Parallel Analysis of Cystic Fibrosis Sputum and Saliva Reveals Overlapping Communities and an Opportunity for Sample Decontamination

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ABSTRACT Culture-independent studies of the cystic fibrosis (CF) airway microbiome typically rely on expectorated sputum to assess the microbial makeup of lower airways. These studies have revealed rich bacterial communities. There is often considerable overlap between taxa observed in sputum and those observed in saliva, raising questions about the reliability of expectorated sputum as a sample representing lower airway microbiota. These concerns prompted us to compare pairs of sputum and saliva samples from 10 persons with CF. Using 16S rRNA gene sequencing and droplet digital PCR (ddPCR), we analyzed 37 pairs of sputum and saliva samples, each collected from the same person on the same day. We developed an *in silico* postsequencing decontamination procedure to remove from sputum the fraction of DNA reads estimated to have been contributed by saliva during expectoration. We demonstrate that while there was often sizeable overlap in community membership between sample types, expectorated sputum typically contains a higher bacterial load and a less diverse community compared to saliva. The differences in diversity between sputum and saliva were more pronounced in advanced disease stage, owing to increased relative abundance of the dominant taxa in sputum. Our effort to model saliva contamination of sputum *in silico* revealed generally minor effects on community structure after removal of contaminating reads. Despite considerable overlap in taxa observed between expectorated sputum and saliva samples, the impact of saliva contamination on measures of lower airway bacterial community composition in CF using expectorated sputum appears to be minimal.

IMPORTANCE Cystic fibrosis is an inherited disease characterized by chronic respiratory tract infection and progressive lung disease. Studies of cystic fibrosis lung microbiology often rely on expectorated sputum to reflect the microbiota present in the lower airways. Passage of sputum through the oropharynx during collection, however, contributes microbes present in saliva to the sample, which could confound interpretation of results. Using culture-independent DNA sequencing-based analyses, we characterized the bacterial communities in pairs of expectorated sputum and saliva samples to generate a model for "decontaminating" sputum *in silico*. Our results demonstrate that salivary contamination of expectorated sputum does not have a large effect on most sputum samples and that observations of high bacterial diversity likely accurately reflect taxa present in cystic fibrosis lower airways.

KEYWORDS microbiome, airway infection, microbial community dynamics

Despite the considerable advances made in the care of persons with cystic fibrosis (CF) and the resultant increase in life expectancy in this population during the last 3 decades, chronic airway infection leading to respiratory failure remains the primary...
cause of death. Initially, studies of CF airway infection focused on human-pathogenic bacterial species (Staphylococcus and Haemophilus) recovered in cultures of respiratory secretions. The bacteria of interest in CF gradually expanded to include several opportunistic pathogens including Pseudomonas, Burkholderia, Achromobacter, Stenotrophomonas, and mycobacteria (1). The advent of culture-independent approaches to study the human microbiome has, more recently, enabled a deeper exploration of CF airway microbiology, and studies employing these methods have indicated that CF airways likely harbor more diverse bacterial communities than previously appreciated (2, 3). These new insights hold promise to translate into novel approaches to better manage airway infection, further reducing respiratory system-associated morbidity and mortality in CF.

Sampling of the lower airways to assess the microbial communities therein presents a challenge, however. Although bronchoscopy has been used to obtain airway secretions (i.e., via bronchoalveolar lavage [BAL]) in several studies of CF microbiology (4–10), this approach is invasive, not without risk, and not feasible for studies requiring analysis of serial samples from individuals across a large subject cohort. Expectorated sputum, therefore, has been most often used to address the dynamics of airway bacterial communities in CF. Because this respiratory tract specimen passes through the non-sterile upper airway and oropharynx, however, it must be assumed to be “contaminated” with microbes residing in these sites, raising concerns about the reliability of sputum in reflecting lower airway microbial communities. The degree to which upper airway microbes contribute to the bacterial communities measured in expectorated sputum—and, by extension, the degree to which these species can be held to account for the conclusion that CF lower airways harbor rich bacterial communities—is difficult to ascertain.

To improve our understanding of the relationship between upper and lower airway bacterial communities, we analyzed pairs of expectorated sputum and saliva samples collected from individuals with CF. We estimated the effect of saliva contamination on expectorated sputum and propose a model to account for the contribution of salivary microbiota to measures of sputum bacterial community composition.

RESULTS

Subjects, samples, and DNA sequencing controls. Ten adult subjects with CF provided 37 same-day pairs of sputum and saliva samples. Subject demographics and sample characteristics are summarized in Table 1. The median subject age was 32 years. The median percent predicted forced expiratory volume in 1 s (ppFEV1) was 61, measured within 1 to 45 days of the collection day. The number of sample pairs per subject ranged from one to five. For subjects with more than one sample pair, the period of time between the first and last sample pair averaged 94 days (range 8 to 330 days), and the time between consecutive samples averaged 32 days (range 7 to 253 days). Six sample pairs were collected during antibiotic treatment (i.e., treatment

| Subject | No. of pairs | Sex | CFTR genotype | Age (yr) at 1st sample | Disease stagea |
|---------|-------------|-----|---------------|------------------------|---------------|
| 1       | 5           | F   | F508del homzygous | 19                    | Int           |
| 2       | 5           | F   | F508del homzygous | 30                    | Early/Int    |
| 3       | 5           | F   | F508del heterozygous | 26                   | Int/Adv      |
| 4       | 5           | M   | F508del heterozygous | 21                    | Early        |
| 5       | 5           | F   | F508del homzygous | 56                    | Early/Int    |
| 6       | 1           | F   | F508del heterozygous | 51                   | Int           |
| 7       | 2           | M   | F508del heterozygous | 32                   | Int           |
| 8       | 3           | F   | F508del homzygous | 29                    | Int           |
| 9       | 3           | M   | F508del heterozygous | 35                   | Adv           |
| 10      | 3           | F   | Other         | 32                    | Int           |

aDisease stage was assigned as described in the text. Int, intermediate; Adv, advanced.
other than maintenance antibiotic use). Only one subject had advanced lung disease, but all spontaneously expectorated sputum on a daily basis.

