Research paper

Randomized study of the safety and pharmacodynamics of inhaled interleukin-13 monoclonal antibody fragment VR942

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A B S T R A C T
Background: Interleukin-13 (IL-13) is a key mediator of T-helper-cell-type-2 (Th-2)-driven asthma, the inhibition of which may improve treatment outcomes. We examined the safety, pharmacokinetics, pharmacodynamics, and immunogenicity of VR942, a dry-powder formulation containing CDP7766, a high-affinity anti-human-IL-13 antigen-binding antibody fragment being developed for the treatment of asthma.

Methods: We conducted a phase 1, randomized, double-blind, placebo-controlled, ascending-dose study at Hammersmith Medicines Research, London, UK, which is now complete. Healthy adults aged 18–50 years (n = 40) were randomized 3:1 to a single inhaled dose of VR942 0.5, 1.0, 5.0, 10, or 20 mg, or placebo. Adults aged 18–50 years who were diagnosed with asthma for ≥6 months before screening, and had forced expiratory volume in 1 s (FEV1) and forced vital capacity (FVC) values ≥70% of the predicted values at screening (n = 45), were randomized to once-daily inhaled VR942 0.5 or 10 mg, or placebo (2:2:1), or VR942 20 mg or placebo (3:2), for 10 days. All participants were randomized to receive VR942 or placebo based on a randomization list prepared by an independent HMR statistician using SAS® software (SAS Institute, Cary, NC). The primary outcome was safety and tolerability of VR942 (safety population, defined as all who received at least one dose of VR942 or placebo). This study is listed on ClinicalTrials.gov (NCT02473939).

Findings: In the VR942 and placebo groups, treatment-emergent adverse events (TEAEs) were reported in 10/30 (33%) and 0/10 (0%) healthy participants, and in 16/29 (55%) and 9/16 (56%) participants with asthma, respectively. Mild intermittent wheezing occurred in 7 participants (VR942 20 mg, n = 4; corresponding placebo, n = 3), resolving spontaneously within 1 h. All TEAEs were mild or moderate; there were no deaths, serious adverse events, or clinically significant changes in vital signs, electrocardiograms, or laboratory parameters. There was no clinically significant immunogenicity, with only one participant with asthma considered positive for treatment-related immunogenicity for CDP7766.

Interpretation: This study, considered to be the only example of a dry powder anti-IL-13 fragment antibody being administered via inhalation, demonstrated that single and repeat doses were well tolerated over a period of up to 10 days in duration. Rapid and durable inhibition of fractional exhaled nitric oxide (FeNO) (secondary outcome) provided evidence of pharmacological engagement with the IL-13 target in the airways of participants diagnosed with mild asthma. These data, together with the numerical improvements observed for predose FEV1, justify further clinical evaluation of VR942 in a broader population of patients with asthma, and continue to support the development of an inhaled anti-IL-13 antibody fragment as a potential future treatment that is alternative to monoclonal antibodies delivered via the parenteral route.

Abbreviations: AE, Adverse event; ANCOVA, Analysis of covariance; AUC0–75, Area under the concentration–time curve to postdose day 10; DPI, Dry-powder inhaler; FeNO, Fractional exhaled nitric oxide; FEV1, Forced expiratory volume in 1 s; FVC, Forced vital capacity; HMR, Hammersmith Medicines Research; ICS, Inhaled corticosteroid; IL, Interleukin; IL-4Rα, Interleukin-4 receptor; MAb, Monoclonal antibody; NO, Nitric oxide; PD, Pharmacodynamic; PK, Pharmacokinetic; ppb, Parts per billion; SABA, Short-acting β2-agonist; SD, Standard deviation; TEAE, Treatment-emergent adverse event.

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Effective treatments for patients whose asthma symptoms remain uncontrolled despite corticosteroid therapy are lacking. Pro-inflammatory cytokines, such as interleukin-13 (IL-13), play an important role in asthma, and several IL-13 inhibitors have been investigated for the treatment of asthma. Despite IL-13 levels being increased in the lung, these agents have been delivered systemically; the inhaled delivery of VR942 represents an alternative, non-invasive, novel approach offering the potential to deliver greater therapeutic benefit whilst maintaining a good tolerability profile. Results from this study continue to support this rationale and justify continued development of VR942 in a broader population of patients with asthma.

1. Introduction

Asthma has a marked impact on patients and society [1], especially when symptoms are uncontrolled [2]. Approximately 300 million people have asthma globally [3], of whom about 10% have severe and persistent disease [4], characterized by uncontrolled symptoms despite treatment with corticosteroids or other controller medications [1].

