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Repeated nebulisation of non-viral CFTR gene therapy in patients with cystic fibrosis: a randomised, double-blind, placebo-controlled, phase 2b trial


Summary

Background Lung delivery of plasmid DNA encoding the CFTR gene complexed with a cationic liposome is a potential treatment option for patients with cystic fibrosis. We aimed to assess the efficacy of non-viral CFTR gene therapy in patients with cystic fibrosis.

Methods We did this randomised, double-blind, placebo-controlled, phase 2b trial in two cystic fibrosis centres with patients recruited from 18 sites in the UK. Patients (aged ≥12 years) with a forced expiratory volume in 1 s (FEV1) of 50–90% predicted and any combination of CFTR mutations, were randomly assigned, via a computer-based randomisation system, to receive 5 mL of either nebulised pGM169/GL67A gene–liposome complex or 0·9% saline (placebo) every 28 days (plus or minus 5 days) for 1 year. Randomisation was stratified by % predicted FEV1 (<70%, ≥70%), age (<18 vs ≥18 years), inclusion in the mechanistic substudy, and dosing site (London or Edinburgh). Participants and investigators were masked to treatment allocation. The primary endpoint was the relative change in % predicted FEV1. The primary analysis was per protocol. This trial is registered with ClinicalTrials.gov, number NCT01621867.

Findings Between June 12, 2012, and June 24, 2013, we randomly assigned 140 patients to receive placebo (n=62) or pGM169/GL67A (n=78), of whom 116 (83%) patients comprised the per-protocol population. We noted a significant, albeit modest, treatment effect in the pGM169/GL67A group versus placebo at 12 months’ follow-up (3·7%, 95% CI 0·1–7·3; p=0·046). This outcome was associated with a stabilisation of lung function in the pGM169/GL67A group compared with a decline in the placebo group. We recorded no significant difference in treatment-attributable adverse events between groups.

Interpretation Monthly application of the pGM169/GL67A gene therapy formulation was associated with a significant, albeit modest, benefit in FEV1, compared with placebo at 1 year, indicating a stabilisation of lung function in the treatment group. Further improvements in efficacy and consistency of response to the current formulation are needed before gene therapy is suitable for clinical care; however, our findings should also encourage the rapid introduction of more potent gene transfer vectors into early phase trials.

Funding Medical Research Council/National Institute for Health Research Efficacy and Mechanism Evaluation Programme.

Introduction Cystic fibrosis has been a target for gene therapy since the CFTR gene was cloned in 1989.1 Lung disease is the main cause of morbidity and mortality in individuals with cystic fibrosis, with a median age at death of 29 years (95% CI 27–31).2 Early expectations of a rapid breakthrough were based on supposed ease of access to the target respiratory epithelium via inhaled aerosols. These hopes were tempered by the subsequent realisation that the airways are well defended, in keeping with their predominant function as conducting passages, rather than absorptive surfaces.
Various vectors for delivery of the CFTR gene into respiratory epithelial cells have been assessed. Viral approaches, including adenoviruses, adeno-associated viruses, and retroviruses, have faltered because of inefficient transduction from the luminal surface and immune responses restricting the efficacy of repeated application.1 As such, research from the UK Cystic Fibrosis Gene Therapy Consortium has initially focused on non-viral vectors. Formulation and delivery of plasmid DNA–liposome complexes have been refined in a large series of preclinical studies,4,5 and safety,6,7 molecular efficacy, and practical doses have been assessed in several phase 1 and 2a studies in patients with cystic fibrosis.1,3 We did this study to assess the clinical efficacy of the non-viral CFTR gene–liposome complex pGM169/GL67A after repeated delivery to the airways.

Methods

Study design and participants

We did this randomised, double-blind, placebo-controlled, phase 2b trial in two cystic fibrosis centres with patients recruited from 18 sites in the UK. Eligible participants had diagnosed cystic fibrosis, were aged 12 years or older, had a forced expiratory volume in 1 s (FEV1) of 50–90% predicted, and had any combination of CFTR mutations.

