Sputum microbiota in adults with CF associates with response to inhaled tobramycin

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INTRODUCTION

Cystic fibrosis (CF) is caused by mutations in the CF transmembrane conductance regulatory gene ultimately resulting in defective ion transport across epithelial cells. This leads to the buildup of mucus in the respiratory tract—serving as an ideal environment for chronic bacterial infections. Pseudomonas aeruginosa (PA), the archetypal CF pathogen, infects up to 60% of patients with CF and chronic infection is associated with worse outcomes.3 Accordingly, clinicians prescribe inhaled antipseudomonal antibiotics to manage these chronic infections.2 Indeed, inhaled antibiotics enable supratherapeutic drug levels at the site of infection and avoid the systemic toxicity associated with parenteral antibiotics.3–7

The first-line therapy used to treat chronic PA infections in CF is inhaled tobramycin powder/solution (TIP/S).3 Aggregate data collected from large clinical trials shows TIP/S improves lung function, quality of life, nutritional status, reduces pulmonary exacerbations (PEx) and even reduces mortality.3–8–12 Despite the many clinical benefits afforded by these therapies, they do not result in sustained clearance of chronic infection but rather result in a variable decrease in P. aeruginosa bioburden (1–2 log10 decrease Colony Forming Unit/g sputum).13 Moreover, patient outcomes have not reliably correlated with reductions in PA sputum density.14

The last two decades of research has revealed that the airways of individuals with CF are colonised by a diverse collection of organisms beyond the few target antibiotics but rather result in a variable decrease in P. aeruginosa bioburden (1–2 log10 decrease Colony Forming Unit/g sputum).13 Moreover, patient outcomes have not reliably correlated with reductions in PA sputum density.14

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antibiotic exposures. However, little is known about how the CF sputum microbiome may influence response to maintenance therapy. In the previous studies, our group examined the effects of the inhaled antibiotic aztreonam lysine (AZLI) on a cohort of treatment naïve patients, and during a single 28-day cycle of on/off AZLI. In both studies, we found that those who failed to respond to AZLI had microbiomes with a higher relative abundance (RA) of Staphylococcus spp—a Gram-positive known to resist aztreonam. However, both studies were limited by their inclusion of a patient population with generally advanced lung disease given AZLI’s role as a salvage agent. Accordingly, we developed a study focusing on the sputum microbiome of patients with CF and their response to the first-line antipseudomonal, tobramycin. We hypothesised that TIP/S has additional ‘off-target’ effects on the CF sputum microbiome and that the baseline microbiome prior to therapy initiation is associated with subsequent patient response.

METHODS

Patients and sample collection

The Calgary Adult Cystic Fibrosis Clinic has kept and maintained a frozen repository of sputum collected from CF adults since 1997. This biobank includes >18,000 sputum samples collected from >300 individuals. Inclusion criteria for our study were chronic P. aeruginosa infection as defined by the Leeds criteria, receipt of TIP/S for at least 1 year, and ≥1 sputum sample collected within 18 months before and after the initiation of TIP/S. We excluded transplant recipients, and patients who had recent prior TIP/S (defined within 1 year of its index initiation). Similarly, we excluded samples collected from patients if they were collected within 30 days of a PEx and or/receipt of intravenous antibacterials.

Patient demographics including age, sex, CF and non-CF demographics, lung function (as measured by percent predicted forced expiratory volume in 1 s (ppFEV1) and percent predicted forced vital capacity (ppFVC)), nutritional status (as measured by body mass index), cultured classical-CF pathogens and medical therapies were collected for baseline samples (defined as first sample collected prior to treatment initiation).