Cytokines play a cardinal role in the pathogenesis and severity of asthma [5]. One such is interleukin (IL)-13 [6] that signals primarily through the type 2 IL-4 receptor (IL-4R; composed of IL-13Rα1 and IL-4Rα subunits [7]) and contributes to many key features of asthma, including mucus production, eosinophilic airway inflammation, immunoglobulin E synthesis, bronchial fibrosis, and airway hyper-responsiveness [8]. IL-13 is overexpressed in the sputum of patients with asthma, particularly those with severe disease [9], and was recently proposed as a biomarker for evaluating asthma control [10]. Elevated IL-13 levels in patients with uncontrolled asthma despite corticosteroid treatment also support the notion of a potential association between IL-13 and treatment resistance [9], implicating IL-13 signaling as a therapeutic target in severe asthma.

IL-13 stimulates inducible nitric oxide (NO) synthase in vitro, driving production and release of NO from lung epithelial cells [11,12]. Parenteral treatment with monoclonal antibodies (MAbs) that inhibit IL-13 signaling reduces fractional exhaled NO (FeNO) levels [13–15]. Thus, FeNO level may serve as a biomarker for lung inflammation [16]. Clinical studies of anti-IL-13 MAbs have demonstrated FeNO’s potential use as a biomarker for pharmacological activity of these agents in the lungs of patients with mild asthma, including inhaled corticosteroid (ICS)-naïve individuals [15,17]. Randomized, placebo-controlled studies with the anti-IL-13 MAbs lebrikizumab and tralokinumab, and the anti-IL-4Rx MAb dupilumab, showed that parenteral administration of these agents reduced exacerbation rates and improved lung function in patients with uncontrolled T-helper-cell-type 2-driven asthma despite corticosteroid therapy, while being well tolerated [13,14,18]. In replicate phase 3 studies in patients with uncontrolled asthma, lebrikizumab and tralokinumab reduced asthma exacerbations in biomarker–high patients compared with placebo; although the reductions did not consistently show statistical significance in all patients across both studies [19,20].

2. Materials and methods

2.1. Study design and participants

This was a two-part, phase 1, randomized, double-blind, placebo-controlled, ascending-dose study. In part 1, healthy participants received a single dose of VR942; in part 2, participants with asthma received multiple doses of VR942. The trial was performed at Hammersmith Medicines Research (HMR), London, UK, in compliance with European Union Directives 2001/20/EC and 2005/28/EC, the Medicines for Human Use (Clinical Trials) Regulations 2004 and current amendments, and the Declaration of Helsinki (Brazil Revision, 2013). Ethical approval was obtained from the Scotland A Research Ethics Committee. All participants gave written informed consent for participation and for data to be entered into The Over Volunteering Prevention System [23]. This study is listed on ClinicalTrials.gov (NCT02473939).

Eligible for inclusion were men and women (non-child-bearing potential) aged 18–50 years, ≥50 kg in weight, a body mass index of 18.0–31.0 kg/m², and a peak inspiratory flow rate of ≥60 L/min for ≥2 s. Healthy participants had to have forced expiratory volume in 1 s (FEV₁) forced vital capacity (FVC) values >80% of the predicted values at screening, based on reference equations [24].

Participants with asthma were required to have a history of asthma (as defined by the Global Initiative for Asthma guidelines [11]) for ≥6 months before screening, FEV₁ and FVC values ≥70% of the predicted values at screening (assessed without β₂-agonist treatment [no use in the previous 6 hours]) and based on reference equations [24], and an FeNO level ≥35 parts per billion (ppb). Owing to challenges in recruiting ICS-naïve participants, the protocol was amended to include participants receiving stable low-dose ICSs (beclomethasone ≤500 μg/day or equivalent). For those taking ICS, FeNO values at screening and predosing on day 1 had to be within 20% of each other.

Exclusion criteria included current smoker, a smoking history of more than 10 pack-years, respiratory tract infection within 4 weeks of screening, use of prescription or over-the-counter medicines (other than paracetamol) 7 days before trial dosing, and use of a MAB 6 months before dosing. The use of short-acting β₂-agonists (SABAs) was permitted; individuals who had used leukotriene antagonists in the 2 weeks before screening or long-acting β₂-agonists (LABAs) at any time before

Agents that both inhibit IL-13 signaling and are deliverable directly to the lungs are of particular interest. Pitrakinra, an IL-4 mutein that binds to the IL-4Rα subunit and reduces IL-4– and IL-13–mediated inflammation in patients with asthma, was the first such drug to support a role for inhaled delivery [21]. VR942 (UCB4144; UCB, Brussels, Belgium) is formulated for inhalation as a dry powder and is being developed for the treatment of uncontrolled asthma. It contains CDP7766, a humanized, high-affinity, neutralizing, anti-human–IL-13 antibody fragment that binds to IL-13, preventing binding to the IL-13Rα1 subunit. In cynomolgus monkeys, nebulized CDP7766 was well tolerated at all doses (0.1–60 mg/animal/day) and caused dose-dependent inhibition of bronchoconstriction and reduction in levels of proinflammatory cytokines induced by an inhaled allergen (Ascaris suum), for up to 24 h [22].

