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Characterization and quantification of the fungal microbiome in serial samples from individuals with cystic fibrosis

Sven D Willger¹, Sharon L Grim^{2,4}, Emily L Dolben¹, Anna Shipunova², Thomas H Hampton¹, Hilary G Morrison², Laura M Filkins¹, George A O'Toole¹, Lisa A Moulton³, Alix Ashare³, Mitchell L Sogin² and Deborah A Hogan^{1*}

Abstract

Background: Human-associated microbial communities include fungi, but we understand little about which fungal species are present, their relative and absolute abundances, and how antimicrobial therapy impacts fungal communities. The disease cystic fibrosis (CF) often involves chronic airway colonization by bacteria and fungi, and these infections cause irreversible lung damage. Fungi are detected more frequently in CF sputum samples upon initiation of antimicrobial therapy, and several studies have implicated the detection of fungi in sputum with worse outcomes. Thus, a more complete understanding of fungi in CF is required.

Results: We characterized the fungi and bacteria in expectorated sputa from six CF subjects. Samples were collected upon admission for systemic antibacterial therapy and upon the completion of treatment and analyzed using a pyrosequencing-based analysis of fungal internal transcribed spacer 1 (ITS1) and bacterial 16S rDNA sequences. A mixture of *Candida* species and *Malassezia* dominated the mycobiome in all samples (74%–99% of fungal reads). There was not a striking trend correlating fungal and bacterial richness, and richness showed a decline after antibiotic therapy particularly for the bacteria. The fungal communities within a sputum sample resembled other samples from that subject despite the aggressive antibacterial therapy. Quantitative PCR analysis of fungal 18S rDNA sequences to assess fungal burden showed variation in fungal density in sputum before and after antibacterial therapy but no consistent directional trend. Analysis of *Candida* ITS1 sequences amplified from sputum or pure culture-derived genomic DNA from individual *Candida* species found little (<0.5%) or no variation in ITS1 sequences within or between strains, thereby validating this locus for the purpose of *Candida* species identification. We also report the enhancement of the publically available Visualization and Analysis of Microbial Population Structures (VAMPS) tool for the analysis of fungal communities in clinical samples.

Conclusions: Fungi are present in CF respiratory sputum. In CF, the use of intravenous antibiotic therapy often does not profoundly impact bacterial community structure, and we observed a similar stability in fungal species composition. Further studies are required to predict the effects of antibacterials on fungal burden in CF and fungal community stability in non-CF populations.

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Background

Subjects with the genetic disease cystic fibrosis (CF) often have complex, chronic mixed bacterial-fungal lung infections that cause severe lung damage that can progress to respiratory failure. Molecular profiling studies strongly support a model in which CF lung infections contain multiple bacteria [1-3], and recent work has revealed that these communities are often surprisingly stable within subjects over time [2,4-7]. Fungal species, such as Candida albicans and Aspergillus fumigatus, are also commonly detected in sputum cultures. Culturebased studies have detected C. albicans in ~40%-75% CF subjects [8-21] and described its association with reduced lung function [16,22]. A large European study found that the presence of fungi by laboratory culture correlated with a 5%-10% lower lung function in CF [20]. Two prospective studies have linked the detection of either C. albicans or A. fumigatus to lower lung function and increased frequency of disease exacerbations [16,22]. Other fungi have also been detected in the CF lung [19,23-27]. Challenges associated with the culturing of fungi, due to slow growth, specific medium requirements [28], and the lack of quantitative methods due in part to filamentous morphologies [29], have made it difficult to describe fungal species and their relative burdens in the CF lung. Consequently, despite data that suggest that the mycobiome poses a health threat to individuals with CF [16,22], there is not yet a comprehensive standard of care upon the detection of fungi in sputum.

Individuals with CF-associated lung infections usually experience periods of stable lung function interspersed with episodes of disease exacerbation, which is loosely defined as a period of impaired lung function or the worsening of other health metrics that requires hospitalization for the administration of one or more intravenous antibiotics. The effects of antibiotic treatment on bacterial numbers and community structure are not fully understood and variable across subjects. One recent study found that antibiotic therapy does not cause reproducible shifts in bacterial community structure or bacterial density within sputum [30], while another study reported that antibiotic use was associated with decreased diversity and short-term shifts in community structure [4]. The effects of antibiotic therapy on fungal loads or fungal species composition in the lung are not well known. Several studies have indicated that antibiotic usage precedes the increased detection of fungi in sputum [31-36]. Furthermore, antibiotic therapy in other systems correlates with an increased incidence of fungal infections, such as bloodstream infections [37-40], oral candidosis [41], and vaginal candidiasis [42], and increased fungal loads in the gastrointestinal tract [43-45]. A full characterization of the fungal species within sputum and other host-associated microbial communities will require the community-wide implementation of lysis methods that are effective for all bacterial and fungal species and the use of a combination of molecular methods that provide information on all of the species present. Most molecular analyses of host-associated fungi have relied upon amplification and sequencing of the internal transcribed spacer 1 (ITS1) region within the ribosomal RNA operon [46-49]. Due to a high degree of variation between even closely related species [50-52], this region can serve as a unique taxonomic identifier in both traditional capillary and next generation DNA sequencing studies [46-49].

