Original research

Emulated trial investigating effects of multiple treatments: estimating combined effects of mucoactive nebulisers in cystic fibrosis using registry data

Emily Granger, Gwyneth Davies, Ruth H Keogh

ABSTRACT

Introduction People with cystic fibrosis (CF) are often on multiple long-term treatments, including mucoactive nebulisers. In the UK, the most common mucoactive nebuliser is dornase alfa (DNase). A common therapeutic approach for people already on DNase is to add hypertonic saline (HS). The effects of DNase and HS used alone have been studied in randomised trials, but their effects in combination have not. This study investigates whether, for people already prescribed DNase, adding HS has additional benefit for lung function or use of intravenous antibiotics.

Methods Using UK CF Registry data from 2007 to 2018, we emulated a target trial. We included people aged 6 years and over who were prescribed DNase without HS for 2 years. We investigated the effects of combinations of DNase and HS over 5 years of follow-up. Inverse-probability-of-treatment weighting was used to control confounding. The period predated triple combination CF transmembrane conductance regulator modulators in routine care.

Results 4498 individuals were included. At baseline, average age and forced expiratory volume in 1 s (FEV1%) predicted were 21.1 years and 69.7 respectively. During first year of follow-up, 3799 individuals were prescribed DNase alone; 426 added HS; 57 switched to HS alone and 216 were prescribed neither. We found no evidence that adding HS improved FEV1% at 1–5 years, or use of intravenous antibiotics at 1–4 years, compared with DNase alone.

Conclusion For individuals with CF prescribed DNase, we found no evidence that adding HS had an effect on FEV1% or prescription of intravenous antibiotics. Our study illustrates the emulated target trial approach using CF Registry data.

BACKGROUND

Randomised controlled trials (RCTs) are the gold standard for evaluating the effects of treatments. However, there are many more questions relating to treatments than can reasonably be evaluated within RCTs. When there are multiple potential treatment strategies, for example, it can be challenging to recruit enough individuals in each treatment arm for sufficient power. An alternative is to use observational data to study treatment effects, and it is increasingly recognised that emulating a target trial when using observational data helps to clarify the research question and avoid common biases.

WHAT IS ALREADY KNOWN ON THIS TOPIC

People with cystic fibrosis (CF) are often prescribed multiple long-term treatments, including mucoactive nebulisers such as DNase and hypertonic saline. The effects of DNase and hypertonic saline used alone on health outcomes have been studied in randomised trials, but the combined effects of both treatments have not been studied.

WHAT THIS STUDY ADDS

We emulated a hypothetical target trial using UK CF Registry data to compare multiple treatment strategies involving DNase and hypertonic saline. The primary interest was in estimating the effect of adding hypertonic saline to DNase after 2 years, compared with continued use of DNase alone, on long-term clinical health outcomes. We found no evidence that adding hypertonic saline has an effect on forced expiratory volume in 1 s or prescription of intravenous antibiotics.

HOW THIS STUDY MIGHT AFFECT RESEARCH, PRACTICE OR POLICY

We provide an example of target trial emulation to answer a clinical question for which there is no evidence from randomised controlled trials. This approach may be used to address clinical questions that are unlikely to be answered in randomised trials in the field of CF and more widely.
Cystic fibrosis

consequences of CFTR dysfunction. The most commonly prescribed mucoactive nebuliser is dornase alfa (DNase), which is recommended as the first choice of mucoactive agent if there is clinical evidence of lung disease. DNase works by reducing viscosity in the lungs, which helps to clear lung secretions. In contrast, mucoactive nebulisers such as hypertonic saline (HS), help to clear lung secretions by rehydrating the airway surface liquid.

RCTs have investigated the effects of DNase or HS alone in CF. A recent Cochrane systematic review found evidence to show that DNase alone may improve lung function and decrease pulmonary exacerbations in people with CF. Another Cochrane systematic review found evidence that HS alone can reduce pulmonary exacerbations and improve lung function, although the evidence for lung function was deemed low quality. Although DNase and HS are often prescribed in combination in clinical practice, their effects in combination have not been studied. Answering questions about the effects of different combinations of these two treatments using an RCT would be challenging, particularly if we are interested in studying long-term effects. We use UK CF Registry data to emulate a target trial designed to compare multiple treatment strategies involving DNase and HS and assess their long-term effects on health outcomes in people with CF. The focus is on patients already established on DNase as defined by CF Registry documentation of current prescription, and the primary aim is to investigate the causal effect of adding and continuing HS versus continuing to use DNase only on two health outcomes: lung function (measured using forced expiratory volume in 1 s, FEV, %) and prescription of intravenous antibiotics. We compare outcomes measured at 1, 2, 3, 4 and 5 years of follow-up under each treatment strategy, where each treatment strategy is to be sustained up to the outcome measurement time. Previous studies have investigated the long-term effects of DNase using registry data and the findings suggest that DNase may improve the rate of decline in lung function and that it may be more beneficial for people with lower lung function. However, HS has not been previously studied using registry data either alone or in combination with DNase.

Our study is undertaken using data which predates the widespread introduction of triple combination CFTR modifier therapies into routine clinical care. The question we address is relevant to the CF population overall, although results from the premodulator period may not translate to a modulator-treated population. This will be able to be investigated using the same framework once more years of data are available. Furthermore, access to modulators is not universal globally, and in those countries with access, patients ineligible or unable to tolerate them represent an important minority in whom this question is particularly relevant.

There are several examples of using the target trial framework to compare treatment strategies across disease areas, including CF, however, they have tended to focus on treatment strategies involving a single treatment. In this study, we use this approach to compare treatment strategies that involve a combination of two treatments.

METHODS
Study design and data source

Our study was designed to emulate a hypothetical RCT (ie, the ‘target trial’). The target trial framework involves describing the protocol for a randomised trial we would like to conduct if it were feasible, and then emulating that trial using the available observational data. A key element of the emulation of the trial involves controlling for confounding of the treatment-outcome relationship. The emulation is performed using the target trial framework.

Table 1 Components of the target trial we aim to emulate in this study

<table>
<thead>
<tr>
<th>Protocol component</th>
<th>Target trial</th>
<th>Emulation of the target trial using UK CF registry data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eligibility criteria</td>
<td>Include: UK individuals with CF who have been taking DNase 2 years and are aged at least 6 years. Exclude: Individuals who have received an organ transplant, been treated with hypertonic saline within the last 2 years, or are taking mannitol, lumacaftor/ivacaftor or tezacaftor/ivacaftor.</td>
<td>Include: Individuals observed in the UK CF Registry who meet the criteria in the target trial between 2007 and 2017, and who had at least 1 year of follow-up after baseline. Exclude: As in the target trial. We also exclude individuals with missing data on time-invariant confounders or FEV, % at baseline.</td>
</tr>
<tr>
<td>Treatment strategies</td>
<td>1. Continue DNase only and do not start hypertonic saline (DN) 2. Continue DNase and add hypertonic saline immediately (DN&amp;HS) 3. Stop DNase and start hypertonic saline (HS) 4. Stop DNase and do not start hypertonic saline (Nil) The treatment strategy is sustained for the duration of follow-up.</td>
<td>As in the target trial.</td>
</tr>
<tr>
<td>Assignment procedures</td>
<td>Participants will be randomly assigned to a treatment strategy when they are recruited to the trial. Participants and doctors will be aware of the treatment strategy they have been assigned to.</td>
<td>In the emulated trial individuals are not randomly assigned to the treatment strategy. This is accounted for in the analysis.</td>
</tr>
<tr>
<td>Follow-up period</td>
<td>1–5 years from randomisation.</td>
<td>1–5 years post-baseline.</td>
</tr>
<tr>
<td>Outcome</td>
<td>Lung function (measured using FEV, %) and use of intravenous antibiotics (yes/no)</td>
<td>As in the target trial.</td>
</tr>
<tr>
<td>Causal contrasts of interest</td>
<td>Per-protocol</td>
<td>As in the target trial.</td>
</tr>
<tr>
<td>Analysis plan</td>
<td>Estimate the mean difference in outcome between treatment strategies at follow-up for FEV, % and corresponding OR for use of intravenous antibiotics. Estimated using regression models for the outcome, with an indicator for treatment group and baseline measure of the outcome as explanatory variables. Confounding by measured baseline and time-varying covariates is addressed using IPTW of MSM (see ‘Treatment effect estimands and statistical analysis’).</td>
<td></td>
</tr>
</tbody>
</table>

CF, cystic fibrosis; DNase, dornase alfa; FEV1%, forced expiratory volume in 1 s; IPTW, inverse-probability-of-treatment weighting; MSM, marginal structural model.