The protocol was approved by the National Research Ethics Committee and the Local Research Committees at the two dosing sites and the 16 other referral centres. Each participant, or a parent, provided written informed consent, and children provided assent.

Randomisation and masking

We randomly assigned patients (1:1), via a computer-based randomisation system, to receive nebulised pGM169/GL67A or 0·9% saline (placebo). Randomisation was stratified by % predicted FEV1 (<70 vs ≥70%), age (<18 vs ≥18 years), inclusion in the mechanistic substudy, and dosage site (London or Edinburgh). Participants in the mechanistic substudy were randomly assigned (2:1) to receive nebulised pGM169/GL67A or placebo, and could participate as part of either a nasal or bronchoscopy group, or both. Participants and investigators were masked to assessing changes in lung function in patients with a broad range of CFTR mutations. Additionally, our study shows that monthly repeated application of non-viral gene therapy can be safely administered to the lungs over a 1 year period.

Procedures

Patients received 5 mL of either 0·9% saline or pGM169/GL67A complex nebulised through a Trudell AeroEclipse II device (Trudell Medical International, London, ON, Canada) at 28 day intervals (plus or minus 5 days) for 12 months. Each 5 mL dose of pGM169/GL67A contained 13·3 mg of plasmid DNA and 75 mg of the GL67A lipid mixture. Routine treatments were continued throughout the study, except for DNase, which was withheld for 24 h before and after dosing. In addition to the nebulised dose, patients in the nasal group of the mechanistic substudy received 2 mL of placebo or pGM169/GL67A divided between nasal cavities via a nasal spray device at the time of each lung dose. Patients in the bronchoscopy group followed the standard protocol, but also underwent a bronchoscopy under general anaesthesia before the first dose and 28 days (plus or minus 5 days) after the final dose.

Outcomes

The primary efficacy endpoint was the relative change in % predicted FEV1, calculated from the mean of two baseline values (at screening and before dosing on day of the first dose) to the mean of two values (2 and 4 weeks after last dose) at study completion. Secondary outcomes included additional measurements of lung function, CT scans, and Cystic Fibrosis Questionnaire-Revised (CFQ-R) scores.1 Exploratory endpoints included exercise testing, activity monitoring, and sputum inflammatory markers. Mechanistic endpoints were nasal or bronchial vector-specific DNA, mRNA, and electrophysiological assessment of CFTR function. We did extensive safety assessments.
patients would provide 90% power to detect a 6% difference between groups in the mean change from baseline at a two-sided 5% significance level. This power calculation was conservative because covariate adjustment can be expected to increase statistical power. We did analyses in the per-protocol population (primary analysis), predefined as participants who received at least nine doses of pGM169/GL67A or placebo, and in the intention-to-treat population, who received at least one dose of pGM169/GL67A or placebo.

We compared outcomes between groups with an ANCOVA model, with inclusion of the relevant baseline value, treatment allocation, and stratification factors (baseline predicted FEV₁, age, dosing site, inclusion in substudy). Results are reported as adjusted mean differences with corresponding 95% CIs. We assessed subgroup effects by including the relevant interaction term in the ANCOVA model. To allow results from different endpoints to be plotted on a common scale, the estimated treatment effects were standardised and presented as multiples of the underlying SD. No adjustment was made to the p values to allow for multiplicity because the secondary endpoints were supportive and the corresponding p values were interpreted conservatively. We assessed bronchial and nasal biomarkers with a Mann–Whitney U test. A two-sided p value less than 0.05 was considered statistically significant.

The trial was overseen by an independent Data Monitoring and Ethics Committee and a Trial Steering Committee. This trial is registered with ClinicalTrials.gov, number NCT01621867.

Role of the funding source
The funder of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report. The corresponding author had full access to all the data in the study and had final responsibility for the decision to submit for publication.

Results
Figure 1 shows the trial profile. Between June 12, 2012, and June 24, 2013, we randomly assigned 140 patients to receive placebo (n=62) or pGM169/GL67A (n=78), of whom 136 (97%) patients comprised the intention-to-treat population and 116 (83%) patients comprised the per-protocol population (figure 1). Reasons for discontinuation in the intention-to-treat population were similar between groups (appendix). Baseline characteristics were similar between the two groups (table 1). Unless indicated otherwise, all subsequent details relate to the per-protocol population.