Definitions

Definitions of lung disease stage were classified based on baseline ppFEV1; mild ≥70%, moderate ≥40–<70%, severe <40%. The primary outcome of interest was patient response. Our definition of response, selected prior to microbiome analysis, was achieving stable lung function measured by ppFEV1 in the year following TIP/S initiation (no net decline, ie, zero change). This was chosen based on clinical trial data associated with TIP/S where improvements in lung function of 0%–8% were observed following treatment—and which generated two comparable groups. Median ppFEV1 per person was determined for each patient for the 18 months prior to and following TIP/S initiation. Change in ppFEV1 was calculated from pre vs post calculations and patients were classified as responders if they had no change or improvements in lung function. Additionally, a supplemental analysis was conducted to define response using changes in the slope of a linear regression in ppFEV1, pre-TIP versus post-TIP/S per patient. Exploratory analyses were performed on samples collected pre vs post TIP/S initiation. Samples associated with TIS and TIP were analysed in aggregate (as TIP was designed specifically to recapitulate the pharmacokinetics of TIS) and separately. Patients were analysed in aggregate and separately by exposure to TIP/S as treatment naïve, or not exposed to TIP/S for at least 5 years.

Sputum DNA extraction, amplification and analysis

Bacterial DNA was extracted from 300 µL of homogenised sputum as previously described. In brief, the V3 hyper-variable region of the 16S rRNA gene was amplified using reverse and forward barcoded primers on the Illumina MiSeq platform. qPCR was performed to determine the absolute abundance of S. aureus and the total bacterial load (defined as the number of 16S rRNA copies/mL) in the sputum samples. Statistical analysis of operational taxonomic unit tables was carried out using phyloseq, vegan and DESeq2 packages in R (V.3.3.1). The Wilcoxon rank-sum test was used to describe changes in Shannon alpha-Diversity Indices (SDIs) in R. Community structures were assessed using Bray-Curtis (BC) beta-diversity measures after proportionally normalising all samples by dividing read counts by the total number of reads. Permuted multivariate analyses of variance (PERMANOVAs) were used to analyse statistical differences in beta-diversity. Results were visualised using principal coordinate analysis plots. To account for repetitive sampling of patients (pre or post) we conducted a sensitivity analysis by setting constraints to permutations by patient ID. Restricted analyses were performed when assessing pre vs post samples to include the first presample (baseline—with the least prior exposure to TIP/S) and the last post sample (in order to observe the effects of prolonged TIP/S exposure). DESeq2 (test=Wald fitType=parametric and Benjamin Hochberg multiple test correction) was used to determine genera that are differentially abundant, log of normalised reads were used to visualise data. Samples with an abundance value of 0 were changed to a value of 1 since log(0) is undefined and log(1) = 0.

RESULTS

Cohort characteristics

Forty-one patients contributing 151 sputum samples (median 4/patient, IQR 3–4) representing before (n=77) and after (n=74) the initiation of TIP/S (figure 1) were analysed. Pre-TIP/S samples were collected at median day −91 (IQR −184 to −1) relative to initiation and samples collected after TIS/TIP at median day +146 (IQR 87–218). At baseline, 17 patients were naïve to TIP/S having never been exposed, and 24 patients had remote exposure (a median 7 years free of TIP/S; IQR 2–8). Of the 41 patients, 26 initiated treatment with TIS and 15 TIP. TIS and TIP usage did not differ by response status or patient demographics including age or genetics. Patient demographics, chronic therapies and cultured pathogens at treatment initiation are presented in table 1 and online supplemental table S1. Patients were a median 30.4 years of age (IQR 24.2–35.2) and had a ppFEV1 57% (IQR 44–74) and ppFVC 87% (IQR 72–93).

Sputum microbiome composition by sex, response and patient ID

A total of 6,139,490 sequences (median 32,667 per sample (IQR 15,351–63,128)) were generated from 151 sputum samples. Intrapatient and interpatient variability is observed in taxonomic summary plots for taxa present in >20% of all samples (figure 1). When assessing all samples, we found patient ID explained 55% of the variation in beta-diversity (between sample diversity) as measured by BC dissimilarity (PERMANOVA, R²=0.55, p=0.001). In addition, Pseudomonas...
Cystic fibrosis

The predominate pathogen abundance explains 36% of the variance (PERMANOVA, $R^2=0.36$, $p=0.001$).