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Cystic fibrosis association, as treatments are not randomly assigned in the observational data.

The key components of the protocol for our target trial are outlined in table 1. We emulate the target trial using data from the UK Cystic Fibrosis Registry.18 This is a national database managed by the Cystic Fibrosis Trust. Data are collected on time-invariant variables, such as sex, ethnicity and genotype, and variables that change over time. Longitudinal data are collected at approximately annual visits on over 250 variables covering several domains, including hospital admissions, pulmonary function, chronic medications, health complications, and these data have been recorded in a centralised database since 2007. For this study, data were available from 2007 to 2018. Further details on the registry are provided elsewhere.18

Table 1 specifies the components of the target trial and summarises how the registry data are used to emulate the target trial. Individuals meeting the inclusion criteria are those aged 6 years or older, who have been prescribed DNase but not HS for two consecutive years between 1 January 2007 and 31 December 2017. The baseline year is defined as the first year the inclusion criteria were met, so the earliest possible baseline year was 2008 and the latest was 2017. It is possible for individuals to meet the

![Flowchart of participant selection into the study sample. CF, cystic fibrosis; DNase, dornase alfa; DN&HS, DNase and adding hypertonic saline.](http://thorax.bmj.com/)

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using the Global Lung Initiative equations.19 At each annual visit
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by spirometry, with %-predicted values for FEV1% calculated

treatment and receipt of intravenous antibiotics. Both outcomes are
used to impute any missing data on the treatments from the
criteria are listed in table
2013 and 2014, we would choose 2013. Exclusion
years of potential follow-
which was most recent, but which allowed for the maximum
we defined the baseline year to be the year from 2008 to 2017
inclusion criteria in more than 1
Our treatments of interest are DNase and HS. At each
an annual review visit it is recorded whether individuals have been
Continuous variables are summarised using mean (SD) and categorical variables are summarised using percentages (%).
*High-risk and low-risk genotype classifications previously defined in Franklin et al.21 Genotypes which do not fall within either category were labelled 'none assigned'.
†Intravenous hospital admissions: number of people with at least one intravenous hospital admission since the last review.
‡Infection data: indicator for any positive culture since the last review.
<table>
<thead>
<tr>
<th>Female, n (%)</th>
<th>DN (N=3799)</th>
<th>DN&amp;HS (N=426)</th>
<th>HS (N=57)</th>
<th>Nil (N=216)</th>
<th>Whole cohort (N=4498)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1733 (45.6)</td>
<td>223 (52.3)</td>
<td>31 (54.4)</td>
<td>106 (49.1)</td>
<td>2093 (46.5)</td>
<td></td>
</tr>
<tr>
<td>21.3 (11.6)</td>
<td>18.7 (10.3)</td>
<td>19.7 (9.6)</td>
<td>24.2 (11.0)</td>
<td>21.1 (11.5)</td>
<td></td>
</tr>
<tr>
<td>Genotype risk group, n (%)</td>
<td>High</td>
<td>2994 (78.8)</td>
<td>358 (84.0)</td>
<td>46 (80.7)</td>
<td>154 (71.3)</td>
</tr>
<tr>
<td>301 (7.9)</td>
<td>24 (5.6)</td>
<td>3 (5.3)</td>
<td>22 (10.2)</td>
<td>350 (7.8)</td>
<td></td>
</tr>
<tr>
<td>None assigned</td>
<td>504 (13.3)</td>
<td>44 (10.3)</td>
<td>8 (14.0)</td>
<td>40 (18.5)</td>
<td>596 (13.3)</td>
</tr>
<tr>
<td>White ethnicity, n (%)</td>
<td>3653 (96.2)</td>
<td>409 (96.0)</td>
<td>56 (98.2)</td>
<td>208 (96.3)</td>
<td>4326 (96.2)</td>
</tr>
<tr>
<td>No of intravenous days over the past year, n (%)</td>
<td>0</td>
<td>1665 (43.8)</td>
<td>151 (35.4)</td>
<td>24 (42.1)</td>
<td>100 (46.3)</td>
</tr>
<tr>
<td>1–14</td>
<td>718 (18.9)</td>
<td>75 (17.6)</td>
<td>12 (21.1)</td>
<td>36 (16.7)</td>
<td>841 (18.7)</td>
</tr>
<tr>
<td>15–28</td>
<td>490 (12.9)</td>
<td>66 (15.5)</td>
<td>12 (21.1)</td>
<td>24 (11.1)</td>
<td>592 (13.2)</td>
</tr>
<tr>
<td>29+</td>
<td>926 (24.4)</td>
<td>134 (31.5)</td>
<td>9 (15.8)</td>
<td>56 (25.9)</td>
<td>1125 (25.0)</td>
</tr>
<tr>
<td>Intravenous hospital admissions, n (%)</td>
<td>1638 (43.1)</td>
<td>220 (51.6)</td>
<td>28 (49.1)</td>
<td>91 (42.1)</td>
<td>1977 (44.0)</td>
</tr>
<tr>
<td>FEV1%, mean (SD)</td>
<td>70.1 (22.8)</td>
<td>67.9 (22.7)</td>
<td>66.2 (17.9)</td>
<td>66.6 (24.6)</td>
<td>69.7 (22.8)</td>
</tr>
<tr>
<td>Rate of decline in FEV1%, mean (SD)</td>
<td>−1.09 (1.50)</td>
<td>−1.33 (1.63)</td>
<td>−1.48 (1.28)</td>
<td>−1.23 (1.67)</td>
<td>−1.12 (1.52)</td>
</tr>
<tr>
<td>BMI z-score, mean (SD)</td>
<td>−0.05 (1.13)</td>
<td>−0.21 (1.13)</td>
<td>−0.31 (0.98)</td>
<td>−0.23 (1.26)</td>
<td>−0.08 (1.13)</td>
</tr>
<tr>
<td>Pseudomonas aeruginosa infection, n (%)</td>
<td>2286 (60.2)</td>
<td>258 (60.6)</td>
<td>39 (68.4)</td>
<td>144 (66.7)</td>
<td>2727 (60.6)</td>
</tr>
<tr>
<td>Staphylococcus infection, n (%)</td>
<td>1529 (40.2)</td>
<td>169 (39.7)</td>
<td>22 (38.6)</td>
<td>95 (44.0)</td>
<td>1815 (40.4)</td>
</tr>
<tr>
<td>Non-tuberculous, mycobacteria infection, n (%)</td>
<td>173 (4.6)</td>
<td>25 (5.9)</td>
<td>4 (7.0)</td>
<td>11 (5.1)</td>
<td>213 (4.7)</td>
</tr>
<tr>
<td>Pancreatic insufficiency, n (%)</td>
<td>3367 (98.9)</td>
<td>392 (92.0)</td>
<td>48 (84.2)</td>
<td>186 (86.1)</td>
<td>3993 (88.8)</td>
</tr>
<tr>
<td>Prescribed ivacaftor, n (%)</td>
<td>42 (1.1)</td>
<td>4 (0.9)</td>
<td>0 (0.0)</td>
<td>4 (1.9)</td>
<td>50 (1.1)</td>
</tr>
</tbody>
</table>

Inclusion criteria in more than 1 year. When this was the case, we defined the baseline year to be the year from 2008 to 2017 which was most recent, but which allowed for the maximum possible follow-up time up to 5 years. For example, a person meeting the inclusion criteria in 2012 and 2013 has 6 and 5 years of potential follow-up, respectively, so we would choose 2013 as their baseline year. For a person meeting the inclusion criteria in 2012, we would choose 2012. Inclusion criteria are listed in table 1.