In this report, we assessed the fungi within the CF lung in subjects hospitalized for treatment with antibacterial antibiotics and advanced the development of tools for the analysis of the fungal component of the microbiome (mycobiome) within clinical samples. Using primers that amplify the ITS1 region, we characterized the mycobiome and bacterial microbiome in expectorated sputum samples from six subjects before and after antibiotic therapy. Candida species dominated the mycobiomes, and their relative abundances were stable over time in contrast to less stable bacterial community structures. Using these samples, we also quantified the numbers of fungal genomes within each sample and found variability in how fungal densities in sputum changed over time and after treatment with antibacterial drugs. Lastly, towards facilitating future mycobiome studies, we assess intra-strain variability in Candida ITS1 sequences and describe improvement of the fungal ITS1 reference database associated with VAMPS (Visualization and Analysis of Microbial Population Structures) [53] and its use in analyses of fungal ITS1 sequences amplified from host-associated microbial communities.

Results

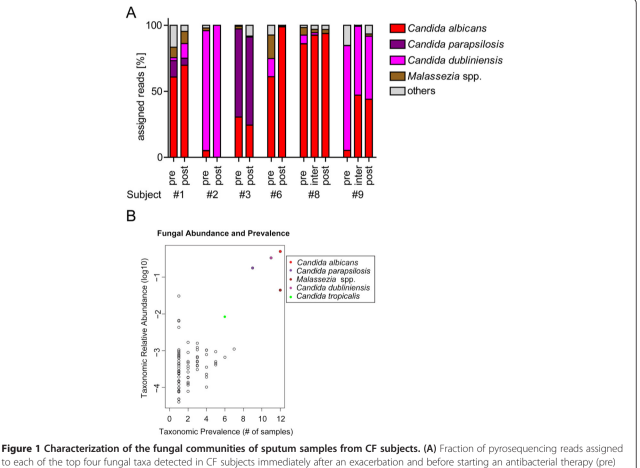
Adapting a publicly available tool for mycobiome analysis To aid the analysis of fungal ITS1 sequences, particularly those amplified from clinical samples, we improved the ITS reference database for Global Alignment for Sequence Taxonomy (GAST) [54] analyses within VAMPS, a widely used web-based tool [53] that resolves phylogenetic affinity of marker gene sequences from diverse bacterial communities ([2,55-57] as examples). The reference database for assigning the taxonomic affinity of fungal ITS1 amplicon sequences derives from the UNITE database [58]. Our modifications included a number of corrections to the taxonomic information within records for medically relevant fungi, and the removal of some sequences that were exact matches to well-characterized plant sequences that had been incorrectly annotated as being of fungal origin. We also increased representation of the humanassociated fungi *Candida* and *Malassezia* in the database by adding all Genbank database entries that contained the keywords "ITS", "Candida", and "Malassezia" and had complete annotations. This step increased the number of *Candida* species from 251 to 284 and the number of unique accession IDs containing the keyword "*Candida*" from 930 to 2021. For *Malassezia*, 72 unique accession IDs containing the keyword "*Malassezia*" from Genbank with complete genus and species information were added.

Overall assessment of the fungi identified in sputum from inpatient subjects

Using VAMPS, we analyzed the fungal community in CF sputum from six hospitalized subjects that received subject-specific antibacterial treatments (Additional file 1: Table S1). Using primers that match conserved flanking sequences, we PCR-amplified the fungal ITS1 sequences from sputum DNA samples for pyrosequencing analyses. We determined the reproducibility of the fungal ITS1 amplification and deep sequencing protocol by analyzing technical replicate libraries created from the same DNA

that had been extracted from a de-identified sputum sample collected for the purposes of the method development. The results from the two technical replicates were very similar in terms of the fraction of reads assigned to each fungus (<1% difference in percent abundance per sample) (Additional file 2: Figure S1).