114 (98%) patients had paired pre-treatment and post-treatment measurements of % predicted FEV₁. Of the two patients (both in the placebo group) who did not have paired measurements, one patient could not do the test because of a surgery-related pneumothorax and one withdrew because of time commitments and was unavailable for follow-up measurements. We recorded a significant ANCOVA-adjusted treatment effect in the pGM169/GL67A group versus placebo at 12 months’ follow-up (3.7%, 95% CI 0.1–7.3; p=0.046; figure 2). The relative changes within each of the individual groups were −4.0% (95% CI −6.6 to −1.4) in the placebo group and −0.4% (−2.8 to 2.1) in the pGM169/GL67A group (figure 2). Post-hoc analysis showed that 21 (18%) patients (n=6 in the placebo group and n=15 in the pGM169/GL67A group) had an improvement in % predicted FEV₁ of 5% or more of their individual baseline values. For comparison, the treatment effect in patients in the intention-to-treat
population who had spirometry measurements both before dosing and within the protocol-defined window after their final dose (n=56 in the placebo group and n=65 in the pGM169/GL67A group) was 3.6% (95% CI 0.2–7.0; p=0.039), with the 20 patients included in the intention-to-treat, but not per-protocol, analysis, receiving a mean of 3.7 doses (SD 1.9).

Figure 3 summarises changes in a range of secondary outcomes. The treatment effect was significant for FVC (p=0.031; appendix) and CT gas trapping (p=0.048), but not for other measures of lung function, imaging, and quality of life (figure 3). We assessed whether a responder subgroup could be identified; the appendix summarises the prespecified subgroups. We noted no significant differences in the primary outcome treatment effect with respect to sex, age, CFTR mutation (phe508del homozygous vs other), Pseudomonas colonisation, predominant smaller or larger airway disease on CT at presentation, concurrent drugs, or treatment-associated adverse events (appendix). Although some subgroups had larger treatment effects than others, these results were typically due to a greater decline in FEV1 in the placebo group, rather than to any difference of effect in the pGM169/GL67A group (appendix). Stratification by baseline % predicted FEV1 suggested a difference, albeit non-significant, in treatment effect between patients with more severe disease (FEV1 49.6–69.2% predicted), who had a treatment effect of 6.4% (95% CI 0.8–12.1; pinteraction=0.065; appendix), and those with less severe disease (69.6–89.9% predicted), who had a treatment effect of 0.2% (−4.6 to 4.9; pinteraction=0.065; appendix). In patients with more severe disease, post-trial and pre-trial changes in both the placebo group (−4.9%) and the pGM169/GL67A group contributed to the treatment effect. Secondary outcomes showed a similar trend favouring the more severe category (appendix).

Patients in both treatment groups received a median of three (IQR one to five) courses of oral or intravenous antibiotics during the trial. Specifically, we assessed co-administered antibiotics during the critical analysis...
period from day 11 to the end of the trial. Numbers of patients receiving any additional antibiotics were 26 (48%) in the placebo group and 30 (51%) in the pGM169/GL67A group (p²=0.774). Thus, the observed FEV₁ treatment effect was considered to be independent of concurrent antibiotic courses.