Our treatments of interest are DNase and HS. At each annual review visit it is recorded whether individuals have been prescribed these treatments over the past year. Start and stop dates for long-term treatments are also available and these were used to impute any missing data on the treatments from the annual review visits. The two outcomes of interest are lung function and receipt of intravenous antibiotics. Both outcomes are recorded annually. Lung function is measured at annual visits by spirometry, with %-predicted values for FEV1% calculated using the Global Lung Initiative equations.19 At each annual visit the number of days on intravenous antibiotics (at home or in hospital) is recorded. In this study, use of intravenous antibiotics was treated as a binary variable indicating whether the individual has any recorded days on intravenous antibiotics since the last annual visit.

The emulation of the target trial uses existing observational data, reflecting data collected during routine clinical care and without any randomisation to treatment strategy. It is important that the analysis accounts for the lack of randomisation, as far as possible. In the observational data, the association between treatment and the outcome is suspected to be confounded by several factors. We used directed acyclic graphs to show the assumed relationships between the relevant variables in our data and to inform which variables should be considered as confounders (see online supplemental figures S.1 and S.2) in our analyses. The following variables were considered as potential confounders: sex, CF genotype, ethnicity, age, respiratory infections, intravenous hospital admissions, body mass index z-score, pancreatic insufficiency, use of CFTR modulators, past FEV1%, past intravenous antibiotic use and rate of decline in FEV1%. Except for sex, genotype and ethnicity, which we take to be fixed over time, these covariates are recorded annually. Further details on how these covariates were defined are provided in online supplemental file.

**Treatment effect estimands and statistical analysis**

The target trial specifies four longitudinal treatment strategies involving our two treatments of interest (table 1). Each treatment strategy involves beginning a particular combination of DNase and HS and sustaining that combination throughout follow-up. Our primary interest was in comparing the strategies of continuing DNase and adding HS (DN&HS) and continuing DNase only (DN). For the FEV1% outcome, the main estimands of interest were the mean differences in FEV1% at times 1–5 years.
had all individuals been following treatment strategy DN&HS, vs had all individuals been following treatment strategy DN. For intravenous antibiotics, the main estimands of interest were the corresponding ORs at times 1–4 years. Note that FEV1% is measured on the day of the annual review, whereas intravenous antibiotic use over the past year is recorded. To estimate 1-year, 2-year, 3-year, 4-year and 5-year treatment effects on FEV1%, we use FEV1% recorded at the first, second, third, fourth and fifth follow-up visit, respectively. To estimate the 1-year, 2-year, 3-year and 4-year treatment effects on intravenous antibiotic use, we use information recorded at the second, third, fourth and fifth follow-up visit. Comparisons between other treatment combinations were of secondary interest. We also compared the strategy of switching to and then continuing HS (HS) and the strategy of dropping DNase (Nil) with the strategy of continuing DNase only (DN).

The treatment effect estimands specified above were estimated using marginal structural models (MSM) estimated using inverse-probability-of-treatment weighting (IPTW). An MSM specifies how the outcome at a given time depends on treatment history up to that time, and in our case also on time and baseline covariates. The MSM cannot be fitted directly due to time-dependent confounding. IPTW involves estimating the probability of individuals receiving the treatment they received at each time point conditional on their treatment and covariate history up to that time. Multinomial regression was used to estimate the probability of having a given treatment combination (DN, DN&HS, HS, Nil). The IPTW at a given time is inverse of the product of the probabilities up to that time. Stabilised weights were used to avoid extreme weights.

We assumed that consecutive visits were approximately 1 year apart. Some individuals had less than five follow-up visits after their baseline visit, due to the administrative end of follow-up, death or organ transplant. These individuals were censored at the time of death, transplant or end of follow-up. We did not use data from visits at which an individual had missing data in the outcome, or from visits at which individuals were using certain treatments (mannitol, lumacaftor/ivacaftor or tezacaftor/ivacaftor). Inverse-probability-of-censoring weights were used to address censoring, and inverse-probability-of-observation weights were used to handle exclusions at a given visit due to missing outcome data or use of certain treatments. Each individual had a combined weight at each time point which combines the IPTW with these other weights. Further details on the weights are provided in online supplemental file (see ‘Inverse-probability-of-treatment weighted estimation of marginal structural models’). As well as missing outcome data, there were missing data in some of the confounding variables. Full details

![Figure 2](image-url) Estimated effects of adding hypertonic saline to DNase compared with continuing DNase alone on FEV1% and prescription of intravenous antibiotics. Mean differences are presented for FEV1% and ORs are presented for intravenous antibiotics. DNase, dornase alfa; FEV1%, forced expiratory volume in 1 s.

![Figure 3](image-url) Estimated effects of adding hypertonic saline to DNase compared with continuing DNase alone on FEV1%. Estimated effects are mean differences and are presented for people with high (100) moderate (75) or low (40) FEV1% at baseline. DNase, dornase alfa; FEV1%, forced expiratory volume in 1 s.
Cystic fibrosis

1. In other words, we found no evidence that adding HS would result in a different mean FEV1% or different odds of intravenous antibiotics among individuals who are already established on DNase, compared with continuing to use DNase only. For the effect estimate at year 4 in the low FEV1% group is 4.55. DNase, dornase alfa; FEV1%, forced expiratory volume in 1 s.

2. Details on the amount of missing data by follow-up year, and on the number of people who were censored each year (due to loss to follow-up, death or transplant), or temporarily excluded due to missing outcome data, are provided in online supplemental tables S.1 and S.2. Outcome trajectories by follow-up year and the distribution of weights used in the analysis are also provided in online supplemental figures S.5–S.7.

Estimates of the effects of treatment combinations

Figure 2 shows the expected mean differences in FEV1% (at times 1–5 years) and the odds of intravenous antibiotics versus no intravenous antibiotics (at times 1–4 years) between the two treatment strategies DN&HS vs DN (ie, the effect of adding HS on FEV1% and odds of intravenous antibiotics within an annual review year). Corresponding tabulated values are provided in online supplemental tables S.3 and S.4. For FEV1%, the mean differences are close to 0 at times 1–5, and all corresponding 95% CI contain 0. Similarly for intravenous antibiotics, the ORs are close to 1 at times 1–4, and all corresponding 95% CI contain 1. In other words, we found no evidence that adding HS would result in a different mean FEV1% or different odds of intravenous antibiotics among individuals who are already established on DNase, compared with continuing to use DNase only. For both outcomes, CIs, particularly for the later time points, were wide reflecting the uncertainty in our estimates.

Although not our focus, we also considered the effect of switching from DNase to HS (HS vs DN) and the effect of dropping DNase (Nil vs DNase). Figures for these additional comparisons are provided in the online supplemental figures S.8 and S.9.

Treatment effects by baseline FEV1%

We found no evidence of treatment effect heterogeneity by FEV1% at baseline. Figure 3 shows the expected mean differences in FEV1% at times 1–5 between DNXHs and DN, by each treatment combination by year, and online supplemental figure S.4 describes the flow of individuals between different treatment combinations by year. Of the 2521 individuals who were prescribed DNase alone in the first year and had 5 years of follow-up, 1615 (64.1%) remained on DNase alone for 5 years. Of the 260 individuals prescribed DNase and HS in the first year and who had 5 years of follow-up, 183 (71.2%) remained on DNase and HS for 5 years.