Datasets from each sputum sample contained an average of 21,434 fungal reads ranging from 1,672 fungal reads (the inter-treatment sample of subject #9) and 37,684 (the post-treatment sample of subject #6) (Additional file 3: Table S3), and a rarefaction analysis indicated that the sequencing depth was sufficient to detect the vast majority of fungi present (Additional file 4: Figure S2A). *Candida* represented the dominant genus with 11 different species. The top three *Candida* species (*C. albicans, Candida dubliniensis,* and *Candida parapsilosis*) accounted for 74%–99.9% of all ITS1 reads in all samples (Figure 1A) making these species both the



to each of the top four fungal taxa detected in CF subjects immediately after an exacerbation and before starting an antibacterial therapy (pre) and approximately 2 weeks afterwards (post) or while hospitalized (inter). The legend indicates the color assigned to each indicated fungal species or genus (*Malassezia* sp. unknown). (B) Relative abundance of all fungal genera, extended to the species level for the clinically important *Candida* species (colored dots), is plotted against the prevalence of each genus and *Candida* species in the six CF subject samples (pre- and post-treatment) and show that fungi that are highly abundant in a single patient are also highly prevalent across patients. The colored dots indicate the *Candida* species and the genus *Malassezia* that are both highly abundant and highly prevalent and the empty dots represent all remaining fungal genera identified in our study.

most abundant species and also the most prevalent. C. albicans ranged from 0.1% of reads (the post-treatment sample from subject #2) to 98.8% of reads (the posttreatment sample from subject #6) (Additional file 5: Table S4). In three subjects, the only reported fungi using culture-based detection methods were C. parapsilosis in subjects #1 and #3 and C. dubliniensis in subject #2 (Additional file 1: Table S1). Another clinically interesting Candida species, Candida tropicalis, was detected at lower levels, ranging from 2 to 477 reads, and only in half of the samples (Figure 1B). Aside from C. albicans, only Malassezia spp. (including Malassezia restricta and Malassezia globosa) occurred in all samples, but at levels 10- to 50-fold lower than the Candida spp. reads (Figure 1A,B). We detected low numbers of reads corresponding to three different species of Aspergillus (Aspergillus unguis, Aspergillus versicolor, and Aspergillus sp.) in this study, but not A. fumigatus, another common CF pathogen, which we have previously detected in other CF sputum samples (data not shown). Across all 14 samples, our VAMPSbased analyses detected 111 different taxa at the genus level, and these resolved to 147 species level assignments (Additional file 5: Table S4). The minor members of the mycobiome are discussed in more detail below.

To validate the results obtained by GAST analyses, we employed BLAST to assign fungal ITS1 sequences to species and genus level categories (Additional file 6: Table S5). For the BLAST searches, we batch blasted examples of all unique reads and assigned a genus or species to each set of sequences. Comparison of GAST and BLAST results initially highlighted the need for database improvement (described above), and once the database had been modified, the two methods yielded a similar number of taxa. Because GAST reports the consensus annotation for all of the sequences that match the query, the developmental state of the ITS1 database may limit the assignment of genus or species level information. In this case, sequences clustered by GAST in VAMPS can be easily retrieved for further analysis.

Because the characterization of fungi within hostassociated microbial communities is at an early stage, we sought to determine if there were any ITS1 sequences that have not been previously described. In initial analyses, up to 13.3% of sequences in a sample could not be identified as fungi (BLAST scores of greater than e - 15for the top hit). On average, samples had 3.6% of reads in this category. Further analysis querying Genbank's Nucleotide collection (nr/nt) database revealed that most of these unassigned sequences corresponded to ITS1 sequences in the genomes of food-associated plants such as *Pisum sativum* (peas), *Allium cepa* (onions), *Allium sativum* (garlic), *Solanum tuberosum* (potato), and *Glycine max* (soybean) suggesting contamination of the sputum sample from subjects' diets. The primers used to amplify fungal ITS1 sequences (Additional file 7: Table S2) displayed high levels of similarity to comparable regions in plants. VAMPS allows one to limit analyses to specific taxonomic levels or to specific groups (for instance, fungi or Ascomycetes) [53], and this eliminated the problem of contaminating sequences. A very small number of unassignable sequences (13 out of 300,072 reads) were also identified, but they were most often single reads that were only found within single samples and were not characterized further.

Comparison of intra- and inter-species ITS1 sequence differences in *Candida* spp.

Some fungal genomes may contain up to 600 copies of the ribosomal RNA genes (rDNA) and intraspecific ITS variability can exceed 24.2% [59-62]. In contrast, C. albicans typically has 150-200 copies of the ITS1 locus, with one report estimating less than 0.2% ITS variability within a strain [62]. Since Candida represented the predominant fungal genus in our CF subjects, we used Illumina MiSeq sequencing to investigate levels of ITS1 sequence variation within and among clinical C. albicans isolates. When the ITS1 region was amplified from genomic DNA from pure cultures of C. albicans wild type SC5314 and seven clinical isolates, then analyzed using Illumina MiSeq, the predominant sequence (>99% of reads) for each strain was the same (100% identical across all isolates). The second most abundant sequence accounted for only ~0.02% of reads. Similarly, we observed no evidence for intra-strain variation in the ITS1 sequences from the analysis of genomic DNAs isolated from pure cultures of three other Candida species (C. dubliniensis, C. parapsilosis, and C. kruseii) (data not shown).