Here, we report data from the first-in-human, phase 1, randomized, double-blind, placebo-controlled, ascending-dose study, which aimed to assess the safety and tolerability of VR942 in healthy participants and participants with asthma.

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screening were excluded. See the supplementary appendix for further exclusion criteria.

All participants were randomly assigned to receive VR942 or placebo based on a randomization list prepared by an independent HMR statistician using SAS® software (SAS Institute, Cary, NC). In part 1, 8 healthy participants were allocated to each VR942 dose group (0.5 mg, 1.0 mg, 5.0 mg, 10 mg, 20 mg) and were randomized 3:1 to receive a single dose of VR942 or placebo. In part 2, 9 participants with asthma were allocated to each of the 0.5 mg and 10 mg groups, and were randomized 2:1 to receive VR942 or placebo; 27 participants were allocated to the 20 mg group and were randomized 3:2 to receive VR942 or placebo.

Placebo and active treatments for each group were identical in appearance. Doses were administered as 1–4 separate inhalations, matched for placebo and active treatment within each cohort. HMR site staff enrolled participants, and all trial personnel, participants, and the study sponsor were blinded to treatment allocation.

2.2. Procedures

VR942 (Vectura Ltd, Chippenham, UK) was administered on a single occasion at nominal doses of 0.5 mg (1 × 0.5 mg inhalation), 1.0 mg (2 × 0.5 mg inhalations), 5.0 mg (1 × 5.0 mg inhalation), 10 mg (2 × 5.0 mg inhalations), and 20 mg (4 × 5.0 mg inhalations). The formulation was contained in a unit-dose blister and delivered via the multidose DPI dry-powder inhaler (DPI; Vectura Ltd, Chippenham, UK). See the supplementary appendix for information about the sentinel approach to dosing.

Participants were admitted to the clinic the day before dosing (day −1) following a 4-week screening period. In part 1, healthy participants received a single dose of VR942 or placebo and remained in the clinic until all postdose assessments were complete (72 h postdose, day 4). After the final in-clinic assessment, participants were discharged and followed-up on visits on days 14 and 28. In part 2, participants with asthma received once-daily morning doses of VR942 or placebo for 10 consecutive days (days 1–10) and were assessed in clinic up to 96 h after the final dose (day 14) and as outpatients at day 28 (Fig. 1).

2.3. Outcomes

The primary objective was to evaluate the safety and tolerability of single doses of VR942 (0.5 mg, 1.0 mg, 5.0 mg, 10 mg, and 20 mg) in healthy participants, and the safety and tolerability of multiple ascending doses of VR942 (0.5 mg, 10 mg, and 20 mg once daily for 10 days) in participants with asthma. Secondary aims included determination of the pharmacokinetic (PK) profile and immunogenicity of CDP7766 (all participants), and the pharmacodynamic (PD) effect of VR942 on FeNO level (participants with asthma).

Vital signs (systolic and diastolic blood pressure, heart rate, temperature, and respiratory rate) were recorded, and 12-lead electrocardiograms (ECGs), physical examinations, spirometry (FEV1 and FVC), pulse oximetry, and laboratory assessments (routine hematology, biochemistry, urinalysis) were performed predose and at specified postdose intervals up to the final follow-up visit (Fig. 1). FEV1 and FVC were measured using a Microlab MK8 spirometer according to ATS/ERS guidelines [25]. See the supplementary appendix for further details regarding the assessment of vital signs and ECGs. Tolerability was assessed throughout the study and included recording of all adverse events (AEs), and comments recorded by HMR site staff on an inhalation checklist regarding participants’ tolerability of VR942 during dosing.

In participants with asthma, FeNO levels were measured at screening and on day −1, days 1–10 (predose, 0.5, 2, 6, and 12 h postdose), days 11–14 (24, 48, 72, and 96 h after the last dose), and day 28 (follow-up). FeNO was measured using a NIOX MINO® analyzer according to ATS/ERS guidelines [25]. On days when spirometry was also scheduled at the same time point, the FeNO level was measured first.