Figure 4 Estimated effects of adding hypertonic saline to DNase compared with continuing DNase alone on prescription of intravenous antibiotics. Estimated effects are ORs and are presented for people with high (100) moderate (75) or low (40) FEV1% at baseline. The upper limit of the 95% CI for the effect estimate at year 4 in the low FEV1% group is 4.55. DNase, dornase alfa; FEV1%, forced expiratory volume in 1 s.

on the amount of missing data and our approach to handling it are given in the online supplemental file (see ‘Missing data’). The MSM was fitted using the combined weights. For the FEV1% outcome the MSM is a linear regression model and for the intravenous antibiotic use outcome the MSM is a logistic regression model. The MSms were fitted for all follow-up times combined with follow-up visit included as a covariate. The analysis for each outcome was conducted with and without interaction terms between treatment use and FEV1% at baseline in the MSM. The 95% CIs for the interaction terms were estimated to assess the evidence for treatment effect heterogeneity and we present treatment effects in individuals with low, moderate or high FEV1% at baseline. Figure S.3 shows the expected mean differences in FEV1% at baseline by setting FEV1% to 40, 75 and 100, respectively. Full specification of the MSms is provided in online supplemental file (see ‘Inverse-probability-of-treatment weighted estimation of marginal structural models’). All SEs and 95% CIs were estimated using the non-parametric bootstrap approach.

RESULTS

Study population and descriptive statistics

We identified 5836 individuals in the UK CF Registry who had been documented as having been prescribed DNase and not prescribed HS for at least two consecutive years between 2007 and 2017. Of these, 4498 individuals met our other inclusion and exclusion criteria for the emulated trial. Figure 1 describes the study sample derivation.

Table 2 summarises the characteristics, measured at baseline, of the study population, by the treatment combination they were observed to be using in the first year of follow-up. During the first year, 3799 individuals were prescribed DNase alone and 426 were prescribed both DNase and HS. Far fewer people were prescribed HS alone, or neither treatment (57 and 216, respectively). On average, individuals prescribed both treatments were the youngest (mean age: 18.7 years) and individuals taking neither treatment were the oldest (mean age: 24.2 years). Individuals prescribed HS alone had the lowest lung function (mean FEV1%: 66.2), whereas those prescribed DNase alone had the highest (mean FEV1%: 70.1). The proportion of people who were recorded as taking no intravenous antibiotics in the year prior to baseline was highest for people prescribed neither treatment (46.3%) and lowest for individuals prescribed DNase and HS (35.4%). Individuals were observed to switch between treatment combination during the follow-up. Online supplemental figure S.3 shows the number of individuals prescribed treatment combinations by year. Of the 260 individuals prescribed DNase and HS in the first year and who had 5 years of follow-up, 183 (71.2%) remained on DNase and HS for 5 years.

Figure 1 Study population and descriptive statistics

Details on the amount of missing data by follow-up year, and on the number of people who were censored each year (due to loss to follow-up, death or transplant), or temporarily excluded due to missing outcome data, are provided in online supplemental tables S.3 and S.4. For FEV1%, the mean differences are close to 0 at times 1–5, and all corresponding 95% CI contain 0. Similarly for intravenous antibiotics, the ORs are close to 1 at times 1–4, and all corresponding 95% CI contain 1. In other words, we found no evidence that adding HS would result in a different mean FEV1% or different odds of intravenous antibiotics among individuals who are already established on DNase, compared with continuing to use DNase only. For both outcomes, CIs, particularly for the later time points, were wide reflecting the uncertainty in our estimates.

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FEV% at baseline (low, moderate, high). Figure 4 shows the corresponding ORs at times 1–4. For FEV%, the estimated mean differences increase as baseline FEV% decreases, suggesting that adding HS is more beneficial for individuals with lower FEV%. However, the corresponding 95% CI all contain 0. Similarly, the results for the intravenous antibiotics outcome provide no evidence of an effect of adding HS to DNase at any level of FEV%.

**DISCUSSION**

We used UK CF Registry data to emulate a hypothetical RCT designed to investigate the effects of multiple treatment strategies for mucoactive nebulised treatments on lung function and prescription of intravenous antibiotics in people with CF. Our primary interest was to investigate whether, for individuals already treated with DNase, adding HS has any additional benefit for these clinical health outcomes. We found no evidence of an effect (harmful or beneficial) of adding HS to DNase, either in change of FEV%,% or prescription of intravenous antibiotics. We found some suggestion that adding HS may benefit lung function for people with lower initial FEV%, although the results were not statistically significant. This is in line with results of a previous study based on UK CF Registry data, which suggested that the use of DNase alone could be more beneficial for FEV% for people with lower initial FEV%.13 People on both DNase and HS in the first year of follow-up tended to have lower lung function and more intravenous days, reflecting clinical practice where HS may be added to DNase when there has been clinical deterioration. This was addressed in the analysis by using IPTW to account for potential confounders.

Despite DNase and HS being commonly prescribed together in clinical practice, there have been no RCTs investigating the effects of these treatments used in combination. Hence, we demonstrate the application of target trial emulation to address a clinical question in CF for which there is no RCT evidence. The target trial framework applies the study design principles of RCTs to observational studies which helps to minimise biases that can arise due to study design and analysis choices.1–5 Target trial emulation has been used successfully in other disease areas to replicate the results from existing RCTs, for example, cardiovascular disease,22,23 and diabetes.24 The UK CF Registry is cited as an exemplar patient registry in the NICE real-world evidence framework,25 holds pharmacovigilance credentials and hosts post-authorisation phase IV pharmacovigilance studies.26 It is the largest national CF registry outside of the USA and captures data on almost all the UK CF population.18 These data, coupled with target trial methodology, provide an opportunity for researchers in CF to address important questions for the CF community. We did not perform sample size calculations and there is some debate as to whether such calculations are needed in observational studies such as this.27–29 We used all the available data in the UK CF Registry, giving the biggest possible sample size for the study.

Although our primary interest was to investigate the effect of adding HS when established on DNase, we also investigated the effect of dropping DNase after 2 years and switching to HS after 2 years. Results suggested poorer outcomes in terms of FEV%,% and intravenous antibiotics when dropping DNase, although results were largely non-significant. We found no evidence of an effect (in either direction) of switching to HS on intravenous days, although some evidence that switching to HS after 2 years can improve FEV%. This result is not clinically plausible and may be impacted by unmeasured confounding, which we discuss further below. Switching to HS is a rare decision in clinical practice. The number of people prescribed HS alone reflects this (table 2).

There are several limitations to this study. A key limitation in analyses that use observational data to study treatment effects is the possibility of bias due to uncontrolled confounding. Our analyses crucially assume that we have captured all the reasons for prescribing different treatment combinations that are associated with the outcome. While our analyses have controlled for several factors considered as potential confounders, including indicators of disease severity, it is important to note the possibility of residual confounding due to factors we did not control for. There could be, for example, biological or socioeconomic factors that influence both the treatment strategy and the outcome but are not collected by the registry. A further limitation is that our analyses rely on accurate treatment data being entered into the registry by clinical teams. Recording of long-term treatments within the CF Registry captures whether the treatment has been prescribed over the past year, but there is no information on adherence to treatment or dosing regimen. It is, therefore, possible that some individuals did not take or were poorly adherent to their prescribed medicine, which could bias our results. Additionally, we were interested in the health outcomes FEV%,% and pulmonary exacerbations (since previous trials have shown that DNase and HS can independently improve these outcomes), but we used intravenous antibiotic days as a proxy for exacerbations. The information on intravenous antibiotic days included both planned and unplanned intravenous and is not a direct marker of exacerbations. However, our approach is in line with previous studies using intravenous antibiotic days data from the UK CF Registry.31,32 Finally, the data used are from 2007 to 2018, and outcomes within the UK CF population are evolving rapidly with the introduction of CFTR modulator treatments, and to answer related questions about discontinuing treatment in those using CFTR modulators.

**CONCLUSIONS**

In an emulated trial using observational UK CF Registry data, we saw no additional benefit to lung function or use of intravenous antibiotics when HS was added to DNase.

Our findings show that the UK CF Registry can support methodology for emulated trials. Although RCTs remain the gold standard, this methodology has the potential to address questions relevant to the CF community and could be particularly useful for assessing the long-term clinical effectiveness of multiple treatment strategies, since such questions are difficult to answer using a clinical trial. In future work, UK CF Registry data combined with the target trial framework could be used to repeat our study in a post-modulator population, including in groups with and without access to, or intolerant of, CFTR modulator treatments, and to answer related questions about discontinuing treatment in those using CFTR modulators.