When we compared the most abundant *C. albicans* ITS1 sequence in the sputum-derived datasets to the dominant sequence from the clinical isolates, we found that they were 100% identical (data not shown). Alignments of the three most common ITS1 sequences that resolved to *C. albicans* from each sample, with the ITS1 sequences from the genome information for *C. albicans* strains SC5314 and WO-1, displayed 99% identity. The second most abundant *C. albicans* ITS1 454 sequence amplified from sputum DNA differed from the most abundant ITS1 sequence by one missing "T" at position 60, which most likely reflects a 454 homopolymer sequencing error. By way of comparison, the closely related *Candida* species *C. albicans* and *C. dubliniensis* have ITS1 sequences that have 94.6% identity.

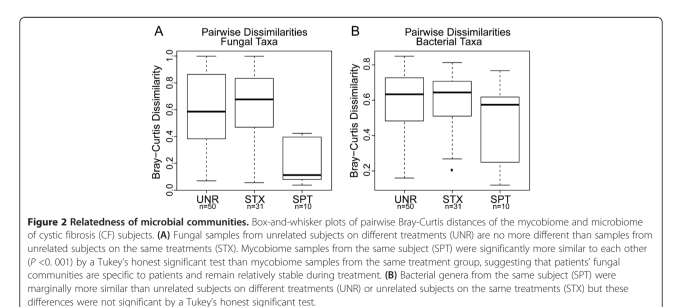
Description of the mycobiome and bacterial microbiome across serial sputum samples

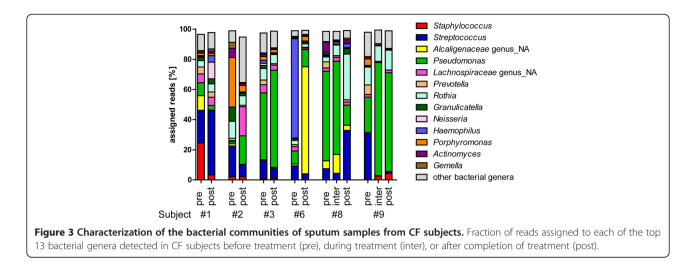
In the collection of serial samples, each subject provided a sputum sample at the time of admission for treatment of exacerbated respiratory disease using two or three antibacterials and one sample upon the completion of treatment; some subjects also provided a third sample during treatment. The drugs administered included Tobramycin, Vancomycin, Doripenem, Meropenem, Ceftazadime, and Linezolid and information which antibacterial drugs were used in each subject is provided in Additional file 1: Table S1. The fungal communities in all samples, grouped by subject, are shown in Figure 1A. The relative abundance of reads corresponding to the predominant Candida spp. appeared to be quite stable over time during treatment, and there was no obvious change upon treatment initiation (Figure 1A). Bray-Curtis dissimilarity and principal coordinate analysis were used to measure and represent taxonomic relatedness between classes of samples in the mycobiome and the microbiome samples. Antibiotic treatment did not establish a specific fungal community. Fungal samples from unrelated people at different points in therapy (unrelated, UNR) were no more different than samples from unrelated people at the same point in treatment (same treatment, STX). However, mycobiome samples from the same subject (same subject, SPT) were significantly more similar to each other (P < 0.001) by a Tukey's honest significant test (Figure 2A) than mycobiome samples from the same treatment point, suggesting that subjects' fungal communities remained relatively stable. Furthermore, as shown in the PCA scores plot (Additional file 8: Figure S3), the samples from subjects #1, #6, and #8 clustered together as did those from subjects #2 and #9. Subject #3 was distinct from the other samples (Additional file 8: Figure S3).

To compare the fungal community profiles to those for the bacteria, we amplified and sequenced the V6 region of the bacterial 16S rDNA gene from the same sputum DNA samples. On average, we recovered 726,787 sequences from each sample (Additional file 3: Table S3), and a rarefaction analysis indicated that this deep sequencing effort identified most members of the sampled bacterial microbiomes (Additional file 4: Figure S2B). Ninety-nine percent of all reads from all samples were assigned to 44 genera, though not all 44 genera were detected in every sample (Additional file 9: Table S6). The remaining 1% of reads corresponded to 397 different genera. As reported by other studies, most samples contained high levels of the genera *Pseudomonas, Staphylococcus, Streptococcus, Prevotella, Rothia, Granulicatella, Neisseria, Haemophilus, Porphyromonas, Gemella,* and members of the families *Alcaligenaceae* and *Lachnospiraceae* (Figure 3) [2,4,63,64].

Consistent with published work [4,6,65], the bacterial community also displayed stability in the major community members in many subjects (Figure 3), though some changes in the bacteria with the highest relative abundance were observed in subjects #2 and #6 (Figure 3). The Bray-Curtis dissimilarity of the bacterial communities showed a less clear picture than for the fungi, but it still was apparent that the samples from the same subject showed some degree of relatedness. Bacterial community profiles from the same subject were marginally more similar (SPT) but the difference was not significant by a Tukey's honest significant test (Figure 2B); like for the fungi, the STX samples were not more similar than the UNR samples. There was less of a clear trend in sample clustering in the PCA scores plot, and the larger number of bacterial genera in each sample may have contributed to this complexity (Additional file 8: Figure S3).

