety population are presented in Table 1. Twenty-two (57.9%) had severe airflow limitation. Mean (standard deviation) bronchodilator reversibility was 0.217 (0.124) L.

| Baseline Characteristic | Total |
|------------------------|-------|
| Number of patients in safety population | 38 |
| Age, years | 65.6 (6.4) |
| Male, n (%) | 22 (57.9) |
| White, n (%) | 38 (100) |
| Body mass index, kg/m² | 27.7 (3.5) |
| Smoking status, n (%) |
| Current | 14 (36.8) |
| Former | 24 (63.2) |
| COPD severity, n (%) |
| Moderate | 22 (57.9) |
| Severe | 16 (42.1) |
| Duration of COPD, years | 9.5 (5.9) |
| Number of COPD exacerbations in previous 12 months, n (%) |
| 1 | 8 (21.1) |
| 2 | 6 (15.8) |
| 3 | 2 (5.3) |
| Postbronchodilator % predicted FEV₁ | 52.0 (12.5) |
| Postbronchodilator FEV₁/FVC | 45.8 (10.2) |
| Bronchial reversibility, % | 19.89 (13.72) |
| FEV₁ absolute reversibility, L | 0.217 (0.124) |

**Table 1** Patient Demographics and Baseline Characteristics (Safety Population)

COPD chronic obstructive pulmonary disease, FEV₁ forced expiratory volume in 1 s, FVC forced vital capacity

Data are mean (standard deviation) unless otherwise stated

### Pharmacodynamics

All active treatments showed statistically significant improvements vs placebo in change from baseline in trough FEV₁ (Fig. 3a; Table 2; least squares [LS] mean treatment difference: navafenterol 400 μg, 0.111 L; navafenterol 1800 μg, 0.210 L; indacaterol 150 μg, 0.141 L; tiotropium 18 μg, 0.145 L; all P < .0001). The magnitude of change in trough FEV₁ was greater with navafenterol 1800 μg compared with both active comparators (LS mean treatment difference 0.065–0.069 L, both P < .05).

A similar pattern of results with a greater effect of navafenterol 1800 μg compared with both active comparators was observed for change from baseline in normalized FEV₁ AUC₀–₁₂, AUC₀–₂₄, and AUC₁₂–₂₄. Navafenterol 400 μg also achieved statistically significant improvements vs tiotropium for AUC₀–₂₄ and AUC₀–₁₂ (Fig. 3b; Table 2).

Mean (SD) time to peak FEV₁ was 3.4 (2.1) h, 3.9 (2.0) h, 3.0 (2.2) h, and 3.3 (2.2) h with navafenterol 400 μg, navafenterol 1800 μg, indacaterol, and tiotropium, respectively. All active treatments showed statistically significant improvements in change from baseline in peak FEV₁ vs placebo (e-Fig. 1; Table 2); navafenterol 400 μg, 0.218 L; navafenterol 1800 μg, 0.285 L; indacaterol 150 μg, 0.227 L; tiotropium 18 μg, 0.139 L; P < .01 for all).

Improvements in peak FEV₁ with navafenterol 1800 μg were significantly greater compared with tiotropium (P < .01) but not indacaterol or navafenterol 400 μg.
Improvements with navafenterol 400 μg were similar to those of the active comparators. At all timepoints from 15 min to 36 h post-dose, all active treatments showed statistically significant improvements in change from baseline in FEV₁ vs placebo (Fig. 4; \( P < .01 \) for all). Additionally, navafenterol 1800 μg significantly improved change from baseline in FEV₁ vs placebo at 5 min post-dose (\( P = .0123 \)). FEV₁ data over time are further described in Additional file 1, along with the proportion of patients achieving ≥100 mL change from baseline in FEV₁ and trough FVC.

Safety and tolerability
Overall, 31 (81.6%) patients reported TEAEs (Table 3). The frequency of TEAEs was similar for placebo and the active comparators (range 34.4–37.5%), slightly higher for navafenterol 400 μg (52.9%), and lowest for navafenterol 1800 μg (22.6%). Overall, the most frequently reported TEAEs across all groups were headache (31.6%) and nasopharyngitis (13.2%). Most TEAEs were mild or moderate in intensity and there were no deaths during the study. Only one incidence of headache was considered to be related to the study treatment by the study investigator, which occurred in the tiotropium group. Five (13.2%) patients discontinued treatment due to TEAEs, including two serious AEs (physical assault and fractured C1 vertebra) and three non-serious TEAEs (COPD exacerbation, pneumonia, and headache); none of these TEAEs were assessed as considered related to treatment.

