Revefenacin: First Global Approval

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Abstract

Revefenacin (YUPELRI™) inhalation solution, a long-acting muscarinic antagonist (i.e. an anticholinergic) developed by Theravance Biopharma and Mylan, is the first and currently the only once-daily, nebulized bronchodilator to be approved in the USA for the treatment of chronic obstructive pulmonary disease (COPD). In November 2018, based on results of three phase III trials, the US Food and Drug Administration granted market authorization to revefenacin for the maintenance treatment of patients with COPD. This article summarizes the milestones in the development of revefenacin leading to this first global approval.

1 Introduction

Chronic obstructive pulmonary disease (COPD) is a progressive disease that is characterized by persistent airflow limitation [1]. COPD is associated with considerable medical, social, and economic burden and is expected to be the third leading cause of death worldwide by 2020. The goal of COPD management is to improve lung function, reduce clinical symptoms of COPD, and prevent future exacerbations. The mainstay maintenance treatment for COPD patients includes inhaled long-acting muscarinic antagonist (LAMAs) and/or long-acting β2-agonists (LABAs), with or without inhaled corticosteroid (ICS) [1]. Long-acting bronchodilators (e.g. LABAs or LAMAs) are often delivered via hand-held devices, such as a dry powder inhaler (DPI) or a metered dose inhaler (MDI). However, some COPD patients [e.g. the elderly, cognitively impaired or physically limited and with suboptimal peak inspiratory flow rates (PIFR)] find it difficult to use such devices, resulting in inaccurate dosing, poor adherence, and potentially poor clinical outcomes [2]. For these patients, nebulized drug delivery, which is easier to use and provides similar efficacy to hand-held inhalers can be an alternative treatment option. However, nebulized bronchodilator therapy options for the maintenance treatment of COPD are currently limited [2].

Revefenacin (YUPELRI™ [3]) inhalation solution is a once-daily nebulized LAMA that has been developed by Theravance Biopharma and Mylan for the maintenance treatment of patients with COPD [4]. Revefenacin is a potent, lung-selective LAMA that prevents bronchoconstriction by inhibiting muscarinic M3 receptors in airway smooth muscle. Revefenacin received its first global approval on 9 November 2018 in the USA for maintenance treatment of patients with COPD [4]. The approved dosage of revefenacin inhalation solution is 175 µg once daily, delivered via a standard jet nebulizer connected to an air compressor [3]. Revefenacin should not be used to relieve acute symptoms of COPD (i.e. as a rescue therapy) and should not be initiated in patients experiencing acutely deteriorating or potentially life-threatening episodes of COPD. Revefenacin should be discontinued immediately if patients develop paradoxical bronchospasm or immediate hypersensitivity reactions [3].

In preliminary development, revefenacin was undergoing clinical investigation in South Africa, New Zealand, Germany and the United Kingdom for the treatment of COPD; however, no recent development reports have been identified outside the USA.

1.1 Company Agreements and Patent Information

GlaxoSmithKline (GSK) licensed revefenacin from Theravance in 2004, but the right was returned to Theravance in
February 2009 due to its incompatibility with GSK’s proprietary inhaler device [5].

In June 2014, Theravance Biopharma was separated from Theravance (now Innoviva) to focus on the discovery, development and commercialization of small molecule medications across multiple therapeutic areas, including respiratory disease [6].

In January 2015, Theravance Biopharma entered into a collaboration with Mylan, for the development and commercialization of nebulized revefenacin for the treatment of COPD and other respiratory diseases [7]. Under the terms of the agreement, Mylan provides reimbursements for the costs of the US registrational program that is led by Theravance Biopharma; following the approval of the first new drug application in the USA, costs will be shared between the two companies [7]. Mylan has the global right to develop and commercialize nebulized revefenacin (except in China where Theravance Biopharma holds the full rights). Theravance Biopharma retains co-promotion rights in the USA under a profit-sharing arrangement (65% Mylan and 35% Theravance Biopharma) [8]. Theravance Biopharma is eligible to receive potential development and sales-based milestone payments (up to $220 million) in the USA plus double-digit royalties on net sales on territories outside of the USA. Theravance Biopharma retains the global right to revefenacin delivered through other dosage forms (e.g. MDI or DPI) although Mylan has certain rights of first negotiation with respect to future development and commercialization of revefenacin delivered other than via a nebulized inhalation product [7, 8]. Theravance Biopharma currently has patent protection for the nebulized formulation of revefenacin in the USA until 2025 (not including any patent term extensions that may be available).