To compare bacterial and fungal richness across samples, we first accounted for differences in the number of

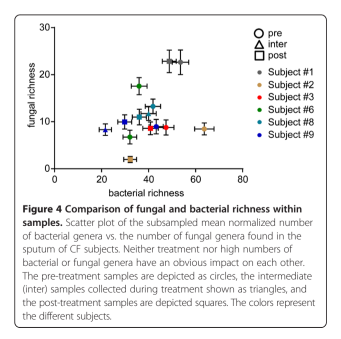




reads by subsampling 1,000 reads for each fungal and bacterial sample and repeatedly subsampling 1,000 times. First, we found that the normalized mean number of fungal genera did not correlate with the number of normalized mean bacterial genera (Figure 4). Using a Wilcoxon matched-pairs signed ranked test, it was found that the post-treatment samples had a lower mean bacterial genus richness (P < 0.05) when compared to pre-treatment samples (Additional file 10: Figure S4). While the normalized mean richness for the fungi trended in the same direction, the differences were not statistically significant (P > 0.1).

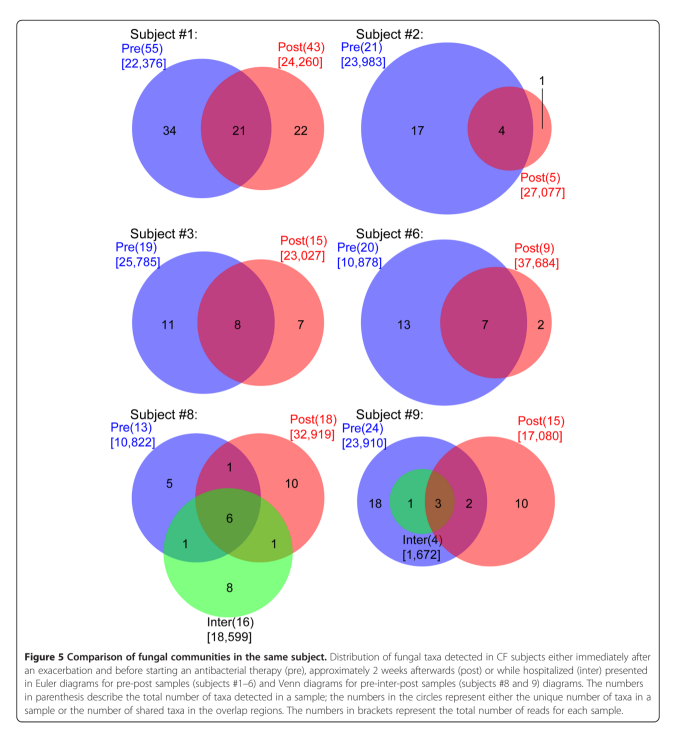
Analysis of minor members of the mycobiome

The remaining fungal genera represented relatively low abundance populations that occurred in one or more



samples (Figure 1B and Additional file 5: Table S4 for identities). When using the BLAST-based mycobiome analysis (see Additional file 11), the number of taxa identified within a sample ranged from 5 taxa (resolved to 3 species, 1 genus, and 1 cluster of unidentified sequences in the post-treatment sample from subject #2) to 60 taxa (resolved to 43 species, 16 genera, and 1 cluster of unidentified sequences in the pre-treatment sample of subject #1) (Additional file 12: Table S5). Minor members (<0.05% of annotated ITS1 sequences) of the mycobiome included other *Candida* species, *Trichosporon* (an emerging pathogen in CF subjects [27,66]), and the human pathogens *Rhodotorula*, *Fusarium*, and *Penicillium* (Additional file 5: Table S4).

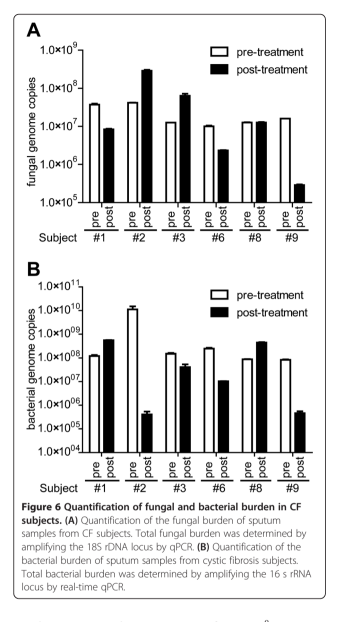
The deep sequencing in this study and the corresponding rarefaction analyses suggest that we have identified the number of taxa that were common to the two (and in some cases three) samples from a given subject (Additional file 4: Figure S2). The overlap in fungal taxa in samples from the same subject is presented as Euler and Venn diagrams (Figure 5 based on data in Additional file 5: Table S4). When only the pre- and post-treatment samples were compared, an average of 44% ±19% of the fungal species detected was common in both samples. This highlights the fact that some of the minor fungal taxa were not common across the serial samples and may represent transient species in the airway or oral community. As an example, 21 taxa were detected in both the pre-treatment and posttreatment samples from subject #1; 34 taxa were unique to the pre-treatment sample and 22 taxa were only in the post-treatment sample (Figure 5). As sequencing depth can impact the number taxa per sample and thus the apparent stability of a taxon, we also considered and presented the number of reads obtained for each sample; there were no differences in sequencing depth between the pre- and post-treatment samples from a given subject (Figure 5). The stability of species across