No clinically relevant pattern of changes could be distinguished in the exploratory outcomes of activity and exercise monitoring and serum and sputum inflammatory markers (appendix). In the bronchoscopy group of the substudy, vector-specific DNA increased in 12 (86%) of 14 patients in the pGM169/GL67A group and was below the limit of quantification in all (n=7) placebo samples (p=0.001; figure 4A); vector-specific mRNA was below the level of sensitivity in both groups (appendix). Changes in basal post-trial and pre-trial potential difference values did not differ significantly in either group (appendix). Figure 4B shows bronchial chloride responses using the mean of all interpretable tracings for each patient; a negative value indicates a change in the non-cystic fibrosis direction. Patients in the placebo group (n=7) had a median change (post-trial minus pre-trial) of 3.1 mV (range 9.3 to –1.2) and those in the pGM169/GL67A group (n=10) had a change of –1.3 mV (4.0 to –5.8; p=0.032; figure 4B). Five (50%) of ten patients in the pGM169/GL67A group had values that were more negative than the largest response in the placebo group (figure 4). In the same analysis with only TLCOc mid-expiratory flow between 25% and 75% of FVC. KCOc=diffusion capacity of the alveolar capillary membrane, corrected for haemoglobin concentrations. TCOc=transfer factor of the lung for carbon monoxide, corrected for haemoglobin concentrations. *Refers to scores from the Cystic Fibrosis Questionnaire-Revised.
In patients in the nasal group of the substudy, vector-specific DNA increased in all the 17 patients given pGM169/GL67A. Despite apparent pGM169 contamination in some samples, the change in pGM169 concentrations differed significantly between the groups (appendix); no vector-specific mRNA was quantifiable in either group. We noted no significant changes in the baseline, zero chloride, or isoprenaline responses (appendix). Four (29%) of 14 pGM169/GL67A patients had mean post-trial minus pre-trial treatment responses (ranging from –3.4 mV to –7.0 mV) that were more negative than the largest response in the placebo group (n=6; appendix). The appendix shows absolute nasal potential difference values.

All patients had adverse events, with no significant difference between groups for either total events or within the nine predefined adverse event categories (table 2). One patient in the placebo group and one patient in the pGM169/GL67A group discontinued study treatment because of adverse events (fatigue and increased respiratory symptoms and flu-like symptoms, respectively). We recorded six serious adverse events, all in the pGM169/GL67A group (appendix). Neither the Data Monitoring and Ethics Committee nor the Trial Steering Committee regarded any serious adverse event as related to study drug; however, one event was considered to be possibly related to a trial procedure (bronchoscopy). We noted no clinically relevant changes in haematology, biochemistry, conversion of anti-CFTR T cells, anti-DNA antibodies, histology, or lipid staining (appendix) and no patients died during the study.

### Discussion

We report the first trial of non-viral based gene therapy for cystic fibrosis, powered to detect clinically relevant pulmonary changes. After monthly dosing for 1 year, we recorded evidence of a beneficial effect of gene therapy versus placebo on FEV1. No effect of sex, age, or whether patients were homozygous for the most common F508del CFTR mutation could be detected. No clinically important adverse events attributable to treatment with pGM169/GL67A were reported.

Although these findings are encouraging, they should be put into perspective. We noted a stabilisation of FEV1 in the pGM169/GL67A group rather than an improvement. This stabilisation took place over a 1 year period and further work will be needed to see if this effect is maintained. The reduction in FEV1 in the placebo group was within the range reported in some other prospective trials10–12 and is consistent with a median survival of 29 years, but is greater than would be expected from registry data.2 Three factors are likely to have influenced this difference. First, the requirement for clinical stability at trial entry meant that patients might have been at their optimum respiratory health at this stage. Second, the enthusiasm of patients to enter the trial, accompanied by a focus on self-care, might have resulted in short-term improvements in lung function during the recruitment period. Both factors are likely to lead to a subsequent decline in lung function as patients regress to their mean values. Third, we included all available data, whether from stable patients or those with exacerbations, by contrast with registry data, which focuses on measurements obtained at annual review. Stabilisation of lung disease in itself is a worthwhile aim and we would caution against the bar being set too high for novel therapeutics in cystic fibrosis populations with an unselected range of mutations. The large response to ivacaftor in patients with class III mutations takes place in the context of correctly localised CFTR protein. By contrast, much smaller improvements in lung function were shown in the ivacaftor–lumacaftor trial for the most common mutation (phe508del) in which the CFTR protein is misfolded and mislocalised.13

<table>
<thead>
<tr>
<th>Change in bronchial DNA score</th>
<th>Placebo (n=7)</th>
<th>pGM169/GL67A (n=14)</th>
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<tbody>
<tr>
<td>LOQ</td>
<td>Limit of quantification</td>
<td>Positive but not quantifiable</td>
</tr>
</tbody>
</table>