There were no clinically significant changes in clinical laboratory tests, blood glucose and serum potassium concentrations (e-Fig. 2), heart rate, or blood pressure. Small increases in mean change from baseline in QT interval corrected for heart rate using the Fridericia
| Change From Baseline | Placebo (n = 32)* | Nafenterol 400 µg (n = 34) | Nafenterol 1800 µg (n = 31) | Indacaterol 150 µg (n = 32) | Tiotropium 18 µg (n = 30) |
|---------------------|------------------|---------------------------|-----------------------------|-----------------------------|--------------------------|
| Trough FEV1 on day 2, L (primary endpoint) |  |  |  |  |  |
| Trough FEV1 value | 0.0171 (0.0191, 0.123) | 0.170 (0.166, 0.224) | 0.101 (0.048, 0.154) | 0.105 (0.051, 0.159) |  |
| Difference vs placebo | 0.111 (0.059, 0.163) | 0.210 (0.156, 0.264) | 0.141 (0.087, 0.195) | 0.145 (0.090, 0.200) |  |
| P-value | < .0001 | < .0001 | < .0001 | < .0001 |  |
| Difference vs indacaterol | -0.030 (0.083, 0.023) | 0.069 (0.015, 0.123) |  |  |  |
| P-value | 2.600 | 0.126 |  |  |  |
| Difference vs tiotropium | -0.034 (0.088, 0.020) | 0.065 (0.011, 0.120) |  |  |  |
| P-value | 2.141 | 0.197 |  |  |  |
| Peak FEV1 on day 1, L |  |  |  |  |  |
| Peak FEV1 value | 0.121 (0.044, 0.198) | 0.339 (0.264, 0.413) | 0.406 (0.328, 0.483) | 0.348 (0.271, 0.424) | 0.260 (0.181, 0.340) |
| Difference vs placebo | 0.218 (0.122, 0.314) | 0.285 (0.186, 0.384) | 0.227 (0.128, 0.326) | 0.139 (0.039, 0.240) |  |
| P-value | < .0001 | < .0001 | < .0001 | < .0001 | .0069 |
| Difference vs indacaterol | -0.009 (0.106, 0.088) | 0.058 (0.041, 0.157) |  |  |  |
| P-value | .8556 | .2476 |  |  |  |
| Difference vs tiotropium | 0.079 (0.020, 0.178) | 0.146 (0.045, 0.246) |  |  |  |
| P-value | .1180 | .0048 |  |  |  |
| Normalized AUC_{0-24} of FEV1, L |  |  |  |  |  |
| Normalized FEV1 AUC_{0-24} value | -0.021 (0.075, 0.082) | 0.168 (0.115, 0.221) | 0.255 (0.201, 0.309) | 0.159 (0.105, 0.212) | 0.126 (0.072, 0.180) |
| Difference vs placebo | 0.189 (0.150, 0.228) | 0.276 (0.236, 0.316) | 0.180 (0.139, 0.220) | 0.147 (0.106, 0.188) |  |
| P-value | < .0001 | < .0001 | < .0001 | < .0001 | < .0001 |
| Difference vs indacaterol | 0.010 (0.030, 0.049) | 0.096 (0.056, 0.136) |  |  |  |
| P-value | .6305 | < .0001 |  |  |  |
| Difference vs tiotropium | 0.042 (0.002, 0.082) | 0.129 (0.088, 0.169) |  |  |  |
| P-value | .0406 | < .0001 |  |  |  |
| Normalized AUC_{12-24} of FEV1, L |  |  |  |  |  |
| Normalized FEV1 AUC_{12-24} value | -0.022 (0.081, 0.086) | 0.233 (0.175, 0.290) | 0.301 (0.242, 0.359) | 0.192 (0.134, 0.250) | 0.156 (0.096, 0.215) |
| Difference vs placebo | 0.255 (0.211, 0.299) | 0.323 (0.278, 0.368) | 0.214 (0.169, 0.260) | 0.178 (0.132, 0.224) |  |
| P-value | < .0001 | < .0001 | < .0001 | < .0001 | < .0001 |
| Difference vs indacaterol | 0.041 (0.004, 0.085) | 0.109 (0.063, 0.154) |  |  |  |
| P-value | .0732 | < .0001 |  |  |  |
| Difference vs tiotropium | 0.077 (0.032, 0.123) | 0.145 (0.099, 0.191) |  |  |  |
| P-value | .0010 | < .0001 |  |  |  |
Table 2 Primary and Secondary Pharmacodynamic Endpoints (Per Protocol Population) (Continued)