Blood samples for PK assessments in part 1 were taken predose and at 0.5, 1, 1.5, 2, 3, 4, 6, 8, 24, 48, and 72 h postdose. In part 2, samples were collected predose and at 0.5, 1, 1.5, 2, 3, 4, 6, and 8 h postdose on days 1 and 10, predose on days 2 and 9, and on days 11–14 (at 24, 48, 72, and 96 h after the last dose on day 10). A validated ligand plasma binding assay method (quantitation range: 99.6–2000 ng/ml; PRA Health Sciences, Assen, Netherlands) was used to measure free CDP7766. In the assay, CDP7766 present in human serum binds to biotinylated human IL-13 immobilized on streptavidin-coated plates. Bound drug was detected using a goat anti-Human Kappa antibody conjugated to ruthenium SULPHO TAG, and the resulting electrochemiluminescent signal was measured on the Meso Scale Discovery Sector Imager 6000. See the supplementary appendix for further information regarding pharmacokinetics and immunogenicity assessments.

Blood samples for evaluating anti-CDP7766 antibodies were taken from all participants predose on day 1, and on days 14 and 28 after the first dose of VR942 or placebo. Immunogenicity tests were performed by PRA International (Groningen, Netherlands), using a validated assay. With a rabbit monoclonal anti-CDP7766, a sensitivity of 1.22 ng/ml was demonstrated. Further details of the methodology for evaluating anti-CDP7766 antibodies can be found in the supplementary appendix.

2.4. Statistical analyses

To assess safety and tolerability while minimizing the number of participants exposed to VR942, the target sample size was 8 participants per dose group for part 1 (VR942, n = 6; placebo, n = 2) and 9 participants for each of the 0.5 and 10 mg dose groups in part 2 (VR942, n = 6; placebo, n = 3). For the 20 mg dose group, the target sample size was 35 participants (VR942, n = 21; placebo, n = 14). This was adjusted (protocol amendment) following a review of data from participants in the VR942 0.5 mg and 10 mg dose groups. The previously assumed common standard deviation (SD) of 30% was reduced to 24%, meaning 25 participants (VR942, n = 15; placebo, n = 10) would be expected to provide >80% power to detect a difference of 30% change from baseline in FeNO level between the VR942 and placebo groups, given a 2-sided t-test with a 5% significance level.

Demographic, safety, and tolerability data were summarized descriptively by treatment group (VR942 and placebo group) and time point for the safety population (all participants who received at least 1 dose of study treatment). Absolute FeNO values and mean changes from baseline were summarized by VR942 and pooled placebo groups for the PD population (all participants who received at least 1 dose of study treatment and who had at least 1 postdose FeNO measurement recorded). Changes from baseline in FeNO level at 0.5 and 2 h post last-dose on day 10 were compared between the VR942 20 mg and placebo groups using an analysis of covariance (ANCOVA), adjusting for treatment group and baseline FeNO level. Post hoc ANCOVAs were performed for (1) the same endpoint and time points for the placebo group and all active treatment groups, and (2) changes from baseline in FeNO level predose on day 10 for the placebo group and all active treatment groups. Weighted least-squares means for FeNO values were obtained from the ANCOVA of change from baseline to postdose day 10 (AUC0–10), adjusting for treatment group and baseline FeNO level. AUC0–10 was normalized before analysis by dividing by the time interval from dosing on day 1 to 12 h postdose on day 10 (nominally 9.5 days).

3. Results

3.1. Participants

In part 1, 50 healthy individuals were screened for inclusion (May 15, 2015 to July 24, 2015). Of these, 40 were randomized to receive treatment; 1 participant in the VR942 0.5 mg group withdrew from
Following administration of a single dose of VR942 in healthy participants, 18 AEs were reported by 11/30 participants (37%) across all VR942 groups (Table 2). In total, 15 treatment-emergent adverse events (TEAEs) were reported by 10/30 participants (33%). Two TEAEs were considered to be related to study medication: mild aphthous ulcer, reported by 1 participant in the VR942 1.0 mg group, and moderate cough, reported by 1 participant in the VR942 10 mg group. All TEAEs were mild or moderate, and included nervous system disorders (n = 4), gastrointestinal disorders (n = 3), musculoskeletal and connective tissue disorders (n = 2), and respiratory, thoracic, and mediastinal disorders (n = 2). There were no AEs or TEAEs in the placebo group, of which 13 were TEAEs. In total, 16 TEAEs were considered by the investigator to be related to study medication. Mild intermittent wheezing was reported by 4 out of 17 participants who received VR942 20 mg and by 3 out of 9 participants in the corresponding placebo group, all of whom received study treatment as 4 separate inhalations. These events were mainly reported on day 1 within 5 min of dosing, and resolved spontaneously within 1 h, although a further 7 episodes of mild, intermittent, and transient wheezing occurred throughout the 10-day dosing period. As on day 1, these events resolved spontaneously. One case of presyncope was reported, which occurred more than 4 h after the first dose of VR942 20 mg and was considered by the investigator to be possibly related to the device owing to the inhalation maneuver used by this participant. Other TEAEs occurring in more than 1 participant in any group were headache and rhinorrhea (Table 3). None of the vital signs, ECG or laboratory values or changes during the study were considered to be clinically significant by the investigator. A summary of the vital signs and ECG data can be found in the supplementary appendix.