The median DNA sequencing error rate, based on mock community analyses, was 0.032% (range 0.025% to 0.036%). The median bacterial load of reagent controls was $4.2 \times 10^3$ (range $3.8 \times 10^3$ to $4.5 \times 10^3$) 16S rRNA gene copies/ml, which was more than 5 logs lower than the median bacterial load of sputum and saliva samples. Neither the reagent nor water controls amplified during library construction. The mean pairwise theta-YC similarity of 59 generous donor control replicates was 0.983 (range 0.931 to 0.998).

**Total bacterial load, bacterial community richness, and community structure.**

In the aggregate, the bacterial load (expressed as 16S rRNA gene copies/ml) in sputum samples (median $1.4 \times 10^3$; range $8.3 \times 10^1$ to $1.0 \times 10^3$) was greater than the bacterial load in saliva samples (median $5.4 \times 10^3$; range $9.3 \times 10^5$ to $7.0 \times 10^9$), although this difference failed to reach statistical significance (linear mixed model, $P = 0.054$; Fig. 1A). For 25 of 37 pairs, the bacterial load of the sputum sample was greater than that of the corresponding saliva sample (Fig. 1B). The median bacterial load difference between paired sputum and saliva samples was $9.3 \times 10^8$ more 16S rRNA gene copies/ml in sputum than in saliva (range, $3.8 \times 10^9$ fewer to $9.8 \times 10^9$ more). Considering all samples (37 saliva and 37 sputum) together, bacterial load was not significantly different between disease stage groups (linear mixed model, $P = 0.655$; Fig. 1C). The bacterial load of all samples collected during antibiotic treatment was significantly lower than that of all other samples (linear mixed model, $P = 0.038$; see Table S1 in the supplemental material). Saliva samples collected during antibiotic treatment had lower bacterial load compared to nontreatment saliva samples, treatment sputum samples, and nontreatment sputum samples; however, there was no significant interaction between sample type and antibiotic treatment status in the model (linear mixed model, $P = 0.176$; Table S1).

The median bacterial richness (number of observed operational taxonomic units [OTUs]) in sputum samples was significantly lower than in saliva (linear mixed model, $P = 0.021$; Fig. 2A). For 25 of 37 pairs, fewer taxa were observed in sputum compared to the respective saliva sample (Fig. 2B). The median difference between saliva and sputum richness was six OTUs (range, $-16$ to $29$). For all sputum and saliva samples in the aggregate, bacterial richness was not significantly different between disease stage groups (Fig. 2C), nor between on- or off-antibiotic-treatment groups (linear mixed model, $P = 0.958$ [disease stage group] and $P = 0.829$ [antibiotic treatment groups]).

A wide range of overlap in community membership and structure was observed in sputum-saliva sample pairs (Fig. 3). The median theta-YC similarity between each sputum and paired saliva sample was 0.400 (range 0.029 to 0.856). For context, the median theta-YC similarity between all sputum samples from all subjects was 0.207, while the median theta-YC similarity between 59 replicate generous donor control samples was 0.985. Community structures were significantly more similar for sample pairs collected from subjects at earlier (early plus early/intermediate) disease stage compared to later (intermediate plus advanced) disease stage (Fig. 4A; $t$ test, $P = 0.007$). Pairs from subjects in the later disease stage group tended to have lower Jaccard similarity, although this difference was not significant (Fig. 4B; $t$ test, $P = 0.105$). The relative abundance of the dominant OTU in sputum was significantly greater than the relative abundance of the dominant OTU in the respective saliva sample for sample pairs in the later disease stage group only (Fig. 4C; paired $t$ tests $P < 0.001$ [later disease stage] and $P = 0.239$ [earlier disease stage]). Community structures (theta-YC and Jaccard) were not significantly different between sputum and saliva sample pairs collected during antibiotic treatment compared to all other pairs ($t$ tests, $P = 0.119$ [theta-YC] and $P = 0.979$ [Jaccard]).

**In silico sputum decontamination.** Bacteria in sputum that originate solely via contamination from saliva during expectoration would be expected to be present in approximately the same relative proportions as are found in the respective saliva sample. In other words, regardless of the volume of saliva that contaminates a sputum...
Bacterial load (16S rRNA gene copies/ml) in sputum and saliva samples. (A) In the aggregate, bacterial load was greater in sputum samples than in saliva samples, although this difference failed to (Continued on next page).
sample, the taxa present in that volume of saliva would add proportionately to the sputum sample. Furthermore, the abundance of each taxon observed in a given volume of expectorated sputum that can be ascribed to saliva contamination is limited by the abundance of that taxon in a comparable volume of saliva. We used this conceptual framework to estimate the maximum possible contribution of taxa from saliva to expectorated sputum samples.