**Figure 4:** Assessment of DNA from bronchial brushings in the placebo (n=7) and pGM169/GL67A (n=14) subgroups (A) and the response of the bronchial epithelium to perfusion with a zero chloride solution containing isoprenaline 10 μM (B, C). Horizontal bars show median values. Each circle in panel A represents an individual patient. Each symbol in panels B and C shows the change in response from trial start to finish for the relevant treatment in an individual patient. Of the 16 participants in the bronchoscopy subgroup, 15 individuals had post-dose bronchoscopies, of whom 14 individuals generated samples for DNA and mRNA molecular analysis. The plotted value in panel B is the mean of all interpretable recordings (range 1–3), and in panel C is the most negative value obtained from all interpretable recordings, at each timepoint for that patient. A more negative value is in the non-cystic fibrosis direction. LOQ=limit of quantification, PBNQ=positive but not quantifiable.

<table>
<thead>
<tr>
<th>Articles</th>
<th>Change in chloride response (mV)</th>
<th>Placebo group (n=54)</th>
<th>pGM169/GL67A group (n=62)</th>
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<tr>
<td></td>
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<table>
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<th>Table 2: Adverse events</th>
<th>Placebo group (n=54)</th>
<th>pGM169/GL67A group (n=62)</th>
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<tr>
<td>Lower airway respiratory symptoms</td>
<td>7 9</td>
<td>9 0</td>
</tr>
<tr>
<td>Gastrointestinal symptoms</td>
<td>2 1</td>
<td>1 8</td>
</tr>
<tr>
<td>Fever or flu-like symptoms</td>
<td>1 1</td>
<td>1 4</td>
</tr>
<tr>
<td>Headache</td>
<td>1 2</td>
<td>1 1</td>
</tr>
<tr>
<td>Upper airway symptoms</td>
<td>2 3</td>
<td>3 4</td>
</tr>
<tr>
<td>Elevated liver function tests</td>
<td>0 3</td>
<td>0 4</td>
</tr>
<tr>
<td>Haematuria</td>
<td>0 2</td>
<td>0 2</td>
</tr>
<tr>
<td>Isolated raised inflammatory markers</td>
<td>0 8</td>
<td>0 7</td>
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<tr>
<td>Other</td>
<td>3 2</td>
<td>3 3</td>
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<tr>
<td>Total</td>
<td>19 1</td>
<td>21 2</td>
</tr>
</tbody>
</table>

Data are mean number of times the respective symptom was experienced by each patient during the trial. Values were calculated by dividing the total number of the relevant adverse event by the total number of relevant patients in that group.
The response in our study was heterogeneous, with apparent responders and non-responders. The data suggest that an approximate doubling of treatment effect was achieved in patients with more severe disease stratified by baseline FEV\textsubscript{1}, supported by trends in other clinically relevant secondary measures. A larger trial with a stratified trial entry design, powered to assess subgroups, and that addresses the mechanisms of response heterogeneity, will be important to verify or refute these data. This differential response could relate to the dose deposited in the airways; in patients with lower baseline FEV\textsubscript{1}, the relatively more obstructed smaller airways result in a larger proportion of the 5 mL dose being deposited in the larger airways. In pre-trial studies we assessed airway deposition in patients with cystic fibrosis with varying FEV\textsubscript{1} severity with technetium-99m labelled human serum albumin of similar droplet size (3–4 μm, using a different nebuliser system) to the pGM169/GL67A formulation. Bronchial airway (generations 2–8) fractional deposition was 2.9% of delivered dose (standard error of the mean [SEM] 0.2; n=33) in patients with 70–90% predicted FEV\textsubscript{1}, and roughly twice as great (6.0%, SEM 1.0; n=23) in those with 50–70% predicted FEV\textsubscript{1}. An additional contributory factor to this enhanced efficacy might be the increased mitotic rate of more severely affected tissues,\textsuperscript{16} which decreases the proportion of time that the nuclear membrane is intact, the membrane acting as a barrier to plasmid DNA entry to the nucleus.