**Twitter** Emily Granger @EGranger90, Gwyneth Davies @daviesgwyneth and Ruth H Keogh @RuthHKeogh

**Acknowledgements** The authors thank people with CF and their families for consenting to their data being held in the UK CF Registry, and NHS teams in CF centres and clinics for the input of data into the Registry. We also thank the UK Cystic Fibrosis Trust and the Registry Steering Committee for access to anonymised UK CF Registry data.

**Contributors** Authorship CRediT statement: EG: methodology, formal analysis,
visualisation, writing—original draft. GD: conceptualisation, methodology, writing—review and editing, supervision. RHK: conceptualisation, methodology, writing—review and editing, supervision, funding acquisition. EG is responsible for the overall content as the guarantor.

Funding GD is supported by a UKRI Future Leaders fellowships (MR/T041285/1). RHK and EG are supported by a UKRI Future Leaders fellowship (MR/S017968/1) awarded to RHK. All research at UCL Great Ormond Street Institute of Child Health is made possible by the NIHR Great Ormond Street Hospital Biomedical Research Centre.

Disclaimer The views expressed are those of the author and not necessarily those of the NHS, the NIHR or the Department of Health.

Competing interests GD reports speaker honoraria from Chiesi Ltd and Vertex Pharmaceuticals. RHK reports a speaker honorarium from Vertex Pharmaceuticals.

Patient consent for publication Not applicable.

Ethics approval This study involves human participants and this work used anonymised data from the UK CF Registry, which has Research Ethics Approval (Reference: 07/Q0104/2). The use of the data for this study was approved by the Registry Research Committee (data request reference 375). This study was also approved by the London School of Hygiene & Tropical Medicine Research Ethics Committee (Reference: 21390). Participants gave informed consent to participate in the study before taking part.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data may be obtained from a third party and are not publicly available. To access the data, an application must be made to the UK CF Registry Research Committee. https://www.cysticfibrosis.org.uk/the-work-we-do/uk-cf-registry/apply-for-data-from-the-uk-cf-registry.

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ORCID iDs Emily Granger http://orcid.org/0000-0003-0134-1467
Gwyneth Davies http://orcid.org/0000-0001-7937-2728

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Granger E, et al. Thorax 2023;0:1–8. doi:10.1136/thorax-2023-220031

Thorax: first published as 10.1136/thorax-2023-220031 on 14 July 2023. Downloaded from http://thorax.bmj.com/ on September 18, 2023 by guest. Protected by copyright.
SUPPLEMENTARY MATERIAL

AN EMULATED TRIAL INVESTIGATING THE EFFECTS OF MULTIPLE TREATMENTS: ESTIMATING COMBINED EFFECTS OF MUCOACTIVE TREATMENTS IN CYSTIC FIBROSIS USING REGISTRY DATA

Emily Granger¹, Gwyneth Davies², Ruth H. Keogh¹

¹Department of Medical Statistics,
Faculty of Epidemiology and Population Health,
London School of Hygiene and Tropical Medicine,
Keppel Street, London, WC1E 7HT

²Population, Policy and Practice Research and Teaching Department,
UCL Great Ormond Street Institute of Child Health (UCL GOS ICH),
London WC1N 1EH, United Kingdom

This document contains additional material on the methods (section 1) and results (section 2).
1. Additional notes on methodology

1.1 Causal estimands

The treatments of interest are dornase alfa and hypertonic saline. In the UK CF Registry, treatment use at each annual review is recorded as a yes/no indicating whether treatment was prescribed over the past year. Time is measured in years since baseline. Let $D_{N,i,k}$ and $H_{S,i,k}$ denote whether dornase alfa and hypertonic saline, respectively, was recorded for the $i$th person at time $k$ ($k=1,2,3,4,5$). Let $A_{i,k}$ denote which treatment combination the $i$th person was on at time $k$ (i.e. between times $k-1$ and $k$). Then $A_{i,k}$ is defined as:

$$A_{i,k} = \begin{cases} 
0 & \text{if } D_{N,i,k} = 0 \text{ and } H_{S,i,k} = 0 \\
1 & \text{if } D_{N,i,k} = 0 \text{ and } H_{S,i,k} = 1 \\
2 & \text{if } D_{N,i,k} = 1 \text{ and } H_{S,i,k} = 0 \\
3 & \text{if } D_{N,i,k} = 1 \text{ and } H_{S,i,k} = 1
\end{cases}$$

Henceforth we suppress the subscript $i$. The outcome at time $k$ is denoted $Y_k$. The outcomes were $\text{FEV}_1\%$ (continuous, measured on the day of the annual review visit) and IV antibiotic use (binary, denoting whether or not any IV antibiotics were prescribed since the last annual review visit). Recall that at baseline (time 1) all individuals had been using DNase for 2 years, according to our inclusion criteria. Let $\bar{A}_k = \{A_1, ..., A_k\}$ denote the treatment history from time 1 to point $k$ and let $Y_{\bar{A}_k}^k$ denote the potential outcome that would be observed for an individual with a particular treatment history $\bar{A}_k$. Our primary aim was to compare the strategies of adding hypertonic saline to DNase (denoted DN&HS) up to follow-up time of interest and continuing DNase alone up to the follow-up time of $\text{FEV}_1\%$ interest (denoted DN). Using our above notation, for the continuous outcome of the main estimands of interest are defined as:

- 1 year: $E(Y_{\bar{A}_1}^1=3) - E(Y_{\bar{A}_1}^1=2)$
- 2 year: $E(Y_{\bar{A}_2}^2=3,3) - E(Y_{\bar{A}_2}^2=2,2)$
- 3 year: $E(Y_{\bar{A}_3}^3=3,3,3) - E(Y_{\bar{A}_3}^3=2,2,2)$
- 4 year: $E(Y_{\bar{A}_4}^4=3,3,3,3) - E(Y_{\bar{A}_4}^4=2,2,2,2)$
- 5 year: $E(Y_{\bar{A}_5}^5=3,3,3,3,3) - E(Y_{\bar{A}_5}^5=2,2,2,2,2)$

Comparisons between other treatment strategies were of secondary interest. We also compared the treatment strategies of switching from DNase to hypertonic saline and continuing HS alone to the follow-up time of interest (denoted HS) versus continuing DNase alone (denoted DN):

$$E(Y_{\bar{A}_k}^k=1) - E(Y_{\bar{A}_k}^k=2), k = 1, ..., 5$$

We also compared the treatment strategies of dropping DNase and not adding hypertonic saline (denoted Nil) versus continuing DNase alone (DN):

$$E(Y_{\bar{A}_k}^k=0) - E(Y_{\bar{A}_k}^k=2), k = 1, ..., 5$$

For the binary outcome of whether a person was prescribed any days of IV antibiotic treatment the estimands are odds ratios instead of mean differences. These are discussed in more detail in section 2.1.

1.2 Confounding variables

To obtain unbiased estimates of the treatment effects, we needed control for both time-invariant and time-varying confounders. Figures S.1 and S.2 are the directed acyclic graphs (DAGs) which show the assumed relationships between variables in our data for the analyses with $\text{FEV}_1\%$ and IV days as the outcome, respectively. Both DAGs are simplified versions of reality. We have not included long-
term arrows (e.g. from a variable recorded at time $k - 2$ to one recorded at time $k$), or relationships between the time-dependent variables measured at a given visit. FEV$\%_i$ and BMI are recorded at the annual review visit. The following covariates (included together in a box in the DAGs) at a given annual visit refer to whether infection, pancreatic insufficiency, IV hospitalisation or ivacaftor prescription were recorded since the previous annual visit: pancreatic insufficiency, *Pseudomonas aeruginosa* infection, *Staphylococcus aureus* infection, *Nontuberculous Mycobacteria* infection, IV hospitalisation and Ivacaftor use.

Figure S. 1: Directed Acyclic Graph depicting the assumed short-term confounding paths of the treatment-outcome association when FEV$\%_i$ was the outcome ($Y_i$ denotes FEV$\%_i$ at time $i$).