Baseline sputum microbiome and its association with clinical response to TIP/S

The primary goal for our study was to determine if baseline sputum microbiome features correlated with clinical response to TIP/S, which might suggest that microbiome may be used as a biomarker to predict response. Nineteen (46%) patients met our a priori definition of response (R), exhibiting no decline in lung function in the year following initiation. The median ppFEV1 change in responders was +2.5% (IQR 0.5%–6.5%). Twenty-two patients (54%) were defined as non-responders (NR) experiencing a median ppFEV1 drop of −5.5% (IQR −3 to −9.5). When assessing baseline samples (collected closest to day 0) we found that treatment response explained 6% of the variation in beta-diversity (between sample diversity), as measured by BC dissimilarity between responders and non-responders (PERMANOVA, $R^2=0.061$, $p=0.046^*$) (figure 2A). However, no differences were observed in SDI or other alpha-diversity (within sample bacterial richness and evenness) metrics (figure 2B). Similar findings were observed for both BC and SDI metrics when we assessed baseline samples collected from patients exposed to TIS and TIP separately (data not shown).

Next, we sought to identify genus-level differences based on response. We observed non-responders to have −5.28 log2 lower RA of Staphylococcus spp compared with responders at baseline ($p<0.001$) (figure 2C and online supplemental table S2). No other genera were significantly associated with treatment response although Pseudomonas spp trended towards a higher abundance in non-responders. As a sensitivity analysis, we assessed increasingly stringent definitions of response (online supplemental table S2). Definitions of response in our sensitivity analysis ranged from −2.5% to 3% changes in lung function post

Figure 1  Microbiome as a function of days from the initiation of TIP/S. Taxonomic summaries for taxa present in >20% of all samples (n=151) for a cohort of 41 patients with chronic Pseudomonas aeruginosa infection. Days from the initiation of TIP/S are represented on the X-axis. Individual patient subplots are organised by response (R/NR), sex (F/M) and patient ID. NR, not responder; TIP/S, tobramycin inhalational powder/solution.
Cystic fibrosis treatment initiation. We chose negative values as a definition of response based on the median (IQR) decrease in lung function of non-responders (patients who fell below the 25th percentile were considered as responders) and to generate a comparable number of patients in each group. A higher abundance of *Staphylococcus* spp continued to associate with response status in the majority of these alternate definitions for baseline pretreatment sputum. Furthermore, a linear regression comparing *Staphylococcus* RA as a function of change in lung function (following the initiation of TIP/S) trended towards a statistical correlation with a $R^2$ of 0.09, $p=0.05$). To confirm the observed difference of *Staphylococcus* spp RA between responders and non-responders, we

### Table 1  Cohort demographics and culture data for baseline samples collected prior to the initiation of chronic TIP/S