For each sputum-saliva sample pair, the following procedure was performed, as illustrated in Fig. 5. OTUs with at least 1% relative abundance in the saliva sample (top saliva OTUs) were identified; OTUs with <1% relative abundance in the saliva sample were not included to avoid false negatives at the lower limits of sequence detection. The absolute abundance of each of the top saliva OTUs was calculated by multiplying the total bacterial load in the saliva sample (quantified by 16S rRNA droplet digital PCR [ddPCR]) by the relative abundance of the OTU in that sample. The absolute abundance of each of the top saliva OTUs present in the corresponding sputum sample was similarly calculated. The ratios of the top saliva OTUs in sputum-saliva sample pairs were calculated by dividing the absolute abundance of each of these OTUs in sputum by its absolute abundance in saliva. The lowest resulting ratio value among all top saliva OTUs in each sample pair was identified as the contamination constant for that sample pair. For pairs where the lowest resulting value was greater than 1 (i.e., the abundances of all top saliva OTUs in sputum were greater than their corresponding abundances in saliva), the contamination constant was set at 1. For sample pairs where a top saliva OTU was absent in sputum, the contamination constant for the pair was set at 0. Next, for each shared OTU in a saliva-sputum sample pair, the absolute abundance of the OTU in the saliva sample was multiplied by the contamination constant for that pair, resulting in the calculated contamination value for that OTU. This value was subtracted from the absolute abundance of that OTU in the respective sputum sample to yield a corrected abundance of the OTU in sputum. The resulting community structure, based on the corrected abundances of all shared OTUs, is the corrected community in the sputum sample.

The mean contamination constant among the 37 sputum-saliva sample pairs was 0.230 (range 0 to 1) (Table S2). The mean theta-YC similarity of pairs of sputum samples (each pair consisting of a sample pre- and post-in silico decontamination) was 0.970 (range 0.309 to 1). There was no significant effect of disease stage group, antibiotic treatment status, sputum alpha diversity, or sputum bacterial load on the similarity of sputum before and after decontamination (linear mixed model, \( P = 0.375 \) [disease stage group], \( P = 0.589 \) [antibiotic treatment status], \( P = 0.427 \) [sputum alpha diversity], and \( P = 0.886 \) [sputum bacterial load]). For the five subjects who each contributed five pairs of samples, the decontamination procedure did not significantly change the bacterial community structure in sputum samples (Fig. 6, analysis of molecular variance [AMOVA] \( P > 0.01 \)).

**DISCUSSION**

Although chronic lung infection and inflammation is the primary cause of morbidity and mortality in CF, the microbiology of CF lung infection is not well understood. Whereas culture-based methods to study CF airway microbiology have focused on a narrow set of opportunistic bacterial pathogens, culture-independent analyses of CF
FIG 2  Bacterial community richness (number of observed OTUs) in sputum and saliva samples. (A) In the aggregate, bacterial richness was greater in saliva samples than in sputum samples (paired t test, (Continued on next page)
respiratory specimens have revealed bacterial communities that can be quite diverse (11–15). Expectorated sputum is most commonly used in these studies, but the reliability of this specimen in reflecting lower airway microbiota is a concern, considering that expectoration involves passage of sputum through the nonsterile upper airway and oral cavity.

Few studies have addressed this concern directly. Goddard and colleagues (16) used sterile technique to obtain endobronchial samples from the airways of explanted CF lungs from several patients undergoing lung transplantation, thereby circumventing

FIG 2 Legend (Continued)
P = 0.021). (B) For 25 of 37 pairs, the bacterial richness of the saliva sample was greater than that of the corresponding sputum sample. Sample pairs collected during antibiotic treatment are marked with a “T.” The vertical black bars at 0 indicate samples where sputum and saliva samples have equal richness. (C) Richness of all samples (37 saliva and 37 sputum samples) was not significantly different between subjects with early or early/intermediate disease stage (n = 15 sample pairs from 3 subjects) and subjects with intermediate, intermediate/advanced, or advanced disease stage (n = 22 sample pairs from 7 subjects). The top and bottom boundaries of each box indicate 75th and 25th quartile values, respectively; black lines inside each box represent the median values.

FIG 3 Relative abundance of taxa observed in 37 pairs of expectorated sputum and saliva samples collected from 10 subjects, showing individual OTUs with an average relative abundance >1% across all samples. "Other" taxa combine remaining OTUs in each sample. Disease stage is indicated above each subject’s plots.
FIG 4  Community similarity of 37 pairs of expectorated sputum and saliva. (A) Community structures were significantly more similar (theta-YC similarity) in sample pairs from subjects with early or early/ (Continued on next page)
the possibility of sample contamination by upper airway secretions. Using DNA sequencing, they compared the bacterial species detected in lower respiratory tract airways to the species found in paired upper airway samples from the same patients and observed airway communities with very low diversity, heavily dominated by “typical” CF pathogens. The paucity of oropharyngeal microbiota in the endobronchial samples led the authors to question the reliability of expectorated sputum in reflecting the lower airway microbiome in CF. As the authors acknowledge, however, analysis of explanted lungs is, by definition, representative of end-stage CF lung disease, wherein very low bacterial community diversity, owing to the overwhelming dominance of a single species, is typically observed (12, 13, 17). Thus, the relevance of this study to airway microbiology in the majority of CF patients who are at earlier stages of lung disease is limited. In a subsequent investigation, Brown and colleagues (18) dissected the diseased lung tissue of a 3-year-old child who underwent lobectomy for severe localized CF lung disease and similarly investigated the microbiota therein. In addition to typical CF pathogens, including *Staphylococcus* and *Haemophilus*, a diverse bacterial population was detected that included *Ralstonia* and anaerobic species.

More recently, Jorth and colleagues (4) reported that DNA sequence analysis of BAL fluid samples from children and young adults with mild CF lung disease showed no evidence of oral bacterial communities in samples that lacked typical CF pathogens. They concluded that established CF pathogens are primarily responsible for CF lung infections. The BAL fluid samples from these relatively healthy young CF patients (mean ppFEV₁ of 92; mean age 13 years; median age not provided) generally had bacterial DNA concentrations (median ~10³ bacterial genome copies/ml) several orders of magnitude lower than the levels found in sputum of expectorating adults with CF (12, 19–21), making it difficult to extrapolate these findings more broadly to the CF population.