| Change From Baseline | Placebo (n = 32)$^a$ | Navafenterol 400 μg (n = 34) | Navafenterol 1800 μg (n = 31) | Indacaterol 150 μg (n = 32) | Tiotropium 18 μg (n = 30) |
|----------------------|----------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|
| Normalized FEV1 AUC_{0–24} value | \(-0.029 (-0.083, 0.025)\) | \(0.104 (0.052, 0.156)\) | \(0.212 (0.158, 0.266)\) | \(0.128 (0.075, 0.181)\) | \(0.100 (0.045, 0.154)\) |
| Difference vs placebo | \(0.133 (0.085, 0.180)\) | \(0.241 (0.192, 0.289)\) | \(0.157 (0.108, 0.205)\) | \(0.129 (0.079, 0.178)\) |
| \(P\)-value | < .0001 | < .0001 | < .0001 | < .0001 |
| Difference vs indacaterol | \(-0.024 (-0.071, 0.023)\) | \(0.084 (0.037, 0.131)\) | | |
| \(P\)-value | 0.3172 | < .0001 | | |
| Difference vs tiotropium | \(0.004 (-0.044, 0.052)\) | \(0.112 (0.064, 0.160)\) | | |
| \(P\)-value | 0.8591 | < .0001 | | |

$^{a}$Two patients from the placebo group were excluded from the normalized FEV1 AUC_{0–24} analysis due to missing data. One patient was missing data from 12 to 24 h post-dose and the other from 14 h post-dose. For all other endpoints in this table, all patients exposed to each treatment were included in the analyses.

AUC_{0–24} area under the curve from 0 to 24 h post-dose, AUC_{0–12} area under the curve from 0 to 12 h post-dose, AUC_{12–24} area under the curve from 12 to 24 h post-dose, FEV1 forced expiratory volume in 1 s. Values are least squares means (95% confidence interval).
formula (QTcF) were observed in all active treatment groups compared with placebo; the largest increase in LS mean (90% confidence interval) change from baseline in QTcF vs placebo was observed 3 h post-dose with navefenterol 400 μg (3.05 [0.106, 6.00] ms) and navefenterol 1800 μg (5.04 [2.01, 8.06] ms), 2 h post-dose with indacaterol (4.38 [1.84, 6.92] ms), and 36 h post-dose with tiotropium (3.98 [1.27, 6.69] ms). Seven male patients met the criteria for potentially clinically significant QTcF increase, including 6 patients with QTcF values > 450 ms and 1 patient with an increase from baseline > 30 ms; none of these abnormalities were considered clinically significant by the investigator or reported as TEAEs. QTcF values > 450 ms were observed sporadically in all active treatment groups at all timepoints (including baseline) with no clear pattern observed (Fig. 5).

**Discussion**

Single doses of navefenterol 400 μg and 1800 μg produced sustained bronchodilation over 24 h in patients with moderate to severe COPD, with significant improvements of 0.111 L and 0.210 L, respectively, in trough FEV₁ vs placebo. The magnitude of the changes was significantly greater with navefenterol 1800 μg compared with the active comparators, indacaterol (0.069 L) and tiotropium (0.065 L). Both doses of navefenterol significantly

![Fig. 4 Mean change from baseline in FEV₁ over 36 h (per protocol population). Data are LS means ± standard error. The number of patients exposed to each treatment differed from the number of non-missing observations for navefenterol 1800 μg (at 45 min and 2 h; both n = 30) and placebo (at 12 and 14 h; n = 31 and n = 30, respectively). FEV₁ = forced expiratory volume in 1 s; LS = least squares](image)