2 Scientific Summary

2.1 Pharmacodynamics

Revefenacin is a potent LAMA that binds competitively and reversibly to the M3 receptors in the airway smooth muscle, thereby inhibiting bronchoconstriction and increasing bronchodilation [9]. In vitro, in Chinese hamster ovary-K1 cell membrane fractions stably expressing human recombinant muscarinic receptors M1–M5, revefenacin shows similar, high affinity (pKi 8.2–9.8) and displays kinetic functional selectivity (i.e. slower receptor dissociation) for human M3 compared with M2 receptors. The dissociation half-life was significantly (p < 0.05) longer for M3 than M2 receptors (82 vs. 6.9 mins) [9]. Following inhalation of revefenacin, bronchodilation is mainly site-specific [3].

In vitro, revefenacin antagonized muscarinic acetylcholine cholinergic receptor-mediated contractions for up to 24 h in rat and guinea pig tracheal tissues, as well as in human airway muscle tissues [9]. In anesthetized dogs and rats, single or repeated dosing (once daily for 7 days) of inhaled revefenacin provided a dose-dependent bronchoprotection against acetylcholine- or methacholine-induced...
bronchoconstriction [10]. The bronchoprotective effect was seen as early as 5 min post-dose and was sustained for up to 24 h. In these animal models, repeated once daily dosing of revefenacin also exhibited superior functional lung selectivity index over repeated dosing of tiotropium (i.e. once daily LAMA therapy) and glycopyrronium (i.e. twice daily LAMA therapy). After repeated dosing, the bronchoprotective potency of revefenacin and tiotropium was maintained, unlike glycopyrronium [10].

In a thorough, positive-controlled QT study in healthy subjects (n = 48), a therapeutic (175 µg) or supratherapeutic (700 µg) dose of inhaled revefenacin did not prolong QTc interval to any clinically meaningful extent [11]. In two 12-week phase III efficacy trials and a 52-week safety phase III trial in patients with moderate to very severe COPD (Sect. 2.3), long-term use of inhaled revefenacin (up to 52 weeks) had no clinically relevant effect on QTc interval compared with placebo [11].

### 2.2 Pharmacokinetics

After inhaled administration in COPD patients, revefenacin is rapidly metabolized in the liver to its major active metabolite (THRX-195518), with its plasma exposure [based on area under the plasma concentration-time curve (AUC)] being 4- to 6-fold greater than revefenacin [3, 12]. The potency of the metabolite THRX-195518 at muscarinic receptors is one-third to one-tenth lower relative to revefenacin [3]. Exposures to revefenacin and its active metabolite increased in a slightly greater than dose-proportional manner across a dose range of 88–700 µg when administered as a single or multiple inhaled doses [3]. The pharmacokinetic profile of revefenacin was best described with a 2-compartment model with first-order absorption and elimination [13].

In healthy volunteers and in COPD patients, peak plasma concentrations (C\text{max}) of revefenacin and its active metabolite occurred 14–41 min after start of nebulization; the absolute bioavailability of revefenacin following oral administration is very low (<3%) [3]. Following repeated administration of inhaled revefenacin, steady state was reached within 7 days, with <1.6-fold accumulation. In COPD patients, exposure (i.e. C\text{max} and AUC) to revefenacin was ≈60% lower but exposure to active metabolite was ≈2-fold higher than healthy volunteers [3].

Following intravenous administration of revefenacin in healthy volunteers, the mean steady-state volume of distribution was 218 L [3]. In vitro human plasma protein binding for revefenacin and its active metabolite averaged 71 and 42% [3].

After intravenous administration of revefenacin in healthy volunteers, ≈ 54% of the radioactivity was recovered in faeces (≈19% as the active metabolite) and 27% was recovered in urine [3, 14]. Following oral administration of a single dose of radiolabelled revefenacin, 88% of the radioactivity was recovered in faeces and < 5% was recovered in urine, suggesting poor oral absorption. Following inhaled administration in COPD patients, revefenacin and its active metabolite were minimally excreted via the renal route (< 1%) [3, 14]. The elimination half-life of revefenacin and its active metabolite after once daily dosing of inhaled revefenacin in COPD patients is 22–70 h [3].