Following administration of VR942 0.5–20 mg, 29 AEs were reported by 16/29 participants with asthma (55%), 28 of which were TEAEs (Table 3). In all, 15 AEs in 9/16 participants (56%) were reported in the placebo group, of which 13 were TEAEs. In total, 16 TEAEs were considered by the investigator to be related to study medication. Mild intermittent wheezing was reported by 4 out of 17 participants who received VR942 20 mg and by 3 out of 9 participants in the corresponding placebo group, all of whom received study treatment as 4 separate inhalations. These events were mainly reported on day 1 within 5 min of dosing, and resolved spontaneously within 1 h, although a further 7 episodes of mild, intermittent, and transient wheezing occurred throughout the 10-day dosing period. As on day 1, these events resolved spontaneously. One case of presyncope was reported, which occurred more than 4 h after the first dose of VR942 20 mg and was considered by the investigator to be possibly related to the device owing to the inhalation maneuver used by this participant. Other TEAEs occurring in more than 1 participant in any group were headache and rhinorrhea (Table 3). None of the vital signs, ECG or laboratory values or changes during the study were considered to be clinically significant by the investigator. A summary of the vital signs and ECG data can be found in the supplementary appendix.

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3.3. PD effects

VR942 treatment markedly decreased FeNO levels in participants with asthma. The mean FeNO level decreased by approximately 22%
from baseline to predose on day 2 (10 mg and 20 mg groups) compared with a reduction of approximately 4% in the placebo group (Fig. 3). Near-maximal reductions were observed on day 4 predose with VR942 10 mg and 20 mg (approximately 42% and 46% reductions, respectively), and on day 6 predose with placebo (approximately 28% reduction; Fig. 3). In all groups, reductions in mean FeNO level were maintained for 96 h after the final dose on day 10, and returned to baseline after the 14-day no-treatment period.

Statistically significant reductions were observed when comparing the effects of VR942 20 mg and placebo on FeNO levels from baseline to 0.5 h and 2 h postdose on day 10 (prespecified ANCOVA; p = .015 and p = .028, respectively, for VR942 versus placebo; Table 4). In a post hoc analysis comparing the effects of all active treatments versus placebo at these 2 time points, statistically significant FeNO reductions from baseline were demonstrated for the VR942 10 mg and 20 mg groups versus placebo (p = .010 and p = .016, respectively; Table 4).

Statistically significant reductions in FeNO were seen from baseline to predose on day 10 for the VR942 10 mg and 20 mg groups versus placebo (VR942 10 mg mean between-group difference [95% confidence interval]: −17.5 [−29.5, −5.5] ppb; p = .005; VR942 20 mg: −11.6 [−20.5, −2.7] ppb; p = .012; post hoc ANCOVA; Table 5). No statistically significant changes were observed between VR942 0.5 mg and placebo in either analysis. The change from baseline in FeNO AUC0–17.5 was significantly greater with VR942 10 mg and 20 mg than with placebo (supplementary appendix Table 2).

3.4. PK measurements and immunogenicity

Plasma levels of CDP7766 in healthy participants and in participants with asthma receiving VR942 20 mg were below the lower limit of quantification for the PK analytical assay (≤99.6 ng/ml), so the PK analysis could not be performed.

Seven of 40 healthy participants tested positive for CDP7766 antibodies, but none were considered positive for treatment-related immunogenicity; these participants also tested positive predose (inhibition ≥36.5%; an observation not considered unusual), and samples taken on days 14 and 28 did not show an increase of at least fourfold in titer. Of these 7 participants, only 1 had an AE, which was mild in severity (musculoskeletal pain in the right foot). In all, 9 of 45 participants with asthma tested positive for CDP7766 antibodies: 3/16 (18.8%) in the placebo group, 1/6 (16.7%) in the VR942 10 mg group, and 5/17 (29.4%) in the VR942 20 mg group. Only 1 of these had a negative predose sample (inhibition <36.5%) and was therefore considered positive for treatment-related immunogenicity for CDP7766 (20 mg). The other samples that tested positive predose did not show an increase of at least fourfold in titer on days 14 and 28. The participant considered positive for treatment-related immunogenicity for CDP7766 experienced one AE of mild continuous wheezing on day 1, which started within 3 min of dosing and resolved after 20 min, and one AE of mild intermittent wheezing, which started within 1 min of dosing on day 7 and resolved after 2 days. Of the 8 participants who had a positive predose sample, 6 (placebo, VR942 10 mg and VR942 20 mg all n = 2) experienced one or more AEs, which were all mild or moderate in intensity (headache, n = 4; bruise at canula site, drowsiness, intermittent cough, intermittent wheezing, itchy eyes, oral haemorrhage, pain at canula site, sore throat, all n = 1).