samples from the same subject was also assessed for bacteria (Additional file 12: Figure S5). Again, comparing the pre- and post-treatment samples, we found that an average of $77\% \pm 7\%$ genera were in common in the two samples (Additional file 12: Figure S5 based on data in Additional file 9: Table S6). Additional file 13: Table S7 highlights those fungal taxa, including low abundance taxa that were detected in more than one sample.

Analysis of the changes in community structure and absolute abundance before and after antibacterial therapy

To complement the data on the relative abundances pre- and post-treatment, we also analyzed the density of bacteria and fungi within each sputum sample by qPCR. Results were normalized to sputum dry weight (genome copies per mg dry weight) (Figure 6A). On average, we could detect $\sim 4.2 \times 10^7$ fungal genome copies/mg dry



weight sputum with a maximum of 2.8×10^8 copies in the post-treatment sample of subject #2 and minimum of 2.9×10^5 copies in the post-treatment of subject #9 (Figure 6A). Overall, we did not observe a specific trend in fungal burden corresponding to the antibacterial therapy. In three subjects, the fungal burden dropped (subjects #1, #6, and #9); in two subjects, the fungal density increased (subjects #2 and #3); and subject #8 exhibited no difference. We designed primers to quantify levels of the four Candida species (Additional file 7: Table S2), and the primers were shown to be species specific with respect to one another in studies performed using genomic DNA from purified cultures (data not shown). When the absolute abundances of fungi, determined using 18S rDNA quantification was compared to the sum of the levels found using the Candida species specific primer sets, they were within the same order of magnitude (Additional file 14: Figure S6) confirming that the *Candida* species were numerically dominant among the fungi within these samples. The qPCR approach also confirmed that relative abundance of *C. tropicalis* levels was much lower than *C. albicans* (data not shown). This finding may parallel observations that *C. tropicalis* is often less abundant than the other *Candida* species in oral samples [67-70].

We determined total bacterial burden without preamplification by targeting the 16S rRNA locus [2] (Figure 6B). On average, we detected $\sim 9.2 \times 10^8$ bacterial genome copies/mg dry weight sputum with a maximum of 1.1×10^{10} copies in the pre-treatment sample of subject #2 and minimum of 4.0×10^5 copies in the post-treatment of subject #2. As with the fungal quantification, we did not observe a specific trend in bacterial density in the pre- and post-treatment samples. The bacterial sputum density dropped in four of the patients, most significantly in subject #2 in which the bacterial burden within sputum dropped ~27,500fold to a level close to the detection limit. In two subjects, the bacterial level went up approximately five-fold. There was no correlation between the changes in bacterial density and changes in fungal density (Figure 6).

Discussion

Dominant members of fungal communities, which we defined as those that account for more than 80% of all assigned reads per sample (Candida and Malassezia species) (Figure 1A) persisted throughout treatment without significant change in relative abundance suggesting that there is stability within the mycobiome. Many minor members of the fungal community appeared to be ephemeral or sporadic, and thus did not show changes in prevalence or abundance that could be attributed to antibacterial treatment (Figure 5 and Additional file 13: Table S7). Based on the depth of sequencing and the rarefaction analyses, the apparent instability of minor members is unlikely an artifact of sequencing depth, but rather likely a reflection of their transient nature. The fungal species we detected within CF sputum mirrors results in other recently published work. For example, denaturing high-performance liquid chromatography analysis of the ITS1 region amplified from fungi within CF sputum also found C. albicans, M. restricta and M. globosa [71]. In a study of four stable CF subjects by Delhaes et al. [19], these fungi were also well represented. While C. albicans is commonly observed in CF sputum, the presence of Malassezia is less well understood in part because Malassezia cannot be cultured without the use of specialized medium. Investigation of the fungal communities on the human skin found M. restricta and M. globosa to

be very common [72], and future work will be required to determine if Malassezia are living in the lung or if the signal is coming from dead cells or DNA. Mournier et al. [71] also detected more fungi using molecular methods when compared to culture-based analyses, and Candida and Malassezia were among them. In addition, they found that A. fumigatus was detected by culture in some samples but not by molecular techniques [71]. While A. fumigatus was not seen in clinical microbiological analyses of the sputa in this study, we were surprised by its complete absence from our datasets. We have detected A. fumigatus in other samples indicating that our methods can amplify its ITS1 region. A recent shotgun metagenomic analysis of CF sputum DNA also found significant levels of C. albicans and A. fumigatus by culture but found no DNA evidence for A. fumigatus and a small number of other Candida sequences [73]. In contrast to what was observed in CF sputum, Charlson et al. [74] analyzed the fungi in the lavages of healthy individuals, and they found that healthy volunteers' bronchoalveolar lavage (BAL) yielded scant fungal amplification. Although *Candida* was identified in several healthy subjects' oropharyngeal washes, it was absent in their BAL.