<table>
<thead>
<tr>
<th>Characteristics of patient cohort</th>
<th>Response*</th>
<th>Non-R†</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Responder (n=19 (46%))</td>
<td>Non-R (n=22 (54%))</td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>12 (63)</td>
<td>11 (50)</td>
<td>0.4</td>
</tr>
<tr>
<td>Median age, years (IQR)</td>
<td>29.2 (24.2–38.6)</td>
<td>30.4 (23.8–34.5)</td>
<td>0.6</td>
</tr>
<tr>
<td>Median BMI (Kg/m²)</td>
<td>22.1 (20.4–23.6)</td>
<td>21.1 (19.3–22.4)</td>
<td>0.3</td>
</tr>
<tr>
<td>Lung stage‡</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Forced expiratory volume (FEV) predicted</td>
<td>50 (41–69)</td>
<td>64 (48–83)</td>
<td>0.2</td>
</tr>
<tr>
<td>Mild</td>
<td>5 (26)</td>
<td>10 (45)</td>
<td>0.4</td>
</tr>
<tr>
<td>Moderate</td>
<td>9 (47)</td>
<td>8 (36)</td>
<td></td>
</tr>
<tr>
<td>Severe</td>
<td>5 (26)</td>
<td>4 (18)</td>
<td></td>
</tr>
<tr>
<td>Genotype§</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ΔF508 homozygous</td>
<td>9 (47)</td>
<td>12 (55)</td>
<td>0.5</td>
</tr>
<tr>
<td>ΔF508 heterozygous</td>
<td>7 (37)</td>
<td>6 (27)</td>
<td>0.4</td>
</tr>
<tr>
<td>CF comorbidities</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pancreatic insufficiency</td>
<td>15 (79)</td>
<td>21 (95)</td>
<td>0.1</td>
</tr>
<tr>
<td>CF-related diabetes</td>
<td>5 (26)</td>
<td>7 (32)</td>
<td>0.7</td>
</tr>
<tr>
<td>Liver disease</td>
<td>6 (32)</td>
<td>3 (14)</td>
<td>0.2</td>
</tr>
<tr>
<td>Sinus disease</td>
<td>5 (26)</td>
<td>6 (27)</td>
<td>0.9</td>
</tr>
<tr>
<td>Traditional cultured CF pathogens</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>Pseudomonas aeruginosa</em></td>
<td>19 (100)</td>
<td>22 (100)</td>
<td>&gt;0.99</td>
</tr>
<tr>
<td>Methicillin susceptible <em>Staphylococcus aureus</em></td>
<td>6 (32)</td>
<td>2 (9)</td>
<td>0.07</td>
</tr>
<tr>
<td>Methicillin resistant <em>S. aureus</em></td>
<td>0</td>
<td>0</td>
<td>NA</td>
</tr>
<tr>
<td>Other¶</td>
<td>2 (11)</td>
<td>3 (14)</td>
<td>0.8</td>
</tr>
</tbody>
</table>

X² tests were used to determine differences between dichotomous clinical variables. Wilcoxon rank-sum tests were conducted for analysis of continuous variables.

*Response was a priori defined as achieving stable lung function as measured by FEV1 per cent predicted in the year following TIP/S initiation (no net decline).†Non-R=non-responder.‡Definitions of lung stage based on forced expiratory volume in one second; mild ≥70%, moderate ≥40 to <70%, severe <40%.§Genotypic data were unknown for one patient in the non-responder group.¶Other bacterial species include *Escherichia coli, Enterobacter cloacae, Serratia marcescens, Streptococcus Group C, Stenotrophomonas maltophilia.*

BMI, body mass index; CF, cystic fibrosis; NA, not applicable; TIP/S, tobramycin inhalational powder/solution.

treatment initiation. We chose negative values as a definition of response based on the median (IQR) decrease in lung function of non-responders (patients who fell below the 25th percentile were considered as responders) and to generate a comparable number of patients in each group. A higher abundance of *Staphylococcus* spp continued to associate with response status in the majority of these alternate definitions for baseline pretreatment sputum. Furthermore, a linear regression comparing *Staphylococcus* RA as a function of change in lung function (following the initiation of TIP/S) trended towards a statistical correlation with a $R^2$ of 0.09, $p=0.05$). To confirm the observed difference of *Staphylococcus* spp RA between responders and non-responders, we

![Figure 2](Heirali A, et al. Thorax 2020;75:1058–1064. doi:10.1136/thoraxjnl-2019-214191)