4. Discussion

This study evaluated the safety and tolerability of a single inhaled dose of VR942 in healthy participants (part 1) and the effects of VR942 once-daily dosing for 10 days in participants with asthma receiving SABA treatment only (with 1 exception receiving SABA plus low-dose ICS; part 2). VR942 0.5–20 mg was generally well tolerated in both populations, and rapid, dose-related sustained FeNO reductions were observed for VR942 10 and 20 mg doses in participants with asthma. There were no detrimental effects on predose FEV1, with VR942 treatment relative to placebo, and numerically greater increases in predose FEV1 were observed from baseline for VR942 0.5 and 20 mg compared with placebo.
VR942 was administered directly to the lungs via a unit-dose blister DPI. Safety findings from this study align with findings from phase 2 studies of systematically administered anti-IL-13 MAbs lebrikizumab, tralokinumab, GS679586, and the anti-IL-4R α MAb dupilumab, which were also well tolerated [13,18,26,27]. Compared with subcutaneous administration, inhalation may provide a more rapid onset of action at lower doses than systemic treatments and may be suitable for use in a broader range of patients [1]. Systemic exposure to CDP7766 was below the limit of quantification following repeat dosing of VR942 20 mg; further studies are needed to explore whether this translates into a reduced potential for systemic-related AEs.

In participants with asthma, a reduction of over 10% in FEV1 from predose to 10 min postdose on day 1 was reported in 6 participants in the VR942 20 mg group and 5 in the corresponding placebo group. All these FEV1 reductions had returned to normal by 60 min postdose without the need for treatment. The reductions in FEV1 were also associated with mild wheezing in 3 participants in the VR942 20 mg group and 2 receiving placebo, which resolved spontaneously. These decreases may have been due to the mass of powder and/or frequency of inhalation, as participants received study treatment as 4 separate inhalations.

### Table 1
Baseline demographic characteristics.

| Characteristic | Healthy participants (N = 40) | Placebo (n = 10) | VR942 0.5 mg (n = 10) | VR942 1 mg (n = 10) | VR942 5 mg (n = 10) | VR942 10 mg (n = 10) | VR942 20 mg (n = 10) |
|----------------|-------------------------------|------------------|----------------------|---------------------|---------------------|---------------------|---------------------|
| Age (years), mean (SD) | 34 (8.4) | 35 (11.0) | 32 (14.3) | 32 (5.4) | 29 (9.7) | 33 (7.3) | 29 (9.3) |
| Men | 100% | 100% | 100% | 100% | 100% | 100% | 100% |
| Race | White | 7 (70.0%) | 4 (66.7%) | 4 (66.7%) | 3 (50.0%) | 2 (33.3%) | 5 (83.3%) | 10 (62.5%) |
| Black/African American | 1 (10.0%) | 1 (16.7%) | 1 (16.7%) | 1 (16.7%) | 2 (33.3%) | 0 | 2 (12.5%) | 2 (12.5%) |
| Asian | 0 | 1 (16.7%) | 0 | 1 (16.7%) | 1 (16.7%) | 0 | 0 | 0 |
| Other | 2 (20.0%) | 0 | 1 (16.7%) | 1 (16.7%) | 0 | 1 (16.7%) | 1 (6.7%) | 0 |
| BMI (kg/m²), mean (SD) | 23.5 (2.51) | 27.2 (2.35) | 23.0 (3.04) | 25.6 (2.56) | 25.1 (1.97) | 26.0 (3.24) | 24.8 (3.06) |
| Smoking status | Former smoker | 4 (40.0%) | 1 (16.7%) | 1 (16.7%) | 1 (16.7%) | 2 (33.3%) | 3 (50.0%) | 11 (68.8%) |
| Never smoked | 6 (60.0%) | 5 (83.3%) | 5 (83.3%) | 5 (83.3%) | 4 (66.7%) | 5 (83.3%) | 5 (83.3%) | 14 (82.4%) |
| Spirometry, mean (SD) | | | | | | | | |
| FEV1 (% predicted) | 106 (8.7) | 100 (17.2) | 94 (9.5) | 107 (16.5) | 89 (4.8) | 102 (10.9) | 89 (17.1) | 86 (11.7) | 77 (13.5) |
| FEV1/FVC | 0.80 (0.045) | 0.81 (0.085) | 0.77 (0.062) | 0.82 (0.041) | 0.80 (0.048) | 0.79 (0.042) | 0.75 (0.081) | 0.68 (0.087) | 0.72 (0.054) |
| FEV1 (L) | 4.23 (0.435) | 4.10 (1.012) | 3.80 (0.580) | 4.27 (0.818) | 3.40 (0.318) | 4.27 (0.385) | 3.68 (0.895) | 3.71 (0.718) | 3.16 (0.723) |
| ICS-naïve | 100% | 100% | 100% | 100% | 100% | 100% | 100% |