Figure S. 2: Directed Acyclic Graph depicting the assumed short-term confounding paths of the treatment-outcome association when IV days was the outcome ($Y_i$ denotes binary indicator for IV prescription recorded at time $i$).

The subscripts denote follow-up year and subscript 0 denotes baseline. PA: *Pseudomonas aeruginosa* SA: *Staphylococcus aureus*; NTM: *Nontuberculous Mycobacteria*; IV: Intravenous antibiotics; BMI: Body Mass Index; A: Treatment combination.

As can be seen from Figures S.1 and S.2, the variables included as time-invariant confounders were: age at baseline, genotype, sex, ethnicity, rate of decline in FEV$\%$, BMI z-score at baseline and FEV$\%_0$. The variables included as time-varying confounders were: pancreatic insufficiency, ivacaftor use, *P. aeruginosa* infection, *Staphylococcus aureus* infection, *Nontuberculous mycobacteria* infection, hospital admissions for intravenous antibiotics, days on intravenous antibiotics, past BMI z-score and past FEV$\%_0$.

Genotype was classed as either high risk, low risk or not assigned as previously defined. Ethnicity was classed as white or non-white due to small numbers in non-white ethnic groups in this population. Rate of decline in FEV$\%$ represented the change in FEV$\%$ observed prior to baseline. We defined the following linear mixed model with random slope and intercept:

$$FEV\%_{ij} = (\alpha_0 + \delta_{0i}) + (\alpha_1 + \delta_{1i})j + e_{ij}$$
Where \( j \in \{0, 1, 2, 3, 4\} \) is the number of years before baseline (\( j = 0 \) is the baseline year). The estimate of the slope parameter \((\alpha_j + \delta_i)\) for each individual is used as a time-invariant variable representing rate of change in FEV\(^1\%\).

Pancreatic insufficiency was a yes/no indicator where individuals were assigned “yes” if they were prescribed pancreatic enzyme supplements. IV hospital admissions was a yes/no indicator were yes indicated individuals had at least one hospital admission for IV antibiotics over the past year. IV days included home and hospital admissions and was categorised as: 0, 1-4, 15-28 and 29+. BMI z-scores were calculated using the WHO reference distribution\(^2\) and FEV\(_1\)% was calculated using the Global Lung Initiative equations\(^3\).

### 1.3 Inverse-probability-of-treatment weighted estimation of marginal structural models.

Let \( \mathbf{L}_B \) denote the set of time-invariant confounders and \( \mathbf{L}_k \) denote the set of time-varying confounders recorded at time \( k \). In both the FEV\(_1\)% and IV days analyses,

\[
\mathbf{L}_B = \{\text{Age}_0, \text{Genotype}, \text{Sex}, \text{Ethnicity}, \text{Rate of decline in FEV1%, FEV1%, BMI}_0\}
\]

For the FEV\(_1\)% analysis, \( \mathbf{L}_k \) is defined as:

\[
\mathbf{L}_k = \{\text{FEV1%}_k-1, \text{BMI}_k-1, \text{IV days}_k, \text{IV Hospital Admission}_k, \text{NTM}_k, \text{SA infection}_n, \text{PA infection}_n, \text{Ivacaftor use}_n, \text{Pancreatic insufficiency}_n\}
\]

For the IV days analysis, \( \mathbf{L}_k \) is defined as:

\[
\mathbf{L}_k = \{\text{FEV1%}_k-1, \text{BMI}_k-1, \text{IV days}_k, \text{NTM}_k, \text{SA infection}_n, \text{PA infection}_n, \text{Ivacaftor use}_n, \text{Pancreatic insufficiency}_n\}
\]

Then, the stabilised inverse-probability-of-treatment weights for individual \( i \) at time \( k \) (\( \text{IPT}.w_{ik} \)) were defined as:

\[
\text{IPT}.w_{ik} = \frac{\prod_{j=0}^{\tilde{N}} \Pr(A_j = a_{ij}| \tilde{A}_{j-1} = \tilde{a}_{ij-1}, \mathbf{L}_B = \tilde{I}_i) \prod_{j=0}^{\tilde{L}} \Pr(A_j = a_{ij}| \tilde{A}_{j-1} = \tilde{a}_{ij-1}, \mathbf{L}_B = \tilde{I}_i)}{\prod_{j=0}^{\tilde{L}} \Pr(A_j = a_{ij}| \tilde{A}_{j-1} = \tilde{a}_{ij-1}, \mathbf{L}_B = \tilde{I}_i)}
\]

Weights were also used to account for missing data in FEV\(_1\)% or BMI (\( \text{MISS}.w_{ik} \)), loss-to-follow-up (\( \text{LTFU}.w_{ik} \)), censoring due to death or transplant (\( \text{CENS}.w_{ik} \)) and time-varying eligibility due to use of lumacaftor/ivacaftor or tezacaftor/ivacaftor (\( \text{LUTE}.w_{ik} \)) or mannitol (\( \text{MANN}.w_{ik} \)).

The probabilities required for each set of weights were obtained using logistic regression. For \( \text{LUTE}.w_{ik} \), and \( \text{MANN}.w_{ik} \), the outcomes were indicators for use of the relevant treatments.

For each individual, we excluded time-points with missing data for FEV\(_1\)% or BMI. To account for the missing data, the remaining individuals were re-weighted by the inverse of their probability of remaining in the study at a given time. The weights, \( \text{MISS}.w_{ik} \), were defined using a similar equation as the one for \( \text{IPT}.w_{ik} \), but the outcome was an indicator for missingness in FEV\(_1\)% or BMI for the \( i \)th individual at time \( k \).

Individuals who were lost to follow-up, died or had an organ transplant were censored at the time of the event. For the loss to follow-up weights, the outcome at time \( k \) was an indicator for whether the individual was lost to follow-up at time \( k + 1 \). For the censoring weights due to death or organ transplant (whichever occurred first), the outcome at time \( k \) was an indicator for whether the individual died or had a transplant between times \( k \) and \( k + 1 \).

All weights were stabilised and probabilities were conditioned on the same variables as the probabilities defined in the inverse-probability-of-treatment weights.

The combined weight for individual \( i \) at time point \( k \) (\( \text{COMBINED}.w_{ik} \)) was defined as a product of all of the above weights:

\[
\text{COMBINED}.w_{ik} = \text{IPT}.w_{ik} \times \text{LUTE}.w_{ik} \times \text{MANN}.w_{ik} \times \text{MISS}.w_{ik} \times \text{CENS}.w_{ik} \times \text{LTFU}.w_{ik}
\]
For our main analysis, we specified the following linear marginal structural model (MSM) for the continuous outcome of FEV₁%:

\[ Y_{ik} = \beta_0 + \sum_{j=1}^{k} \sum_{c=1}^{3} \beta_{cj} I(a_j = c) + \beta_{k} L_{Bi} + \beta_{k}(k + 1) + \epsilon_{ik}, k = 1, \ldots, 5 \]

The parameters of the MSM are estimated by fitting the model using the observed data weighted using the combined weight. This enables estimation of the estimands specified in supplementary section 1.1. We note that these are marginal mean differences, as the conditional and marginal mean differences coincide for the linear MSM above.

For the binary outcome of whether the individual was prescribed any IV antibiotics over the past year, the marginal structural model (MSM) used for the main analysis was:

\[ \log \left( \frac{\Pr (Y_{ik+1} = 1 | L_{Bi})}{\Pr (Y_{ik+1} = 0 | L_{Bi})} \right) = \beta_0 + \sum_{j=1}^{k} \sum_{c=1}^{3} \beta_{cj} I(a_j = c) + \beta_{k} L_{Bi} + \beta_{k}(k + 1) + \epsilon_{ik}, k = 1, \ldots, 4 \]

This can be fitted using the observed data weighted using the combined weight. This results in estimates of conditional odds ratios. For example, our primary odds ratios of interest are

\[ OR_{k}^{DN&HS vs DN} = \frac{\Pr (Y_{ik+1} = 1 | L_B = 1)}{\Pr (Y_{ik+1} = 0 | L_B = 0)} / \frac{\Pr (Y_{ik+1} = 1 | L_B)}{\Pr (Y_{ik+1} = 0 | L_B)} \]

For the analyses investigating whether the treatment effects differed by FEV₁% measured at baseline, the above MSMs were extended to include an interaction between FEV₁% (a component of \( L_B \)) and \( I(a_j = c) \).