**Figure 2** Baseline microbiome and associated response status for samples collected prior to TIP/S initiation. Responders were defined as patients who experienced no net decrease in lung function with TIP/S use. (A–C) Microbiome analyses of baseline samples only. (A) PcoA of Bray-Curtis dissimilarities showing community-wide differences between samples collected from R (n=19) vs non-R (n=22) (PERMANOVA, $R^2=0.061$, $p=0.046$*). (B) Shannon alpha-Diversity Index (SDI) between R and non-R for baseline samples were not significantly different as demonstrated by Wilcoxon RANK sum test. (C) Log 2-fold of normalised reads reveal responders have a higher abundance of *Staphylococcus* as measured by a Wald’s test using DESeq2. NR, not responder; PERMANOVA, permutational multivariate analysis of variance; TIP/S, tobramycin inhalational powder/solution.
Our next objective was to determine if changes in long-term microbial community structure associated with the initiation of TIP/S. No significant differences in sputum microbiota (BC) were identified when comparing all samples collected prior (n=77) to and post (n=74) initiation of TIP/S (PERMANOVA, $R^2=0.012$, $p=0.84$). Similar findings were observed when analyses were performed to account for multiple samples/patient and constrained by patient ID (PERMANOVA, $R^2=0.0033$, $p=0.85$) (online supplemental figure S2). No differences were observed in SDI for samples collected preinitiation versus postinitiation of TIP/S (online supplemental figure S2). A secondary analysis using a generalised estimating equation was conducted to determine if baseline SDI and *Staphylococcus* RA influenced changes in the SDI before and after TIP/S initiation, however, no significant differences were observed (data not shown). These findings were sustained when restricted analyses were performed assessing SDI using samples collected at baseline pre (n=41) vs last post (n=41) initiation of TIP/S (Wilcoxon, $p=0.54$). When we assessed the first pre and last post samples we found a −3.8 log fold decrease in *Paremononas* spp ($p=0.0001$) although the base mean reads were low and organisms belonging to this genus may be less biologically relevant (data not shown). Similar findings were observed when TIS and TIP exposed patients were assessed separately. No significant differences were observed in either BC or SDI indices pre vs post initiation of therapy (data not shown).

Next, we assessed microbiome changes in baseline pre versus the last available sample collected separately for each of responders and non-responders. No significant differences were noted for beta-diversity of responders pre-TIP vs post-TIP/S (PERMANOVA, $R^2=0.026$, $p=0.33$). Similar findings were observed SDI where no differences were noted in non-responders pre-TIP vs post-TIP/S ($p=0.20$). Notably, however, non-responders had a −3.1 log 2-fold decrease *Pseudomonas* spp ($p<0.001$) and −2.8 log 2-fold decrease in *Leptotrichia* spp ($p=0.027$) post-TIP/S initiation.

As the sputum microbiome of individuals with mild lung disease (≥70% FEV1) generally trends towards higher SDI and is potentially more susceptible to deviations with medical interventions, we did a subgroup analysis where we assessed only those 15 patients with mild lung disease. No differences were observed in baseline pretherapy and last post-therapy samples collected from patients with mild lung disease in BC beta-diversity, SDI, and no taxa were differentially abundant (data not shown).

Finally, we performed a subgroup analysis in those patients who were TIP/TIS naïve as well as in those who had not had exposure in ≥5 years (in order to eliminate any potential confounding from past prior exposures) and results were similar in diversity measures in both of these subgroups. No significant differences were observed in samples collected pre versus last post for BC beta-diversity or SDI in either the naïve group or the group with no TIP/S exposure in ≥5 years. Similarly, no taxa were differentially abundant in samples collected pre versus last post in patients that were naïve to TIP/S. However, patients who had not had TIP/S exposure in ≥5 years showed significant decrease in *Paremononas* and *Mogibacterium* in the samples collected pre versus last post (data not shown).