In our study, we similarly observed bacterial DNA concentrations in sputum (median ~10⁹ 16S gene copies/ml) far exceeding those measured by Jorth and colleagues (4) in BAL fluid samples from younger patients. Whereas sample contamination introduced by laboratory reagents and/or during sample collection is a critically important concern in microbiome analyses of low-biomass samples (22–24), the impact would be expected to be considerably less in the analysis of samples with very large biomass. In fact, we observed that bacterial DNA levels in our reagent controls were several logs lower than those found in sputum and saliva samples. More relevant to our study, however, was our observation that bacterial loads in sputum were generally greater than those in saliva (Fig. 1A and B), which would considerably mitigate the impact of salivary contamination of expectorated sputum relative to what would be expected with salivary contamination of very-low-biomass BAL fluid samples. Consistent with previous work (25), we found that antibiotic treatment was not a significant factor contributing to differences in bacterial load or bacterial richness.

Our analyses of bacterial community structures revealed considerable overlaps in membership within pairs of sputum and saliva samples. This is not surprising considering that sputum and saliva samples each likely represent a mixture of the two. Nevertheless, detecting taxa in sputum that are known to inhabit the upper airways and oropharynx raises reasonable concerns about the source of these species. Assessing lower airway infection in CF from an ecological theory perspective, however,
provides a cogent conceptual framework whereby the upper airway serves as a source of microbes that may migrate to and persist in the lower airways (26). Microbiologic continuity of the aerodigestive tract has been identified in healthy individuals (27) as well as in persons with CF (28), where similarity between oral and gastric communities indicates connectivity between mouth, lung, and upper gastrointestinal microbiomes. In this biogeography context, continuously mixed microbial communities inhabit these contiguous anatomic sites, with community composition in each determined by rates of migration, colonization/infection, and extinction (26, 29).

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**FIG 5** Illustration of *in silico* sputum decontamination procedure using a hypothetical pair of sputum and saliva samples.

For each sputum-saliva pair:
- Determine OTU relative abundances in sputum ($R_{spu}$) and saliva ($R_{sal}$) by 16S rRNA gene sequencing
- Measure total bacterial loads in sputum and saliva by ddPCR
- Calculate OTU absolute abundances as ($R_A$) x (total bacterial load) for sputum ($A_{spu}$) and saliva ($A_{sal}$)

For each top saliva OTU with >1% $R_{sal}$:
- Divide $A_{spu}$ by $A_{sal}$
- The minimum $A_{spu}/A_{sal}$ value in the sputum-saliva pair is the contamination constant ($K_C$) for that pair
- If the minimum value is > 1, then $K_C = 1$; if a saliva OTU is absent in sputum, then $K_C = 0$

For each shared OTU in a sputum-saliva pair:
- Multiply $A_{sal}$ by $K_C$ for that pair to yield the calculated contamination ($C_C$) value of that OTU in sputum

For each shared OTU in a sputum-saliva pair:
- Subtract $C_C$ from $A_{spu}$ to yield the corrected abundance ($C_A$) of the OTU in sputum
FIG 6 (A) Theta-YC based PCoA of bacterial community structures in sputum from five subjects who each provided five pairs of samples. Sputum before (closed circles) and after (open circles) in silico (Continued on next page)
As has been shown in several previous studies (12, 13, 17), we found that sputum bacterial community diversity decreased with lung disease progression, primarily due to reduced community evenness resulting from the progressive dominance of a single species. This trend was not observed in the corresponding saliva samples, however. Further, we found, as have others (30), that sputum and saliva bacterial loads remained relatively constant with advancing lung disease. Given this consistency in bacterial load and salivary community diversity across disease stages, it is improbable that salivary contamination would account for high sputum community diversity during early stages of lung disease but not during late disease stages.

We found that despite the overlap in bacterial community membership between paired sputum and saliva samples, we observed, in some pairs, taxa that were disproportionately represented in one sample or the other. We noted, for example, some sample pairs with comparable bacterial loads in which a taxon was present in high relative abundance in saliva but present in very low relative abundance or not detected at all in the corresponding sputum sample. Conversely, we observed some sample pairs in which a specific taxon was present in much greater relative abundance in sputum than in the corresponding saliva sample. Our in silico decontamination model is premised on the principle that the contribution of species from saliva to sputum must be proportionate to the relative abundance of species in saliva. Taking total bacterial load and taxon relative abundance into account in this model allows an estimation of the maximum salivary contamination of sputum for each sample pair, providing a liberal estimate of potential contamination and a means to “decontaminate” expectorated sputum in silico.

While the thrust of our study was to estimate the impact of salivary microbiota on measures of microbial communities in sputum, this effort allowed us to consider previously described software packages for estimating the proportion of contaminants in microbial communities of interest. Programs such as SourceTracker (31) and decontam (32) seek to objectively remove entire OTUs or genera based on their a priori identification as contaminants, or their low abundance, or their presence in control samples. In contrast, our approach considers the absolute abundances of taxa and estimates contamination separately for each pair of sputum and saliva samples. This strategy is more similar to microDecon (33), which also addresses overlapping OTUs (those OTUs expected to occur in both the sample and the contamination source) based on the principle that contamination from a common source will be proportionate and can be calculated using a “constant” OTU that is assumed to be entirely contamination. The difference of our approach is that for each sample and source pair, we use the estimation of maximum contamination to establish an upper limit of contamination, rather than make a probabilistic estimation of contamination (SourceTracker) or remove taxa entirely (decontam). The primary goal of our method was to evaluate the potential for saliva contamination of expectorated sputum, which in turn establishes the utility of sputum as a measure of lower airway microbiota in CF. The calculated saliva contamination of sputum samples therefore represents a “worst-case scenario.”