A population pharmacokinetic analysis indicated that systemic exposure of revefenacin and its active metabolite are not significantly affected by age, gender, smoking status or weight [3]. Revefenacin is not recommended in patients with any degree of hepatic impairment as the safety of revefenacin has not been studied in COPD patients with mild to severe hepatic impairment [3]. Dosage adjustments are not required in patients with renal impairment. However, C\text{max} and AUC of revefenacin and its active metabolite increased up to 2.5-fold in patients with severe renal impairment; thus, monitoring for systemic antimuscarinic adverse events (AEs) is recommended in this population [3].

Coadministration of revefenacin with other anticholinergic medications should be avoided as concomitant use of these drugs may increase the risk of anticholinergic AEs (e.g. acute narrow-angle glaucoma, urinary retention) [3]. Coadministration of revefenacin with OATP1B1 and OATP1B3 inhibitors (e.g. rifampicin, cyclosporine) is also not recommended as it may increase systemic exposure of the active metabolite of revefenacin [3].

### 2.3 Therapeutic Trials

#### 2.3.1 Phase III Trials

Twelve weeks of revefenacin treatment effectively improved lung function [as assessed by forced expiratory volume in 1 s (FEV\text{1})] in patients aged >40 years with moderate to very severe COPD in two replicate, randomized, double-blind, placebo-controlled trials (NCT02459080 and NCT02512510) [15, 16]. Pooled data showed that, compared with placebo (n = 428), patients treated with once daily revefenacin 88 µg (n = 425) or 175 µg (n = 402) via a standard jet nebulizer experienced significantly (p ≤ 0.001) greater improvements in the 24 h change from baseline in trough FEV\text{1} on day 85 [between-group differences (BGDs) 118 mL (95% CI 86–150 mL) and 145 mL (95% CI 113–178 mL)] (primary endpoint) and the overall treatment effect in trough FEV\text{1} over 12 weeks (88 µg: BGD 112 mL, 95% CI 106–118 mL; 175 µg: BGD 139 mL, 95% CI 133–145 mL) [15, 16]. The treatment benefits of either dosage of revefenacin were consistently seen in subgroup of patients with higher risk of COPD exacerbations, including in the elderly (>65 years), those receiving concomitant therapy (LABA or ICS) and those with very severe COPD.
[classified as Global Initiative for COPD category D (GOLD D) \((p \text{ value not reported})\) [17]. According to a 24 h serial spirometry substudy \((n = 264)\) on day 84, revefenacin 88 or 175 µg were associated with significant \((p < 0.01)\) improvements in trough FEV\(_1\) at all time points compared with placebo (BGDs 111–185 and 154–269 mL) [18], indicating bronchodilation was sustained for at least 24 h post-dose. Both dosages of revefenacin were associated with a nominal reduction in the annualized rate of COPD exacerbations but the differences compared with the placebo group were not statistically significant [19].

Significant \((p \leq 0.0203)\) improvements from baseline in health-related quality of life indicators [assessed by St. George’s Respiratory Questionnaire (SGRQ) and COPD Assessment Test (CAT)] with either revefenacin dosage versus placebo were only shown in one (NCT02459080) of the two replicate trials. Despite improvements from baseline in the revefenacin 175 µg group, a greater than expected response in the placebo group prevented statistically significant differences in the second trial (NCT02512510) [20]. At entry, patients in the two replicate trials had a mean age of 63.7 years, a mean smoking history of 52.3 pack-years \((49\% \text{ of patients were current smokers [16]}\) and the mean predicted FEV\(_1\) was 54.4%; 38.1% of patients were taking concomitant LABA or LABA/ICS therapy and 34.5% had very severe COPD (classified as GOLD D) [15]. These trials included patients with moderate to very severe COPD (post-ipratropium FEV\(_1\) \(\leq 80\%\) of predicted and \(\geq 700\) mL), a smoking history of \(\geq 10\) pack-years and a FEV\(_1\)/forced vital capacity (FVC) ratio of < 0.7 [3]. Patients with narrow-angle glaucoma, unstable cardiac disease or symptomatic prostatic hypertrophy or bladder outlet obstruction were excluded [3].