In this study, we explored whether inpatient treatment, which included multiple antibacterials, impacted the fungal community in a consistent way, perhaps indicating changes in biological niches or environmental conditions upon perturbation of the bacterial community. Our data suggest that the use of multiple antibiotics did not necessarily have strong effects on either the bacterial or fungal communities. While the number of bacterial genera (richness) declined slightly but significantly (Additional file 10: Figure S4), there was not a consistently change to the structure of the community with similar relative abundances of dominant species in four out of six subjects (Figure 2 and Additional file 12: Figure S5). In these same samples, we did not observe a consistent increase in the number of fungal genera (Additional file 10: Figure S4) and the fungal species that dominated the samples did not change markedly (Figure 1A). This lack of difference between pre- and post-samples may well be due to the fact that these subjects have been exposed to years of inhaled and intravenous antibacterial therapy, and the bacteria in their lungs have adapted or evolved in this context. Thus, future studies in subjects without routine exposure to antibiotics are needed in order to assess the effects of antibacterials on the resident fungi. We and others have characterized the interactions between Pseudomonas aeruginosa and C. albicans, including effects on fungal morphology [75,76], viability [77,78], metabolism [76,79], and virulence factor regulation [80,81], and the high relative abundance of *P. aeruginosa* in the bacterial communities and C. albicans in the fungal communities supports the hypothesis that these species have the potential to impact one another in the lung.

The therapy used in the treatment of disease exacerbation also had no consistent impact on fungal density in patient sputum samples. Measurements of fungal and bacterial genomes per milligram of sputum showed an inconsistent trend across serial samples collected before and after antibacterial treatment (Figure 6) suggesting the need for either a larger number of subjects or alternative methods for the measurement of live microbes including bacteria and fungi. Previous reports in other body sites have linked antibiotics to increased fungal burden [82-88]. In a recent study, Dolville et al. [89] investigated the gut microbiome of mice over 76 days of treatment with the antibiotics vancomycin, ampicillin, neomycin, and metronidazole and subsequent recovery. They observed that bacterial numbers dropped in abundance more than 3-fold while fungi increased ~40-fold in abundance upon treatment but found that both numbers went back to pre-treatment levels upon recovery. Interestingly, the bacterial microbiome was detectably different from Candida-free controls and Candida became more abundant than at the time point prior to treatment [89]. While we observed that fungal density in sputum had the potential to vary (either increase or decrease), we found no common trend. It is important to note that all of these patients have a significant history of prior antibiotic exposure and thus some adaptation may have occurred. The effects of antibiotic therapy in fungal populations may differ from those obtained upon first exposure to a drug.

Fungal load has been associated with lung disease severity [16,20,22] suggesting that it is important to measure fungal density in sputum in addition to the mere presence of specific species. We found a range of fungal loads across samples (Figure 6), and this range is consistent with previous reports. For example, Bauernfeind et al. [15] found between 10^4 and 10^7 colony forming units per gram of sputum of Candida using culture methods. Charlson et al. [74] highlight the importance of measuring bacterial and fungal loads and to compare it to background contamination by upper respiratory tract flora to identify lung-enriched taxa. Here, we applied qPCR quantification of the 16S rDNA locus for total bacteria, the 18S rDNA locus for total fungi, and species-specific primers to quantify three Candida species. One limitation of DNA-based quantification protocols is the inability to differentiate between nucleic acids from live and dead species. Methods to inactivate DNA from dead or lysed cells would enhance our understanding of the active microbiota in the lung [90].

Finally, we report the use of VAMPS for analysis of the ITS1 sequencing results. VAMPS, which has provided a user-friendly tool for the analysis of bacterial communities [2,91], now has an improved database that will expand the ability to analyze fungal ITS1 sequences from clinical samples. The VAMPS database, derived from the UNITE database [58], contains most of the available ITS1 sequences. However, further curation will improve taxonomic information in existing entries, and the inclusion of additional high-quality, taxonomically resolved sequences that further aid in the analysis and taxon assignment in future human mycobiome studies [53].