Despite this aggressive approach, the impact of saliva contamination on sputum samples was estimated to be minimal. In fact, the mean sputum community structure similarity before and after decontamination was within the range of technical (sequencing) replicate controls, with only three of 37 sputum samples falling outside this range, suggesting that much of the diversity observed in sputum (i.e., due to nontypical CF pathogens) was present prior to passage through the oral cavity during expectoration. While we do not think in silico decontamination should be a necessary step for all CF...

FIG 6 Legend (Continued)
decontamination procedure is shown for each sample (before/after decontamination sputum pairs are connected by lines). Subjects are depicted in different colors. Fifty-nine technical replicates (generous donor controls) are shown as open gray triangles. (B) Using the same ordination data as in panel A, subjects and controls are visualized separately with the axes for each plot set so that the samples fill the ordination space. As a result, the scales between plots are not comparable.
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Microbiome studies employing expectorated sputum, it is nevertheless possible that even the minor changes in community structures observed after decontamination may reveal dynamics of interest that would have been otherwise obscured. In subject 5, for example, the bacterial community structures of four of five sputum samples shifted in a similar fashion after in silico decontamination. In this regard, the impact of salivary contamination on community structure—and whether this constitutes a meaningful change—likely varies between subjects and over time within subjects and ultimately depends on the question at hand. Importantly, this study provides evidence that the diversity of bacterial communities measured in CF sputum cannot be attributed solely to the contamination of sputum by salivary microbiota during expectoration. As such, these results support the use of expectorated sputum as a measure of lower airway microbiota in CF and do not corroborate reports asserting that CF lung microbial diversity is predominantly an artifact of sampling methods.

Materials and Methods

Study design and sample collection. Expectorated sputum and saliva samples were collected from a cohort of persons with CF as part of a long-term study of CF airway microbiota. This study was approved by the Institutional Review Board of the University of Michigan Medical School (HUM0037056), and informed written consent was obtained from all participants. Sputum and saliva samples were collected by subjects at home in 30-ml sterile conical tubes. Subjects were reminded that sputum was the thick material from deep in the lungs and not the watery fluid in their mouth. Subjects were instructed to spit out saliva and rinse their mouth with tap water before taking a deep breath and coughing deeply to move sputum up from their lungs. Sputum was then expectorated directly into a collection tube. For saliva samples, subjects were asked not to eat or drink for 30 min prior to collection. Subjects were instructed to allow saliva to collect in their mouths for at least 1 min and then gently spit or drool into the collection tubes. Samples were immediately stored in a −20°C manual defrost freezer (1 to 10 weeks; mean 4 weeks) before same-day courier transport on dry ice to the University of Michigan and immediate storage at −80°C. We have previously shown that sputum storage at −20°C does not have a significant effect on sequencing-based measures of bacterial community structure (34).

Electronic medical records were reviewed for subject demographic and clinical data. Daily antibiotic use was recorded by subjects on the day the samples were collected, and subjects were classified as either on or not on treatment antibiotics (i.e., antibiotics other than chronic maintenance antibiotics such as inhaled tobramycin and oral azithromycin) at the time of sample collection. Disease stage was assigned based on ppFEV₁ over the period of sample collection: early (ppFEV₁ ≥ 70), intermediate (70 ≥ ppFEV₁, ≥ 40), or advanced (ppFEV₁ < 40) (35, 36). When subjects crossed between disease stage categories during the course of this study, they were classified as either early/intermediate or intermediate/advanced.

Sample processing and 16S rRNA gene sequencing. Sputum and saliva samples were thawed on ice prior to homogenization in 10% Sputolysin (MilliporeSigma, Burlington, MA, USA). Samples were treated with bacterial lysis buffer (Roche Diagnostics Corp., Indianapolis, IN, USA), lysozyme (MilliporeSigma), and lysostaphin (MilliporeSigma) as previously described (12), followed by mechanical disruption by glass bead beating and digestion with protease K (Qiagen Sciences, Germantown, MD, USA). The MagNA Pure nucleic acid purification platform (Roche Diagnostics Corp.) was used to extract and purify DNA according to the manufacturer’s protocol. Reagent control samples were similarly prepared, with UltraPure DNase/RNase-free distilled water (Life Technologies Corp., Grand Island, NY, USA) substituted for the sample.

DNA libraries were prepared by the University of Michigan Microbial Systems Molecular Biology Laboratory as described previously (37). In brief, the V4 region of the bacterial 16S rRNA gene was amplified using touch-down PCR with barcoded dual-index primers. The touch-down PCR cycles consisted of 2 min at 95°C, followed by 20 cycles of 95°C for 20 s, 60°C (starting from 60°C, the annealing temperature decreased 0.3°C each cycle) for 15 s, and 72°C for 5 min, followed by 20 cycles of 95°C for 20 s, 55°C for 15 s, and 72°C for 5 min and a final 72°C for 10 min. Resulting amplicon libraries were normalized and sequenced on an Illumina sequencing platform using MiSeq reagent kit V2 (Illumina Inc., San Diego, CA, USA). The final load concentration was 4.0 to 5.5 pM with a 15% PhiX spike to add diversity.

DNA sequencing controls and DNA sequence analyses. Reagent-only controls were prepared with each change in reagent lot. Mock bacterial community DNA standards (ZymoBIOMICS Microbial Community DNA; Zymo Research, Irvine, CA, USA) and water controls were included in sequencing runs by the University of Michigan Microbial Systems Molecular Biology Laboratory. Sequences from reagent and water controls were analyzed to assess the impact of laboratory contaminants on sputum and saliva sample DNA sequencing results. Mock community sequences were used to measure sequencing error rates. DNA extracted from a “generous donor” CF sputum sample was also sequenced to determine interrun sequencing variability. Sequencing data from 59 replicates of the generous donor control were analyzed.