### Table 2
AEs in healthy participants from predose day 1 to day 28.

| AE | Placebo (n = 10) | VR942 dose | 0.5 mg (n = 10) | 1 mg (n = 16) | 5 mg (n = 6) | 10 mg (n = 6) | 20 mg (n = 6) | Total (n = 30) |
|----|------------------|------------|-----------------|---------------|------------|-------------|-------------|-------------|
| Any AE | 0 | 1 (17%) | 3 (50%) | 2 (33%) | 3 (50%) | 2 (33%) | 11 (37%) |
| Any TEAE | 0 | 0 | 3 (50%) | 2 (33%) | 3 (50%) | 2 (33%) | 10 (33%) |
| Treatment-related TEAE | 0 | 0 | 1 (17%) | 0 | 1 (17%) | 0 | 2 (7%) |
| TEAEs by system order class Nervous system disorders | 0 | 0 | 1 (17%) | 1 (17%) | 1 (17%) | 1 (17%) | 4 (13%) |
| Headache | 0 | 0 | 1 (17%) | 1 (17%) | 1 (17%) | 1 (17%) | 4 (13%) |
| Dizziness | 0 | 0 | 0 | 0 | 0 | 1 (17%) | 1 (3%) |
| Gastrointestinal disorders | 0 | 0 | 1 (17%) | 1 (17%) | 1 (17%) | 1 (17%) | 3 (10%) |
| Asthmatic ulcer | 0 | 0 | 1 (17%) | 0 | 0 | 0 | 0 |
| Diarrhea | 0 | 0 | 0 | 1 (17%) | 0 | 0 | 1 (3%) |
| Toothache | 0 | 0 | 0 | 0 | 0 | 1 (17%) | 1 (3%) |
| Musculoskeletal and connective tissue disorders | 0 | 0 | 0 | 2 (33%) | 0 | 0 | 2 (7%) |
| Arthralgia | 0 | 0 | 0 | 1 (17%) | 0 | 0 | 1 (3%) |
| Back pain | 0 | 0 | 0 | 1 (17%) | 0 | 0 | 1 (3%) |
| Respiratory, thoracic, and mediastinal disorders | 0 | 0 | 0 | 0 | 2 (33%) | 0 | 2 (7%) |
| Cough | 0 | 0 | 0 | 0 | 1 (17%) | 0 | 1 (3%) |
| Rhinorrhea | 0 | 0 | 0 | 0 | 1 (17%) | 0 | 1 (3%) |
| General disorders and administration-site conditions | 0 | 0 | 0 | 1 (17%) | 0 | 0 | 1 (3%) |
| Pyrexia | 0 | 0 | 0 | 1 (17%) | 0 | 0 | 1 (3%) |
| Infections and infestations | 0 | 0 | 0 | 0 | 1 (17%) | 0 | 1 (3%) |
| Urinary tract infection | 0 | 0 | 0 | 0 | 0 | 1 (17%) | 1 (3%) |
| Injury, poisoning, and procedural complications | 0 | 0 | 0 | 0 | 1 (17%) | 0 | 1 (3%) |
| Joint dislocation | 0 | 0 | 0 | 0 | 0 | 1 (17%) | 1 (3%) |

AE: adverse event; TEAE: treatment-emergent adverse event.

Data are n (%). Medical Dictionary for Regulatory Activities preferred terms are used. Participants with at least one AE are counted only once per system organ class and preferred term.
in patients with greater functional impairment. Lung function will be monitored in future studies.

Results of studies in similar populations showing that FeNO levels are responsive to anti-IL-13 therapies [15,17,28] and those in which IL-13 stimulates lung epithelial NO production make FeNO an intuitive endpoint for an inhaled treatment. Our analyses showed a dose-response effect for reduction in FeNO level following VR942 treatment, whereby no effect was observed with VR942 0.5 mg compared with placebo in contrast to the rapid reductions in FeNO with VR942 10 mg and 20 mg, which were significantly greater than the effect of placebo and were sustained for up to 96 h post cessation of treatment administration. FeNO reductions have been reported with other anti-IL-13 and

| Table 3 |
|---------|
| AEs in participants with asthma, from predose day 1 to day 28. |

| AE | Placebo (n = 16) | VR942 dose |
|----|-----------------|------------|
|    | 0.5 mg (n = 6) | 10 mg (n = 6) | 20 mg (n = 17) | Total (n = 29) |
| Any AE | 9 (56%) | 3 (50%) | 2 (33%) | 11 (65%) | 16 (55%) |
| Any TEAE | 9 (56%) | 3 (50%) | 2 (33%) | 11 (65%) | 16 (55%) |
| Treatment-related TEAE | 5 (31%) | 2 (33%) | 2 (33%) | 8 (47%) | 12 (41%) |
| Device-related TEAE | 0 | 0 | 0 | 1 (6%)* | 1 (3%)* |