**Table 3** Frequency of TEAEs Overall and Occurring in ≥2 Patients, by MedDRA® Preferred Term (Safety Population)

| TEAE (Preferred Term)* | Placebo (n=32) | navefenterol 400 μg (n=34) | navefenterol 1800 μg (n=31) | Indacaterol 150 μg (n=32) | Tiotropium 18 μg (n=30) | All (N=38) |
|------------------------|----------------|---------------------------|-----------------------------|--------------------------|------------------------|------------|
| Any event, n (%)       | 11 (34.4)      | 18 (52.9)                 | 7 (22.6)                    | 12 (37.5)                | 11 (36.7)              | 31 (81.6)  |
| Headache               | 7 (21.9)       | 5 (14.7)                  | 3 (9.7)                     | 4 (12.5)                 | 7 (23.3)               | 12 (31.6)  |
| Nasopharyngitis        | 1 (3.1)        | 1 (2.9)                   | 1 (3.2)                     | 2 (6.3)                  | 1 (3.3)                | 5 (13.2)   |
| COPD                   | 0              | 2 (5.9)                   | 1 (3.2)                     | 1 (3.1)                  | 0                      | 4 (10.5)   |
| Erythema               | 1 (3.1)        | 1 (2.9)                   | 0                           | 1 (3.1)                  | 0                      | 3 (7.9)    |
| Medical device site reaction | 1 (3.1) | 0                         | 1 (3.2)                     | 1 (3.1)                  | 0                      | 3 (7.9)    |
| Constipation           | 1 (3.1)        | 1 (2.9)                   | 1 (3.2)                     | 0                        | 0                      | 2 (5.3)    |
| Nausea                 | 1 (3.1)        | 1 (2.9)                   | 0                           | 0                        | 0                      | 2 (5.3)    |
| Rhinitis               | 0              | 1 (2.9)                   | 0                           | 1 (3.1)                  | 0                      | 2 (5.3)    |

COPD: chronic obstructive pulmonary disease, MedDRA: Medical Dictionary for Regulatory Activities, TEAE: treatment emergent adverse event.

*MedDRA version 18.1
improved change from baseline in normalized FEV\textsubscript{1} AUC\textsubscript{0–24} vs placebo and tiotropium, with navafenterol 1800 µg also improving this parameter vs indacaterol.

The changes from baseline in trough FEV\textsubscript{1} with both doses of navafenterol exceeded the minimal clinically important difference of 0.100 L vs placebo [9]. The magnitude of change in trough FEV\textsubscript{1} vs placebo with navafenterol 400 µg (0.111 L) and 1800 µg (0.210 L) was similar to that observed with single doses of the MABA, batefenterol, in a population of patients with moderate COPD (FEV\textsubscript{1} 50 to 80% predicted; patients with > 2 exacerbations in the previous 12 months excluded), at doses of 400 µg (0.141 L) and 1200 µg (0.184 L) [10]. Both doses of navafenterol demonstrated a rapid onset of action (within 5–15 min post-dose), with changes from baseline in peak FEV\textsubscript{1} vs placebo of 0.218 L and 0.285 L with navafenterol 400 µg and 1800 µg, respectively.

One limitation of the study was the inclusion of active comparators as monotherapy rather than combination therapy, meaning that no conclusions can be drawn about the efficacy of AZD8777 compared with LAMA/LABA FDCs. However, the magnitude of the treatment difference between navafenterol 1800 µg and indacaterol 150 µg for trough FEV\textsubscript{1} (0.069 L) was similar to that observed between the LAMA/LABA FDC DPI glycopyrronium/indacaterol 50/110 µg and indacaterol 150 µg (0.07 L) in a 26-week study in patients with moderate to severe COPD [11]. Similarly, the treatment difference between navafenterol 1800 µg and the LAMA tiotropium 18 µg (0.065 L) was similar to that observed between tiotropium/olodaterol 5/5 µg FDC delivered via soft mist inhaler and tiotropium 5 µg (0.050–0.071 L) after 24 weeks’ treatment in patients with moderate to very severe COPD [12]. Whilst it is difficult to draw firm conclusions across studies due to differences in methodologies, these results suggest that navafenterol 1800 µg may provide similar benefits to LAMA/LABA combinations compared with monotherapy. It is important to note that the above-mentioned studies report data following repeat dosing, and the current study was limited by its single-dose design; however, a greater bronchodilatory response for navafenterol has also been achieved with repeat-dosing vs single-dosing [13]. Furthermore, a 4-week study with batefenterol found that greater improvements in FEV\textsubscript{1} were observed on day 28 compared with day 1 for all doses investigated [14].

The pharmacokinetic data generated for navafenterol are consistent with the rapid onset of action and sustained bronchodilation observed in the pharmacodynamic response. Navafenterol is rapidly absorbed into the bloodstream (median time to maximum concentration 1–2 h) and slowly eliminated from plasma.

Single doses of navafenterol were well tolerated and no safety concerns were identified. The only TEAE occurring in more than two patients in the navafenterol treatment groups was headache; however, its frequency was lower than in the placebo group. Small increases in the duration of QT\textsubscript{cF} were observed with both doses of navafenterol; however, the upper limit of the 90% confidence interval for the largest QT\textsubscript{cF} change from baseline vs placebo was < 10 ms. Possible effects of navafenterol on QT\textsubscript{cF} interval require further evaluation in larger clinical trials with repeat dosing. There were no clinically
significant changes in blood glucose and potassium concentrations in the present study.