Raw DNA sequences were analyzed with mothur v1.41.3 (38) following the mothur MiSeq SOP (full list of mothur commands available at https://github.com/um-cf-microbiome/LI/Puma_Sputum-Saliva). Briefly, low-quality and chimeric reads were discarded, and remaining sequences were classified taxo-
nomically against the SILVA database (release 132) using the RDP Bayesian Classifier. Sequences were clustered into OTUs using a 3% dissimilarity cutoff into the OptiClust algorithm (39). To limit the effects of sequencing depth, each sample was rarefied to the lowest number of reads in the sample set ($n = 2,076$). Alpha diversity of the subsampled data was measured as observed OTUs (richness). Beta diversity was calculated based on the Jaccard (40) or Yue and Clayton (“theta-YC”) (41) measures of dissimilarity and, for ease of interpretation, are reported as similarity (i.e., $1 - \text{dissimilarity}$). Theta-YC was chosen to describe dissimilarity between communities, taking into consideration the proportions of both the shared and nonshared members of each community. An advantage of this metric compared to other dissimilarity metrics is that it is less sensitive to outlier values and weighs rare and abundant OTUs more evenly than the Bray-Curtis or Morisita-Horn (42) metric.

ddPCR. Total bacterial load of sample and control DNAs was quantified by 16S rRNA droplet digital PCR (ddPCR) on a QX200 AutoDG droplet digital PCR system (Bio-Rad Laboratories, Inc., Hercules, CA, USA). Primer sequences were 5′-TCTCCAGGGAACGAGCT-3′ and 5′-GGACTACGGGCGGTATCTAATCCT G-3′ (final concentration 900 nM each), and probe sequence was 6-carboxyfluorescein [FAM]-5′-CGTA TTACCGGCCGCTGTTG-3′- (6-carboxytetramethylrhodamine [TAMRA]) (final concentration 250 nM). All reactions were run in duplicate. Reaction mixtures were transferred to an automated droplet generator (Bio-Rad Laboratories, Inc.), followed by gene amplification in a C1000 Touch thermal cycler (Bio-Rad Laboratories, Inc.). Cycling conditions were 10 min at 95°C, followed by 40 cycles at 94°C for 30 s and 58°C for 2 min, and a final 98°C for 10 min, with a ramp rate of 2°C/s per step. DNA quantification was performed with the QX200 droplet reader (Bio-Rad) and data analysis with QuantaSoft Analysis Pro (Bio-Rad) using default parameters for threshold amplification. Reaction mixtures with fewer than 10,000 droplets were omitted from analysis. DNA concentrations of replicates were averaged, adjusted for dilution factor, reported in copies of target gene per microliter of DNA, and then converted to copies of target gene per milliliter of sample based on the DNA extraction steps.

Data analyses. A linear mixed effects model was used to compare bacterial load between sputum and saliva samples (R packages ‘lme4’ [43] and ‘lmerTest’ [44]), with bacterial load as the dependent variable; sample type, disease stage group, antibiotic treatment status, and an interaction term for antibiotic treatment and sample type as fixed effects; and intercept for subject as a random effect. A linear mixed effects model was also used to compare bacterial richness between sputum and saliva samples, with richness (number of observed OTUs) as the dependent variable; sample type, disease stage group, and antibiotic treatment status as fixed effects; and intercept for subject as a random effect. t tests were performed to compare sputum-saliva theta-YC and Jaccard similarities between disease stage groups and between antibiotic treatment status groups. Paired t tests were performed to compare dominant OTU (the OTU with the highest relative abundance in the sample) relative abundances between sample types within disease stage groups.

A theta-YC-based principal-coordinate analysis (PCoA) plot was generated to visualize the effects of in silico saliva decontamination on sputum samples and to compare these effects to repeat sequencing of a generous donor control sample. Analysis of molecular variance (AMOVA) was used to compare differences in centroids of sputum bacterial communities within individual subjects before and after decontamination (mothur, v1.41.3). A Bonferroni-corrected $P$ value of 0.01 was used to assess significance. A linear mixed-effects model was used to model the impact of sample variables on the effect of the sputum decontamination procedure, with theta-YC similarity of sputum before and after decontamination as the dependent variable; sputum alpha diversity (as inverse Simpson index), sputum bacterial load, disease stage group, and antibiotic treatment status as fixed effects; and intercept for subject as a random effect.

Data availability. Raw sequence data are available at NCBI (SRA accession no. PRJNA611611). Detailed protocols, mothur log file, deidentified subject data, 16S rRNA gene sequencing data, 16S ddPCR data, and reproducible code for data analyses and figures are available on GitHub (https://github.com/um-cf-microbiome/LiPuma_Sputum-Saliva).

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

TABLE S1, PDF file, 0.01 MB.
TABLE S2, PDF file, 0.02 MB.

ACKNOWLEDGMENTS

This work was supported by the National Institutes of Health (R01HL136647) and the Cystic Fibrosis Foundation (LIPIUMA15P0).

We thank the study subjects for their generous participation and dedication.