2. Additional results

2.1 Missing data

We found 5360 individuals with CF who were documented as having been prescribed dornase alfa and not hypertonic saline for at least two consecutive years between 2007 and 2017, and who had at least one baseline visit and one follow-up year. After excluding individuals who were under the age of 6 years, had received a solid organ transplant by baseline, or were prescribed mannitol, tezacaftor/ivacaftor or lumacaftor/ivacaftor at baseline, we were left with 4810 individuals who were eligible for inclusion. Table S.1 shows the amount of missing data by year for those individuals. Note that this includes people who were transplanted, or prescribed mannitol, tezacaftor/ivacaftor or lumacaftor/ivacaftor post-baseline.

<table>
<thead>
<tr>
<th>Year (k)</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
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<tr>
<td></td>
<td>(n=4810)</td>
<td>(n=4810)</td>
<td>(n=4471)</td>
<td>(n=4078)</td>
<td>(n=3660)</td>
<td>(n=3261)</td>
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<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
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<tr>
<td>FEV₁%</td>
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<td>175 (3.9%)</td>
<td>149 (3.7%)</td>
<td>164 (4.5%)</td>
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</tr>
<tr>
<td>Number of IV days</td>
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<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
</tr>
<tr>
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<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
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</tr>
<tr>
<td>Genotype</td>
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<td>44 (0.9%)</td>
<td>44 (0.9%)</td>
<td>44 (0.9%)</td>
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<td>0 (0%)</td>
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<td>0 (0%)</td>
</tr>
<tr>
<td>Rate of decline in FEV₁%</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>Ivacaftor use</td>
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<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>P. aeruginosa infection</td>
<td>0 (0%)</td>
<td>3 (0.1%)</td>
<td>10 (0.2%)</td>
<td>13 (0.3%)</td>
<td>6 (0.2%)</td>
<td>5 (0.2%)</td>
</tr>
<tr>
<td>Staphylococcus aureus infection</td>
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<td>3 (0.1%)</td>
<td>10 (0.2%)</td>
<td>13 (0.3%)</td>
<td>6 (0.2%)</td>
<td>5 (0.2%)</td>
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<tr>
<td>NTM</td>
<td>0 (0%)</td>
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<td>11 (0.2%)</td>
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<td>7 (0.2%)</td>
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</tr>
<tr>
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<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
<td>0 (0%)</td>
</tr>
<tr>
<td>Lagged BMI z-score*</td>
<td>-</td>
<td>79 (1.6%)</td>
<td>79 (1.6%)</td>
<td>85 (1.9%)</td>
<td>57 (1.4%)</td>
<td>59 (1.6%)</td>
</tr>
</tbody>
</table>
Pancreatic Insufficiency  0 (0%)  0 (0%)  0 (0%)  0 (0%)  0 (0%)  0 (0%)

*Lagged BMI z-score at visit i refers to the BMI z-score at visit k-1

We excluded individuals with missing data on time-invariant variables (genotype and ethnicity) and individuals with missing FEV\(_1\)% data at baseline (\(k = 0\)).

The last observation carried forward was used to impute the missing infection data (\(P.\ aeruginosa\) infection, \(Staphylococcus aureus\) infection, \(Bukholderia cepa\) infection and NTM). This was considered a valid approach as these infections are usually long-term and there was no missing data in these variables at time 0.

Missing data weights were used to account for missing BMI and FEV\(_1\)% (except for individuals who had missing FEV\(_1\)% at baseline, who were excluded).

### 2.2 Summary of exclusions due to loss to follow-up, death, transplant, ineligibility and missing data.

Table S.2 gives the numbers of individuals who were excluded or censored each year for different reasons. Individuals who were censored due to loss-to-follow-up, death or transplant, and these individuals account for the decreasing number observed by follow-up year. For example, in between visits 1 and 2, 250 individuals were lost-to-follow-up, 58 died and 29 received an organ transplant. By visit 2, 4498-(250+58+29)=4161 individuals remained in the study.

Individuals who were temporarily excluded due to missing data or temporary ineligibility (due to initiating treatment with CFTR modulators or mannitol) were allowed to re-enter the study, and these numbers account for the differences between the number of people observed in each follow up year and the number of people included in the final analysis (final N).

Column percentages are given with respect to the sample sizes in row 1.

<table>
<thead>
<tr>
<th>Follow-up year:</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number observed*</td>
<td>4498</td>
<td>4161</td>
<td>3776</td>
<td>3365</td>
<td>2975</td>
</tr>
<tr>
<td>LTFU**</td>
<td>0 (0%)</td>
<td>250 (5.6%)</td>
<td>303 (7.3%)</td>
<td>305 (8.1%)</td>
<td>286 (8.5%)</td>
</tr>
<tr>
<td>Death**</td>
<td>0 (0%)</td>
<td>58 (1.3%)</td>
<td>57 (1.4%)</td>
<td>79 (2.1%)</td>
<td>74 (2.2%)</td>
</tr>
<tr>
<td>Transplant**</td>
<td>0 (0%)</td>
<td>29 (0.6%)</td>
<td>25 (0.6%)</td>
<td>27 (0.7%)</td>
<td>30 (0.9%)</td>
</tr>
<tr>
<td>Mannitol</td>
<td>20 (0.4%)</td>
<td>42 (1.0%)</td>
<td>56 (1.5%)</td>
<td>88 (2.6%)</td>
<td>120 (4.0%)</td>
</tr>
<tr>
<td>Prescribed CFTR modulators</td>
<td>8 (0.2%)</td>
<td>15 (0.4%)</td>
<td>13 (0.3%)</td>
<td>14 (0.4%)</td>
<td>35 (1.2%)</td>
</tr>
<tr>
<td>Missing data</td>
<td>162 (3.6%)</td>
<td>257 (6.2%)</td>
<td>222 (5.9%)</td>
<td>197 (5.9%)</td>
<td>186 (6.3%)</td>
</tr>
<tr>
<td>Final N***</td>
<td>4308</td>
<td>3847</td>
<td>3485</td>
<td>3066</td>
<td>2634</td>
</tr>
</tbody>
</table>

LTFU: Lost to follow-up; CFTR modulators: these include lumacaftor/ivacaftor and tezacaftor/ivacaftor; Missing data: this is missing data in FEV\(_1\)% or BMI z-score.

*Number observed: this gives the number of individuals who remained in the study by visit k.

**Numbers for visit k denote individuals who were lost-to-follow-up, died or received a transplant between visits k and k+1.

***Final N: final number of people included in the analysis each year after censoring and temporary exclusions.
2.3 Summary of the numbers of people prescribed each treatment combination and flow of participants between treatment combinations by year

Figure S.3 shows the number of people prescribed each treatment combination by year. Across all follow-up years, the percent of individuals using neither DNase nor hypertonic saline ranged between 4.2% and 5.2%. The percentages prescribed DNase only and hypertonic saline only ranged from 51.7%-81.2% and 1.2%-2.8% respectively. The percentage prescribed both DNase and hypertonic saline ranged from 9.1%-29.6%.

Figure S.4 describes the flow of participants between treatment combinations by year. Of the 143 individuals who were using neither DNase nor hypertonic saline in the first year and had 5 years of follow-up, 31 (21.7%) continued to use neither treatment for 5 years. Of the 51 individuals using hypertonic saline only in the first year (i.e, who switched from DNase to hypertonic saline) and had 5 years of follow-up, 16 (31.4%) remained on hypertonic saline only for 5 years. Of the 2521 individuals who continued to be prescribed DNase only in the first year and had 5 years of follow-up, 1615 (64.1%) remained on DNase only for 5 years. Of the 260 individuals who added hypertonic saline to DNase in the first year and had 5 years of follow-up, 185 (71.2%) remained on this combination for 5 years.
Figure S.3: Flowchart showing the number of participants in the study and number of participants prescribed each treatment combination by year.