In a randomized, double-blind phase IIb study (NCT03095456), 207 patients with moderate to very severe COPD and suboptimal PIFR \((< 60 \text{ L/min})\) were randomized to receive revefenacin 175 µg once daily via a standard jet nebulizer or tiotropium once daily via HandiHaler\textsuperscript{\textregistered} for 28 days [17, 21]. On day 29, revefenacin and tiotropium effectively improved trough FEV\(_1\) and FVC from baseline, with improvements numerically favouring revefenacin over tiotropium; however, these differences did not reach statistical significance \((FEV\(_1\): \(p = 0.481\); FVC: \(p = 0.104\)). In a prespecified subgroup analysis, revefenacin significantly \((p \leq 0.0407)\) improved trough FEV\(_1\) and FVC from baseline compared with tiotropium in patients with severe to very severe COPD (i.e. FEV\(_1\) < 50% of predicted), which accounted for \(\approx 80\%\) of enrolled patients [17, 21].

In efficacy analyses from a 52-week, randomized, partially blind, active-controlled safety phase III study (NCT02518139; \(n = 1055\) [17, 22]), relative to placebo, inhaled revefenacin 88 or 175 µg once daily provided long-term improvement in trough FEV\(_1\) from baseline in the subgroups of patients with moderate to very severe COPD who received concomitant LABA or ICS therapy or those aged 65 years \((p \text{ value not reported})\) [17]. The study also included open-label tiotropium (active control) and assessed the impact of revefenacin and tiotropium on the annualized rate of COPD exacerbations; there were no significant differences across treatment groups [19]. At entry, patients had a mean age of 64 years and a mean predicted FEV\(_1\) of 54%; 46% of patients were current smokers and 50% were taking concomitant LABA (2.2%) or LABA/ICS (47.8%) therapy [23].

### 2.3.2 Phase II trials

Compared with placebo, in a randomized, double-blind, placebo-controlled, dose-ranging phase II study in patients with moderate to severe COPD (NCT02040792; \(n = 355\)), least squares mean (LSM) trough FEV\(_1\) from baseline on day 28 significantly improved in the revefenacin 88, 175 and 350 µg once daily groups (134.2–155.0 vs. – 32.4 mL;
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all $p < 0.001), but not with revefenacin 44 μg once daily [2]. At 4 h post-dose, ≥ 100 mL increase in FEV$_1$ from baseline was evident in > 80% of patients receiving a revefenacin dosage ≥ 88 μg/day. Inhaled revefenacin dosages ≥ 88 μg/day also significantly ($p < 0.005$) decreased the use of rescue medication (i.e. albuterol) compared with placebo. Revefenacin 350 μg/day did not show additional efficacy benefit over revefenacin 175 μg/day [2].

NCT02040792 enrolled patients (aged 40–75 years) with moderate to severe COPD (pre-bronchodilator FEV$_1$ 30–80% of predicted and post-ipratropium FEV$_1$/FVC ratio of < 0.7) and a smoking history of > 10 pack-years [2]. At entry, patients had a mean age of 61.9 years, a mean smoking history of 39.5 pack-years (53.7% of patients were current smokers) and a mean predicted FEV$_1$ of 43.6%; 36.7% of patients were currently using ICS. Patients were randomized to receive inhaled revefenacin 44, 88, 175 or 350 μg once daily or placebo for 28 days [2].

Inhaled revefenacin 22–700 μg once daily via a standard jet nebulizer provided rapid and sustained bronchodilation in patients with moderate to severe COPD in two randomized, double-blind, placebo-controlled, dose-finding phase II studies (NCT03064113 and NCT01704404) [12]. In NCT03064113 ($n = 32$), a significant ($p < 0.001$) increase from baseline in the LSM peak (0–6 h) FEV$_1$ was observed with single dose inhaled revefenacin (350 or 700 μg) or ipratropium 500 μg (active control) compared with placebo (1838.9–1867.3 vs. 1676.7 mL). The onset and the magnitude of the bronchodilator effect with revefenacin was similar to that seen with ipratropium. In NCT01704404 ($n = 59$), a 7-day treatment course of revefenacin (22, 44, 88, 175, 350 or 700 μg once daily) significantly ($p \leq 0.006$) increased LSM trough FEV$_1$ from baseline on day 7 compared with placebo (91.2–151.9 vs. 37.8 mL). In both studies, the bronchodilator effect of revefenacin was maintained for at least 24 h post-dose. In addition, a minimal clinically important difference for FEV$_1$ was achieved in patients receiving revefenacin doses ≥ 88 μg and the dose response appeared to plateau at doses ≥ 175 μg [12].