The proportion of reversible patients in this study was possibly due to the small sample size of the trial. This may have contributed to the effects on lung function observed with navafenterol and the active controls. However, the effects on trough FEV1 with navafenterol in this part of the study are in line with the effect size seen in the phase 2a study that only included reversible patients [15]. Since statistical comparisons between navafenterol and the active comparators were made using ANCOVA with no correction for multiple testing, statistical significance should be interpreted with caution, considering the lack of overall control of the type I error. Since this was a first-time-in-human phase I study, additional studies with navafenterol and larger sample sizes will be conducted to elucidate the efficacy of the drug as it progresses through development.

**Conclusion**

Both doses of navafenterol demonstrated rapid onset of action (within 5–15 min post-dose) and sustained bronchodilation over 24 h. The bronchodilatory efficacy of navafenterol 1800 μg was greater than that of both indacaterol 150 μg and tiotropium 18 μg. Overall, navafenterol was well tolerated and no safety concerns were raised. These results support the continued clinical development of navafenterol.

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**Supplementary information**

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**Abbreviations**

AE: Adverse event; AUC0–t: Area under the concentration-time curve from time zero extrapolated to infinity; AUC0–12: Area under the concentration-time curve from zero to the last quantifiable measurable concentration; AUC0–6: Area under the curve from 0 to 6 h post-dose; AUC0–12: Area under the curve from 0 to 12 h post-dose; AUC0–24: Area under the curve from 0 to 24 h post-dose; AUC12–24: Area under the curve from 12 to 24 h post-dose; COPD: Chronic obstructive pulmonary disease; DPI: Dry-powder inhaler; FDC: Fixed dose combination; FEV1: Forced expiratory volume in 1 s; GOLD: Global Initiative for Chronic Obstructive Lung Disease; PVC: Forced vital capacity; ICS: Inhaled corticosteroid; LABA: Long-acting β2-adrenoceptor agonist; LAMA: Long-acting muscarinic receptor antagonist; LSS: Least squares; MABA: Muscarinic receptor antagonist and β2-adrenoceptor agonist; PK: Pharmacokinetic; QTcF: QT interval corrected for heart rate using the Fridericia formula; SD: Standard deviation; TEAE: Treatment-emergent adverse event

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**About this supplement**

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**Authors’ contributions**

D. S. was an investigator in the study and contributed to the interpretation of data. C. A., U. W. H., L. J., B. S., C. V., A. L., and A. A. conceived the study and V. B. managed the study. All authors contributed to the interpretation of data. All authors critically reviewed the manuscript and approved the final version for submission. All named authors meet the International Committee for Medical Journal Editors criteria for authorship of this manuscript, take responsibility for the integrity of the work as a whole, and have given final approval for the version to be published. D. S. takes responsibility for (is the guarantor of) the content of the manuscript, including the data and analysis.

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**Availability of data and materials**

Data underlying the findings described in this manuscript may be obtained in accordance with AstraZeneca’s data sharing policy described at https://astrazenecagrouptrials.pharmacm.com/ST/Submission/Disclosure.

**Ethics approval and consent to participate**

The study protocol was approved by an Independent Ethics Committee (NRES Committee – Cambridgeshire and Hertfordshire, Health Research Authority, Nottingham, UK; Reference No 15/EE/0329; see Additional file 1) and the UK Medicines and Healthcare Products Regulatory Agency. The informed consent form was also reviewed by the Independent Ethics Committee. The study was performed in accordance with the Declaration of Helsinki and the International Conference on Harmonization/Good Clinical Practice guidelines. All patients provided written, informed consent before enrolment in the study.

**Consent for publication**

Not applicable.

**Competing interests**

D. S. has received sponsorship to attend international meetings, honoraria for lecturing or attending advisory boards, and research grants from various pharmaceutical companies including Apellis, AstraZeneca, Boehringer Ingelheim, Chiesi, Cipla, Genentech, GlaxoSmithKline, Glenmark, Johnson and Johnson, Menarini, Mundipharma, Novartis, Pentinnovate, Pfizer, Pulmatrix, Skypharma, Teva, Thevenez, and Verona. B. C., A., C. A., L. J., B. S., A. L., and I. P. are employees of AstraZeneca and may own stock or stock options. C. V. and A. A. were employees of AstraZeneca at the time the study was conducted. C. V. is a current employee of PAREXEL. A. A. is a current employee of Ellex Pharmaceuticals.
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