**TEAEs by system organ class**

- **Nervous system disorders**
  - 2 (13%), 1 (17%), 2 (33%), 5 (29%), 8 (28%) for Placebo, VR942 0.5 mg, VR942 10 mg, VR942 20 mg, and Total, respectively.

- **Respiratory, thoracic, and mediastinal disorders**
  - 4 (25%), 1 (17%), 0, 5 (29%), 6 (21%) for Placebo, VR942 0.5 mg, VR942 10 mg, VR942 20 mg, and Total, respectively.

- **General disorders and administration-site conditions**
  - 2 (13%), 2 (33%), 0, 1 (6%), 3 (10%) for Placebo, VR942 0.5 mg, VR942 10 mg, VR942 20 mg, and Total, respectively.

AE: adverse event; TEAE: treatment-emergent adverse event. Data are n (%). Medical Dictionary for Regulatory Activities preferred terms are used. Participants with at least one AE are counted only once per system organ class and preferred term.

* Considered to be a consequence of the inhalation maneuver performed by this participant.

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Fig. 3. Mean percentage change in FeNO level from baseline to predose in participants with asthma. Placebo n = 16, VR942 0.5 mg n = 6, VR942 10 mg n = 6, VR942 20 mg n = 17. Error bars represent 95% CI. Boxed data (day 10) represent the prespecified analysis timepoint. CI: confidence interval; FeNO: fractional exhaled nitric oxide; FU: follow-up.

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anti-IL-4Rx candidates in participants with asthma not using ICS [15,17,28] and in those with uncontrolled asthma [14,19,27,29], where attenuation was associated with improvements in several clinical outcomes, including lung function and annualized exacerbation rates [19,26]. FeNO reductions observed with placebo in this study may have been due to participants being domiciled for the 10-day treatment period, therefore having a lower exposure to lung NO stimuli (e.g., environmental allergens, pollutants) than at home. A similar effect was reported in patients with asthma undergoing alpine rehabilitation, in whom significant reductions in FeNO levels over the first 2 weeks were attributed to house dust mite allergen avoidance [30–32].

We consider this to be the only evaluation of an inhaled anti-IL-13 treatment effect on daily in-clinic FeNO levels. Study limitations include evaluating VR942 treatment in those with mild asthma, and the duration of the study (with regard to establishing the potential for immunogenicity).

In conclusion, this phase 1, placebo-controlled, dose-escalation study showed that inhaled VR942 (0.5–20 mg) was generally well tolerated in healthy participants and participants with asthma. We showed a convincing PD effect at 10 mg and 20 mg doses, demonstrating rapid and sustained FeNO reductions greater than those observed with placebo. This supports target engagement of VR942 with IL-13 pathways in the airways of patients with asthma, justifying further clinical evaluation in future trials.

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**Conflicts of interest**

GB, MJM, FM, and AS are employees of Vectura Ltd; MJ, PP, FS, PV, MZ, and RP are employees of UCB Pharma. LL is a senior partner of TranScrip LLP. GB, MJM, FM, and AS are shareholders in Vectura Ltd. MB is the owner of HMR, a contract research organization.

**Author contributions**

GB analyzed and interpreted the data, and critically reviewed the manuscript drafts. MB was the principal investigator, conducted the study, and reviewed the protocol and manuscript drafts. MJ provided statistical design, analyzed and interpreted the data, and critically reviewed the manuscript drafts. LL analyzed and interpreted the data, and critically reviewed the manuscript drafts. MJ designed the study, performed literature searching, interpreted the data, and critically reviewed the manuscript drafts. FS designed the study, and critically reviewed the manuscript drafts. PV analyzed and interpreted the data, and critically reviewed the manuscript drafts. AS designed the study, and critically reviewed the manuscript drafts. PP analyzed and interpreted the data, and critically reviewed the manuscript drafts. FM designed the study, analyzed and interpreted the data, and critically reviewed the manuscript drafts. MJM designed the study, performed literature searching, interpreted the data, and critically reviewed the manuscript drafts. MJ provided statistical design, analyzed and interpreted the data, and critically reviewed the manuscript drafts. MJM, MM, and RL are shareholders in Vectura Ltd. MB is the owner of HMR, a contract research organization.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ebiom.2018.07.035.

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