NCT03064113 and NCT01704404 enrolled patients (aged 40–75 years) with moderate to severe COPD (prebronchodilator FEV$_1$ 35–80% of predicted and FEV$_1$/FVC ratio of < 0.7) and a smoking history of > 10 pack-years [12]. At entry, patients in these trials had a mean age of 62.0 and 63.9 years and a mean predicted FEV$_1$ of 50.5 and 47.2%. NCT03064113 was conducted in South Africa and New Zealand and NCT01704404 was conducted in United Kingdom, Northern Ireland and New Zealand [12].

Key clinical trials of revefenacin, sponsored by Theravance Biopharma

| Drug(s) | Indication | Phase | Status | Location(s) | Identifier |
|---------|------------|-------|--------|-------------|------------|
| Revefenacin, placebo | Adults (aged > 40 years) with moderate to very severe COPD | III | Completed | USA | NCT02459080 |
| Revefenacin, placebo | Adults (aged > 40 years) with moderate to very severe COPD | III | Completed | USA | NCT02512510 |
| Revefenacin, tiotropium | Adults (aged > 40 years) with moderate to very severe COPD | III | Completed | USA | NCT02518139 |
| Revefenacin, tiotropium | Adults (aged > 40 years) with moderate to very severe COPD and a low PIFR (< 60 L/min) | IIIb | Completed | USA | NCT03095456 |
| Revefenacin, formoterol, placebo | Adults (aged > 40 years) with moderate to very severe COPD | IIIb | Completed | USA | NCT03573817 |
| Revefenacin, placebo | Adults (aged > 40 years) with moderate to very severe COPD | II | Completed | USA | NCT02040792 |
| Revefenacin, ipratropium, placebo | Adults (aged 40–75 years) with moderate to severe COPD | II | Completed | New Zealand, South Africa | NCT03064113; ANZCTR U1111-1120-8290 |
| Revefenacin, placebo | Adults (aged 40–75 years) with moderate to severe COPD | II | Completed | United Kingdom, Northern Ireland and New Zealand | NCT01704404 |

*COPD* chronic obstructive pulmonary disease, *PIFR* peak inspiratory flow rate

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2.4 Adverse Events

Revefenacin inhalation solution was generally well tolerated in patients with moderate to very severe COPD in clinical trials [3]. In a pooled analysis of NCT02459080 and NCT02512510, at least one any-grade AE was reported in 53.4, 50.7 and 48.3% of patients receiving revefenacin 88 or 175 µg or placebo once daily, respectively [24]. The most common AEs occurring in ≥ 2% of patients with a higher incidence in revefenacin 88 or 175 µg than placebo recipients were cough (4.0 and 4.2 vs. 4.0%), headache (4.9 and 4.0 vs. 2.6%), upper respiratory tract infection (4.7 and 2.7 vs. 2.1%), nasopharyngitis (3.3 and 3.7 vs. 2.1%) and back pain (1.4 and 2.2 vs. 0.7%). The incidence of antimuscarinic AEs and serious AEs were low and similar across treatment groups [24, 25]. Where reported, AEs leading to treatment discontinuation occurred in 13% of revefenacin 175 µg recipients and 19% of placebo recipients [3].

The tolerability profile of revefenacin in patients with moderate to very severe COPD in the 52-week safety phase III study (NCT02518139; Sect. 2.3) was generally similar to that observed in the two 12-week studies and revealed no new safety signals [3]. The frequency and severity of treatment-emergent AEs, as well as instances of major adverse cardiovascular events were generally similar across treatment groups [23].

2.5 Ongoing Trials

Currently, there are no ongoing trials identified for revefenacin inhalation solution.

3 Current Status

Revefenacin inhalation solution received its first global approval on 9 November 2018 in the USA for the maintenance treatment of patients with COPD.

Compliance with Ethical Standards

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