**DISCUSSION**

The CF sputum microbiome consists of a diverse group of organisms. The role of this community in health and disease remains to be fully determined, but already it has been associated with age, sex, lung disease stage and the occurrence of PEX.13,18-22 Our group previously assessed changes in the sputum microbiome following the initiation of AZLI21 and during a single cycle of AZLI therapy.23 Our findings suggested that the sputum microbiome is relatively resistant to AZLI although intrapatient differences were observed ‘on’ vs ‘off’ treatment. In both of these studies non-responders had a higher abundance of...
of *Staphylococcus* spp—an organism intrinsically resistant to aztreonam. As *S. aureus* has now become the most prevalent CF pathogen—these results proved intriguing and suggested that baseline microbiota may influence response to therapies—potentially serving as a biomarker to enable treatment personalisation.32 However, both studies were limited with the inclusion of patients with more advanced lung disease (given AZLI’s use as a salvage agent in Canada)—thereby having a less diverse microbiome—potentially less susceptible to perturbation. To address these limitations, here we completed a similar study to understand the effect and association of TIP/S—the first-line treatment for chronic *P. aeruginosa* infection in CF—on CF sputum microbiome.

Interestingly, patients who responded to TIP/S had baseline differences in sputum bacterial community composition compared with non-responders as determined by BC beta-diversity analyses. Such findings may support a role for sputum microbiota to be used as a biomarker for personalisation of therapies. In particular, these differences were characterised by a higher abundance of *Staphylococcus* in baseline samples collected from responders compared with non-responders. This is an intriguing observation, distinctly contrasting what our group has observed with the monobactam AZLI. However, tobramycin—an aminoglycoside—functions by interfering with bacterial protein synthesis by binding the 3OS-ribosomal subunit.33 Like aztreonam, tobramycin’s spectrum of activity is generally confined to aerobic Gram-negative organisms—with the very notable exception of *Staphylococci*.34 Indeed, both methicillin-sensitive *S. aureus* (MSSA) and methicillin-resistant *S. aureus* are generally susceptible to tobramycin.35 36 When we assessed 57 MSSA isolates collected from 30 individuals from this cohort derived from a concomitant pathogen biobank collected a median 44 days (IQR −104 to 337) from TIP/S initiation we determined the median tobramycin minimal inhibitory concentration of these isolates was 0.75 µg/mL (IQR0.5–1.0)—well below the levels achieved with TIP/S34 37 (data not shown). Given the increasing prevalence of MSSA (infected 53.6% of Canadians in 2017), having recently surpassed *P. aeruginosa* (infected 40.2%) as the most prevalent pathogen—new treatment strategies targeting this organism are required.12 In contrast, responders had lower RA of *Pseudomonas*, the intended target of TIP/S, compared with non-responders. Together, these data suggest that the current practice of only using cycled TIP/S in *P. aeruginosa* chronically infected patients may be inappropriately exclusionary. Future clinical trials assessing the potential clinical benefit of TIP/S for patients with CF with chronic *S. aureus* infection may thus be in order. As this drug is already licensed, and care providers are very familiar with its use—a prospective randomised control trial demonstrating clinical benefit could immediately result in patient access. Indeed, ‘off-label’ TIP/S use in *P. aeruginosa* uninfected individuals may already be common in PEx prone individuals.

Furthermore, responders trended towards higher SDI compared with non-responders at baseline. Concomitant with these findings, responders demonstrated a lower *Pseudomonas* and higher *Staphylococcus* absolute and RA in contrast to non-responders suggesting a propensity towards response to treatment may be linked to baseline community composition. All of these findings suggest that our use of tobramycin exclusively in those with chronic *P. aeruginosa* infection may in fact be too limited and those with earlier disease (and those without chronic *P. aeruginosa* infection) may have the potential for benefit of TIP/S therapies. When we conducted a ROC curve analysis to evaluate how accurately RA of *Staphylococcus* and SDI predict response—we found RA of *Staphylococcus* was superior to SDI.