REFERENCES

1. LiPuma JJ. 2010. The changing microbial epidemiology in cystic fibrosis. Clin Microbiol Rev 23:329–323. https://doi.org/10.1128/CMR.00068-09.
2. Huang YJ, LiPuma JJ. 2016. The microbiome in cystic fibrosis. Clin Chest Med 37:59–67. https://doi.org/10.1016/j.ccm.2015.10.003.
3. O’Toole GA. 2018. Cystic fibrosis airway microbiome: overturning the old, opening the way for the new. J Bacteriol 200:e00561-17. https://doi.org/10.1128/JB.00561-17.
4. Jorth P, Ehsan Z, Rezayat A, Caldwell E, Pope C, Brewington JJ, Goss CH, Benscoter D, Clancy JP, Singh PK. 2019. Direct lung sampling indicates that established pathogens dominate early infections in children with
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cystic fibrosis. Cell Rep 27:1190–1204. https://doi.org/10.1016/j.celrep.2019.03.086.
5. Hogan DA, Willger SD, Dolben EL, Hampton TH, Stanton BA, Morrison HG, Sogin ML, Czum J, Ashare A. 2016. Analysis of lung microbiota in bronchoalveolar lavage, protected brush and sputum samples from subjects with mild-to-moderate cystic fibrosis lung disease. PLoS One 11:e0149998. https://doi.org/10.1371/journal.pone.0149998.
6. Muhlebach MS, Zorn BT, Esther CR, Hatch JE, Murray CP, Turkovic L, Ranganathan SC, Boucher RC, Stick SM, Wolfgang MC. 2018. Initial acquisition and succession of the cystic fibrosis lung microbiome is associated with disease progression in infants and preschool children. PLoS Pathog 14:e1006798. https://doi.org/10.1371/journal.ppat.1006798.
7. Frayman KB, Armstrong DS, Carzino R, Ferkol TW, Grinwood K, Storch GA, Teo SM, Wylie KM, Ranganathan SC. 2017. The lower airway microbiota in early cystic fibrosis lung disease: a longitudinal analysis. Thorax 72:1104–1112. https://doi.org/10.1136/thoraxjnl-2016-209279.
8. Pittman JE, Wylie KM, McEwen ET, Hatch JE, Quante J, Frayman KB, Clarke N, Davis M, Stick SM, Hall GL, Montgomery G, Ranganathan S, Davis SD, Ferkol TW, Australian Respiratory Early Surveillance Team for Cystic Fibrosis. 2017. Association of antibiotics, airway microbiome, and inflammation in infants with cystic fibrosis. Ann Am Thorac Soc 14:1548–1555. https://doi.org/10.1513/AnnalsATS.201712-1201OC.
9. Renwick J, McNally P, John B, DeSantis T, Linnane B, Murphy P, SHIELD Investigators. 2011. Changes in cystic fibrosis airway microbiota at pulmonary exacerbations. PLoS One 6:e172811. https://doi.org/10.1371/journal.pone.0172811.
10. Willner D, Haynes MR, Furlan M, Schmieder R, Lim YW, Rainey PB, Rohwer F, Conrad D. 2012. Spatial distribution of microbial communities in the upper cystic fibrosis lung. ISME J 6:471–474. https://doi.org/10.1038/ismej.2011.104.
11. Cox MJ, Allgaier M, Taylor B, Baek MS, Huang YJ, Daly RA, Karaoz U, Yu C, Marchuk DA, Ranganathan S. 2016. Analysis of lung microbiota in cystic fibrosis. Cell Rep 27:1190–1204. https://doi.org/10.1016/j.celrep.2017.08.007.
12. Zhao J, Schloss PD, Kallinik LM, Carvajal JD, VanDeVanter DR, Murray S, Li JZ, Young VB, LiPuma JJ. 2012. Decade-long bacterial community dynamics in cystic fibrosis airway. Proc Natl Acad Sci U S A 109:5809–5814. https://doi.org/10.1073/pnas.1105771109.
13. Coburn B, Wang PW, Diaz Caballero J, Clark ST, Brahma V, Donaldson S, Zhang Y, Sundeen A, Gony G, Tullis ED, Yau YC, Waters VS, Hwang DM, Gutman DS. 2014. Longitudinal lung microbiota across age and disease stage in cystic fibrosis. Sci Rep 5:10241. https://doi.org/10.1038/srep10241.
14. Whelan FJ, Heirali AA, Rossi L, Rabin HR, Parkins MD, Sogin ML, O’Toole GA. 2013. Unique microbial communities persist in individual cystic fibrosis patient throughout a clinical exacerbation. Microbiome 1:27. https://doi.org/10.1186/2049-2618-1-27.
15. Zemanick ET, Harris JK, Wagner BD, Robertson CE, Sagel SD, Stevens MJ, Accurso FJ, Laguna TA. 2013. Inflammation and airway microbiota during cystic fibrosis pulmonary exacerbations. PLoS One 8:e62917. https://doi.org/10.1371/journal.pone.0062917.
16. Saltor SJ, Cox MJ, Turek EM, Calus ST, Cookson WO, Moffatt MF, Turner P, Parkhill J, Loman NJ, Walker AW. 2014. Reagent and laboratory contamination, can critically impact sequence-based microbiome analyses. BMC Biol 12:87. https://doi.org/10.1186/1471-2199-12-87.
17. Kim D, Hofstader CE, Zhao C, Mattel L, Tanes C, Clarke E, Lauder A, Sillier-Mix S, Chehound C, Kelsen J, Conrad M, Collman RG, Baldassano R, Bushman FD, Bittinger K. 2017. Optimizing methods and dodging pitfalls in microbiome research. Microbiome 5:52. https://doi.org/10.1186/s40168-017-0267-5.
18. Eisenhofer R, Minich JJ, Carzino R, Cooper A, Knight R, Weyrich LS. 2019. Contamination in low microbial biomass microbiome studies: issues and recommendations. Trends Microbiol 27:105–117. https://doi.org/10.1016/j.tim.2018.11.003.
19. Fodor AA, Klem ER, Gilpin DF, Elborn JS, Boucher RC, Tunney MM, Wolfgang MC. 2012. The adult cystic fibrosis airway microbiota is stable over time and infection type, and highly resilient to antibiotic treatment of pulmonary exacerbations. PLoS One 7:e45001. https://doi.org/10.1371/journal.pone.0045001.
20. Whiteson KL, Bailey B, Bergkessel M, Conrad D, Delhaes L, Felts B, Harris JK, Hunter R, Lim YW, Maughan H, Quinn R, Salamon P, Sullivan J, Wagner BD, Rainey PB. 2014. The upper respiratory tract as a microbial source for pulmonary infections in cystic fibrosis. Parallel from island biogeography. Am J Respir Crit Care Med 189:1309–1315. https://doi.org/10.1164/rccm.201312-2129PP.
21. Basisz CM, Erb-Downward JR, Dickson RP, Freeman CM, Schmidt TM, Young VB, Beck JM, Curtis JL, Huffnagle GB. 2015. Analysis of the upper respiratory tract microbiome as the source of the lung and gastric microbiota in healthy individuals. mBio 6:e00037-15. https://doi.org/10.1128/mBio.00037-15.
22. Al-Momani H, Perry A, Stewart CJ, Jones R, Krishnan A, Robertson AG, Sjostedt S, Doherty C, Cummings SP, Anderson A, Forrest T, Griffin SM, Brodmile M, Pearson J, Ward C. 2016. Microbiological profiles of sputum and gastric juice aspirates in cystic fibrosis patients. Sci Rep 6:26985. https://doi.org/10.1038/srep26985.
23. Venkataraman A, Basisz CM, Beck JM, Young VB, Curtis JL, Huffnagle GB, Schmidt TM. 2015. Application of a neutral community model to assess structuring of the human lung microbiome. mBio 6:e00834-14. https://doi.org/10.1128/mBio.00834-14.
24. Eisenhofer R, Minich JJ, Cooper A, Knight R, Weyrich LS. 2019. Contamination in low microbial biomass microbiome studies: issues and recommendations. Trends Microbiol 27:105–117. https://doi.org/10.1016/j.tim.2018.11.003.
25. Davis NM, Proctor DM, Holmes SP, Relman DA, Callahan BJ. 2018. Simple 16S rRNA gene sequencing reveals site-specific signatures of the upper and lower airways of cystic fibrosis patients. J Cyst Fibros 17:204–212. https://doi.org/10.1016/j.jcf.2017.08.007.
26. Goddard AF, Staudinger BJ, Dowd SE, Joshi-Datar A, Wolcott RD, Aitken ML, Fligner CL, Singh PK. 2012. Direct sampling of cystic fibrosis lungs indicates that DNA-based analyses of upper-airway specimens can misrepresent lung microbiota. Proc Natl Acad Sci U S A 109:13769–13774. https://doi.org/10.1073/pnas.1207453109.
27. Carmody LA, Caverley LJ, Foster BK, Rogers MAM, Kallinik LM, Simon RH, VanDeVanter DR, LiPuma JJ. 2018. Fluctuations in airway bacterial communities associated with clinical states and disease stages in cystic fibrosis. PLoS One 13:e0194060. https://doi.org/10.1371/journal.pone.0194060.
28. Brown PS, Pope CE, Marsh RL, Qin X, McNamara S, Gibson R, Burns JL, Deutch G, Hoffman LR. 2014. Direct sampling of the lung of a young child with cystic fibrosis reveals diverse microbiota. Ann Am Thorac Soc 11:1049–1055. https://doi.org/10.1513/AnnalsATS.201311-3830C.
29. Streeßmann FA, Rogers GB, Marsh P, Lilley AK, Daniels TW, Carroll MP, Hoffman LR, Jones G, Allen CE, Patel N, Forbes B, Tuck A, Bruce KD. 2011. Does bacterial density in cystic fibrosis sputum increase prior to pulmonary exacerbation? J Cyst Fibros 10:357–365. https://doi.org/10.1016/j.jcf.2011.05.002.
30. Price KE, Hampton TH, Gifford AH, Dolben EL, Hogan DA, Morrison HG, Sogin ML, O’Toole GA. 2013. Unique microbial communities persist in
2015. Fecal microbiota transplantation eliminates *Clostridium difficile* in a murine model of relapsing disease. Infect Immun 83:3838–3846. https://doi.org/10.1128/IAI.00459-15.