First year of follow-up

Number of individuals who enter the study: 446

- Number of people temporarily excluded: 106 (23.9%)
- Number of people on treatment combination **“N”**: 158 (9.5%)
- Number of people on treatment combination **“D”**: 183 (11.0%)
- Number of people on treatment combination **“N” & “D”**: 410 (6.6%)

Number of people censored: 337 (7.6%)

Second year of follow-up

Number of individuals remaining in the study: 419

- Number of people temporarily excluded: 243 (7.8%)
- Number of people on treatment combination **“N”**: 106 (4.9%)
- Number of people on treatment combination **“D”**: 209 (9.8%)
- Number of people on treatment combination **“N” & “D”**: 361 (8.8%)

Number of people censored: 326 (9.9%)

Third year of follow-up

Number of individuals remaining in the study: 377

- Number of people temporarily excluded: 296 (7.7%)
- Number of people on treatment combination **“N”**: 158 (4.5%)
- Number of people on treatment combination **“D”**: 236 (6.4%)
- Number of people on treatment combination **“N” & “D”**: 441 (11.9%)

Number of people censored: 231 (11.6%)

Fourth year of follow-up

Number of individuals remaining in the study: 325

- Number of people temporarily excluded: 296 (9.8%)
- Number of people on treatment combination **“N”**: 173 (5.2%)
- Number of people on treatment combination **“D”**: 82 (2.5%)
- Number of people on treatment combination **“N” & “D”**: 165 (5.0%)

Number of people censored: 200 (6.1%)

Fifth year of follow-up

Number of individuals remaining in the study: 295

- Number of people temporarily excluded: 341 (11.5%)
- Number of people on treatment combination **“N”**: 122 (4.4%)
- Number of people on treatment combination **“D”**: 82 (2.8%)
- Number of people on treatment combination **“N” & “D”**: 130 (4.1%)

Number of people censored: 181 (6.1%)

The percentages given in the first follow-up year are percentages of the total number of individuals who entered the study. The percentages given in follow-up years 2-5 are percentages of the number of individuals who remained in the study in follow-up years 2-5, respectively.
Figure S.4: Flow of participants prescribed each treatment combination by follow-up year
2.3 Outcome trajectories in the whole cohort

Figure S.5 shows the average FEV\(_1\)% and the proportion of individuals with at least one day on IV antibiotics in the whole cohort, by follow-up visit. The average FEV\(_1\)% decreases by year, whereas the proportion of individuals on IV antibiotics increases by year.

*Figure S. 5: Average FEV\(_1\)% and proportion of people on IV antibiotics in the whole cohort, by follow-up visit. Note that the vertical axes are truncated and the changes over time are small.*

2.4 Distribution of weights

Figures S.6 and S.7 show the distribution of inverse-probability-of-treatment (IPT) weights and combined weights by year, respectively (weights are defined in Section 1.3). Boxplots show that the median weights are approximately 1 for each year, as expected. The variance of weights tends to increase by year but there are no extreme values.

*Figure S. 6: Distribution of inverse-probability-of-treatment (IPT) weights by year. Horizontal line at y=1.*
2.5 Comparing different treatment strategies

Figure S.8 shows the expected mean differences in FEV₁% at times 1-5 years between the following treatment strategies: (1) DN&HS versus DN; (2) HS versus DN and (3) Nil versus DN. The three comparisons can be interpreted as follows: (1) the effect of adding hypertonic saline (2) the effect of switching from DNase to hypertonic saline and (3) the effect of stopping DNase. Figure S.9 shows the estimated odds ratios for IV antibiotic treatment at times 1-4 years in the ‘active’ treatment strategies DN & HS, HS, Nil versus the comparator treatment strategy DN. Odds ratios above 1 indicate a larger odds of having IV therapy in the active treatment strategy.

The results show evidence of a beneficial effect of switching to hypertonic saline in terms of FEV₁%, but no effect (beneficial or harmful) on IV antibiotic use. Estimated effects of stopping DNase are negative for FEV₁% and positive for IV antibiotic use (indicating worse outcomes in people who stop DNase in both cases). However, there is no evidence of an effect on either outcome, with all 95% confidence intervals containing the null value (0 for FEV₁% and 1 for IV antibiotic use).

Figure S.8: Estimated effects (and 95% CIs) for multiple treatment strategy comparisons, for 1-5 years, on FEV₁%
2.7 Tabulated results

Table S.3 presents the estimated mean differences in FEV1%, at times 1-5, between the two treatment strategies DN&HS versus DN, for the whole cohort, and conditional on having high, medium or low FEV1% at baseline. High medium and low are defined as 100, 75 and 45, respectively. Table S.4 presents the odds of IV antibiotics versus no IV antibiotics, at times 1-4, between the two treatment strategies DN&HS versus DN, for the whole cohort, and conditional on having high, medium or low FEV1% at baseline.

Table S.3: Estimated mean difference in FEV1% comparing HS&DN versus DN for 1-5 years for the whole cohort, and conditional on high (100), moderate (75) or low (45) FEV1% at baseline

<table>
<thead>
<tr>
<th>Year</th>
<th>Whole cohort</th>
<th>High FEV1%</th>
<th>Moderate FEV1%</th>
<th>Low FEV1%</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.01 (-0.70, 0.81)</td>
<td>0.51 (-0.93, 1.81)</td>
<td>0.13 (-0.63, 0.91)</td>
<td>-0.40 (-1.44, 0.71)</td>
</tr>
<tr>
<td>2</td>
<td>0.04 (-1.05, 1.09)</td>
<td>-1.78 (-3.59, 0.25)</td>
<td>-0.39 (-1.50, 0.76)</td>
<td>1.56 (0.13, 2.93)</td>
</tr>
<tr>
<td>3</td>
<td>0.89 (-1.77, 2.31)</td>
<td>0.39 (-3.24, 3.45)</td>
<td>0.85 (-1.13, 2.66)</td>
<td>1.48 (-0.76, 3.53)</td>
</tr>
<tr>
<td>4</td>
<td>-1.49 (-3.53, 0.57)</td>
<td>-3.09 (-5.93, -0.02)</td>
<td>-1.89 (-3.91, 0.21)</td>
<td>-0.21 (-3.36, 3.05)</td>
</tr>
<tr>
<td>5</td>
<td>1.85 (-2.84, 5.44)</td>
<td>0.07 (-5.75, 5.47)</td>
<td>1.36 (-3.18, 5.10)</td>
<td>3.41 (-3.19, 8.85)</td>
</tr>
</tbody>
</table>

Table S.4: Estimated odds ratios of HS&DN versus DN for 1-5 years on prescription of IV antibiotics for the whole cohort, and conditional on high (100), moderate (75) or low (45) FEV1% at baseline

<table>
<thead>
<tr>
<th>Year</th>
<th>Whole cohort</th>
<th>High FEV1%</th>
<th>Moderate FEV1%</th>
<th>Low FEV1%</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1.12 (0.99, 1.28)</td>
<td>1.12 (0.87, 1.42)</td>
<td>1.12 (0.98, 1.27)</td>
<td>1.11 (0.87, 1.48)</td>
</tr>
<tr>
<td>2</td>
<td>1.10 (0.92, 1.30)</td>
<td>1.31 (0.96, 1.76)</td>
<td>1.12 (0.95, 1.32)</td>
<td>0.89 (0.64, 1.29)</td>
</tr>
<tr>
<td>3</td>
<td>1.12 (0.89, 1.46)</td>
<td>1.26 (0.87, 1.85)</td>
<td>1.13 (0.91, 1.48)</td>
<td>0.98 (0.63, 1.63)</td>
</tr>
<tr>
<td>4</td>
<td>1.24 (0.88, 1.81)</td>
<td>0.90 (0.44, 1.69)</td>
<td>1.22 (0.84, 1.84)</td>
<td>1.84 (1.01, 4.55)</td>
</tr>
</tbody>
</table>

References