We cannot rule out that the changes recorded in the present study are the result of a non-specific response to the pGM169/GL67A formulation. The placebo was 0.9% saline rather than a scrambled or CFTR-deleted plasmid–liposome complex. We selected 0.9% saline partly on the basis of pragmatic financial considerations, but mainly for ethical considerations, not wishing to expose patients with cystic fibrosis to first-in-man repeated pulmonary dosing of an untested product that might direct the expression of an immunologically active peptide or novel non-coding RNA molecule with deleterious biological functions. Furthermore, we wanted to compare progression on therapy with the natural history of the disease. In terms of alternative explanations for the effects we noted, we know of no evidence that monthly nebulisation of 0.9% saline is deleterious to lung function, nor that liposome alone produces physiological effects we noted, which result in changes in patient-related care.\textsuperscript{23,24} We did not formally assess infective exacerbations in view of the fairly small patient numbers in our study, but use of antibiotic courses as a surrogate identified no obvious treatment advantage. The treatment effect is consistent with a clinically meaningful benefit from the perspective of the European Medicine Agency;\textsuperscript{25} however, further improvements in efficacy and consistency of response to the current formulation, or its combination with CFTR potentiators, are needed before gene therapy is suitable for clinical practice. Furthermore, our findings should encourage the rapid introduction of more potent gene transfer vectors into early phase trials, now that much of the groundwork has been established.
The data reported here provide the first proof of concept that repeated administration of non-viral CFTR gene therapy can safely change clinically relevant parameters, providing another step along the path of translational cystic fibrosis gene therapy.

**Contributors**

EWFVA, ACB, SC, JCD, DMG, DRG, APG, UG, SCH, TEH, JAI, GDM, and DJP conceived, designed, and analysed the overall study. DA and GDM designed and coordinated data collection and statistical analysis. DKA, KJB, DB, PC, GD, NSD, HIE, RFF, [JG, JSRG, DMH, KH, SLH, J], BFK, MM, EKP, ALQ, CJJS, SSb, NJS, NS, EJS, SNS, RPU, and MDW assessed patient outcomes and undertook and analysed individual in-vivo assays. EVB, MHID, and SSo coordinated and undertook the administration of the trial. RB, NJ, PL-E, GR, and KS oversaw receipt, preparation, and dispensing of study drug. JB, RC, MC, HED, AD, JD, SG-S, LH, MPL, AWM, MCM, DM, CM, CAM, HM, LJM, AGN, TO, JP-L, IAP, KMP, BJS, SGS-J, MT, MYW, and JMW designed, undertook, and analysed in-vitro assays. SHC, RKS, and PW-H coordinated the production of lipid 67A, DDSC, LAD, and GM designed, undertook, and analysed studies of study drug delivery.

**Declaration of interests**

ACR, AD, APG, AGN, AWME, BFK, BJS, CM, CJJS, DKA, DA, DB, DM, DHM, DMG, DDSC, DJP, DRG, DKG, EKP, EVB, FWA, GD, GM, GDM, GR, HED, HM, HIE, IAP, JAI, JB, JCD, JD, JG, JSRG, JJ, JP-L, JMW, KJB, KH, KMP, KS, LAD, LH, LJN, MC, MCM, MHD, MM, MT, MYW, MAM, MDW, MPL, NSD, NJS, NJ, NJS, NS, PC, PLE, PW-H, RB, RC, REFF, RPU, SC, SCH, SG-S, SLH, SiS, SSS, SGS-J, SNS, TEH, TO, and UG report grants from the National Institute for Health Research, the Cystic Fibrosis Trust, Just Gene Therapy, Medicor Foundation, and other support from Genzyme, a Sanofi company, related to the submitted work during the conduct of the study. ALQ, JCD, JMWM, NC, NJS, RKS, SG, and SHC report fees, grants, honoraria, and patents outside of the submitted work. ACR, AD, APG, EKF, FWA, JAI, JCD, LAD, SCH, and UG report patents related to the submitted work.

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**References**