38. Kozich JJ, Westcott SL, Baxter NT, Highlander SK, Schloss PD. 2013. Development of a dual-index sequencing strategy and curation pipeline for analyzing amplicon sequence data on the MiSeq Illumina sequencing platform. Appl Environ Microbiol 79:5112–5120. https://doi.org/10.1128/AEM.01043-13.

39. Westcott SL, Schloss PD. 2017. OptiClust, an improved method for assigning amplicon-based sequence data to operational taxonomic units. mSphere 2:e00073-17. https://doi.org/10.1128/mSphereDirect.00073-17.

40. Jaccard P. 1908. Nouvelles recherches sur la distribution florale. Bull Soc Vaudoise Sci Nat 44:223–270.

41. Yue JC, Murray KC. 2005. A similarity measure based on species proportions. Commun Stat Theory Methods 34:2123–2131. https://doi.org/10.1080/STA-200066418.

42. Schloss PD, Gevers D, Westcott SL. 2011. Reducing the effects of PCR amplification and sequencing artifacts on 16S rRNA-based studies. PLoS One 6:e27310. https://doi.org/10.1371/journal.pone.0027310.

43. Bates D, Maechler M, Bolker B, Walker S. 2015. Fitting linear mixed-effects models using lme4. J Stat Softw 67:1–48.

44. Kuznetsova A, Brockhoff PB, Christensen R. 2017. lmerTest package: tests in linear mixed effects models. J Stat Softw 82:1–26.