A recently published manuscript by Nelson et al follows a cohort of non-naive patients through a single TIP cycle, complements and supports the findings herein as they too have observed the greatest short-term impacts of TIP were on non-canonical sputum constituents and its effects independent of the targeted pathogen.38 In addition to differences in study design and tobramycin exposure, our cohorts have different characteristics including microbial community make-up with *P. aeruginosa* being more abundant in our cohort. Whereas, Nelson et al observed differences in community structure at specifically selected time points where TIP usage was controlled, we did not observe consistent and significant long-term changes in the microbiome before and after TIP/S.37 38 Indeed, these findings mirror our prior observations with AZLI. The most important limitation to our observation is our inability to reliably discern whether patients in our cohort were ‘on’ or ‘off’ TIP/S 28-day cycle in our biobanked samples (historically this clinical information was not documented).

We acknowledge a number of other limitations to our work. Our study focused exclusively on adult patients. We determined sputum samples with a higher abundance of *Staphylococcus* spp were more likely to occur in responders to TIP/S. While we are unable to confirm that these reads were specific to *S. aureus* as opposed to another *Staphylococcus* species (difficult to discern from 16s data)—we confirmed this with *S. aureus*-specific qPCR. Furthermore, *S. aureus* was cultivated in real time from many specimens. Using culture-independent technologies meant, we were unable to determine whether patients in our cohort had *S. aureus* small colony variants and if this was associated with response—relevant as this phenotype has previously been associated with exaggerated lung function decline.39 While our study focused exclusively on inhaled tobramycin, it is likely that most individuals in our cohort had been remotely exposed to parenteral formulations of tobramycin during PEx—and these may have transiently influenced the microbiome although we expect these to be transient shifts, long since resolved.17 Furthermore, in this retrospective study, samples were collected at clinical visits irrespective of TIP/S initiation which is a limitation of our study design.

As the management of CF involves multiple complementary therapies all started at different times—each of which may impact the sputum microbiome—attributing changes to one specific therapy is exceedingly complex. Furthermore, previous studies have shown regional diversity in microbiome of lungs, and significant potential for oropharyngeal contamination which may be not have been appreciated based on our study design using expectorated sputum.40 However, invasive procedures such as bronchoalveolar lavage pose patient risk and are not a feasible for routine screening.40 41 While we focused solely on microbial community composition, future prospective studies could assess functional differences in the metagenome (as done by Nelson et al.) and metatranscriptome—unfortunately impossible for biobanked samples such as ours. Additionally, culture enriched metagenomics may be implemented to gain a better understanding of microbial communities specifically low abundant organisms.42 In the design of future clinical trials, it will be key to biobank samples including urine, serum, sputum, faeces and cultured-bacteria to better understand intricate host-microbial changes in response to different therapeutics.
CONCLUSIONS
Our study confirms previous findings suggesting that the long-term CF sputum microbiome is relatively resistant to change following initiation of inhaled antibiotics. Intriguingly, differences in baseline microbiome were observed between responders and non-responders suggesting a potential for sputum to be used as a precision medicine tool to predict patient response to therapies. Interestingly, responders had a higher abundance of S. aureus at baseline suggesting that this organism may play a role in response to TIP/S although this warrants further investigation.

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Contributors All authors meet requirements for authorship having contributed significantly to the project. AH, CT and MDP designed the study. AH, CT and MDP collected clinical data. HR, RS and MDP collected patient samples. AH performed experiments and was responsible for data analysis. Sample processing and statistical analyses were performed by AH, NA, ILL, LR, DS, MS and MDP. AH wrote the initial draft of the manuscript, and all authors contributed to its revision. MDP supervised the study and serves as guarantor of the work.

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Competing interests MDP, HR and MS have received research support from CF Canada, CIHR and Gilead.

Patient consent for publication Not required.

Ethics approval All patients provide prospective consent for the collection and storage of sputum samples for research purposes as approved by the Calgary Health Region Ethics Board (REB15-0854).

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data from this project are available at; https://www.ncbi.nlm.nih.gov/bioproject/662963

REFERENCES
4 Lipuma JJ. Microbiological and immunologic considerations with aerosolized drug delivery. Chest 2001;120:1185–23.