Conclusions

The data presented in this report found stable fungal communities in the sputum of six individuals with CF. Though to a lesser extent, stability was also mirrored in the bacterial communities. We hypothesize that the resident bacteria have developed a tolerance or resistance to antimicrobials due to repeated prior exposure, and thus, little perturbation of co-colonizing fungi is observed. In addition to numerically dominant fungal populations that are predominantly members of the genus Candida, there are less abundant fungi, many of which were less stable across serial samples from the same subject. To enhance future analyses of the mycobiome, we developed and modified a fungal ITS1 database for use in VAMPS, a publically available tool for the analyses of visualization of data on microbial communities. Future studies will continue to investigate the effects of antibacterial drugs on fungal and bacterial organisms and their densities in sputum.

Methods

Study participants and ethics statement

This study was performed in accordance with the guidelines set by the Committee for the Protection of Human Subjects (CPHS) and approved by the Institutional Review Board (IRB) at Dartmouth College. Sputum samples were obtained from six subjects over the age of 18 with a confirmed diagnosis of CF and who had previous cultures positive for *P. aeruginosa* after written, informed consent. Information about the study participants is provided in Additional file 1: Table S1.

Sample collection

Serial sputum samples were collected from six CF subjects at the time of hospitalization for treatment of a respiratory disease exacerbation and after a complete course of intravenous antibacterial therapy, which often included the use of multiple antibiotics (Additional file 1: Table S1). For some subjects, an intermediate sample during therapy was also collected. Study participants expectorated sputum into a sterile specimen cup, and the sputum sample was stored at -80° C. In preparation for DNA extraction, the sample was thawed and homogenized by passing the sputum multiple times through needles of

increasing gauges. Sputum (500 μ l) was dispensed into a 2-ml screw cap tube and frozen at -80° C for at least 1 h prior to lyophilization for at least 12 h. The dry weight of the sputum was determined before DNA extraction.

DNA extraction

Cells within the dried sputum sample were disrupted using a mixture sterile glass beads (equivalent amounts of 0.1-, 0.5-, and 1-mm diameter beads) in a bead beater (Biospec, Bartlesville, OK, USA); the tubes were agitated four times for 30 s each time with a cooling step in between pulses. The ground samples were resuspended in 300 µl of TE + DTT (TE amended with DTT at a final concentration of 0.08% added from a 2% stock solution) containing lysozyme (3 mg/ml) and lyticase (10 U/ml) and incubated for 30 min at 37°C. Cell lysis buffer (500 µl) (Qiagen Puregene Core Kit B, QIAGEN Inc., Valencia, CA, USA) was added, and the mixture was incubated for 15 min at 80°C. To remove RNA, RNase $(1.5 \ \mu l)$ (QIAGEN Inc.) was added and the samples were incubated for 30 min at 37°C. Lysates were chilled on ice for 1 min, 200 µl of Protein Precipitation Solution (Qiagen Puregene Core Kit B, QIAGEN Inc.) was added, and the solutions were mixed vigorously for 20 s. Cell debris was sedimented by centrifugation at 13,000 rpm for 3 min, and the supernatant was transferred to a new 1.5-ml tube prior to addition of 600 μ l of 100% isopropanol. After mixing by inversion, the DNA was precipitated by centrifugation at 13,000 rpm for 20 min. The DNA pellet was washed with 300 µl of 70% ethanol and air dried before resuspension in 100-200 µl of DNA hydration solution (Qiagen Puregene Core Kit B, QIAGEN Inc.). The DNA concentrations were measured using a Nanodrop.

Fungal ITS1 amplification and sequencing

The ITS1 region was amplified from sputum template DNA using the primers ITS1F and ITS1R (Additional file 7: Table S2) in two rounds of PCR. The reaction components were 1× Qiagen PCR buffer, 0.75 mM MgCl₂, 7% (ν/ν) dimethyl sulfoxide (DMSO), 0.2 mM deoxynucleotide triphosphates (dNTPs), 0.4 µM primers, 1.25 units Tag polymerase (QIAGEN Inc.), 100 ng of template DNA, and water in a 25-µl reaction volume. Cycling conditions were 95°C 5 min; 95°C 30 s, 50°C 30 s, 72°C 45 s for 10 cycles; and 72°C 2 min. PCR products were purified using the MinElute PCR Purification Kit (QIAGEN Inc.) and resuspended in 10 µl of EB buffer. The entire 10 μ l volume was used as template for a second round of PCR using primers containing adapter and barcode sequences for Roche GS-FLX amplicon pyrosequencing (Additional file 2: Table S2). Reaction components and cycling conditions were as described above, but with 30 cycles. Amplified DNA was purified

with 0.7 volumes of Agencourt AMPure XP beads (Beckman Coulter, Brea, CA, USA) and quantified on a Bioanalyzer 2100 (Agilent, Palo Alto, CA, USA). Purified amplicon libraries were sequenced using the Titanium amplicon protocol on a GS-FLX (Roche Applied Science, Indianapolis, IN, USA) following the methods outlined in [2]. Sequence data were quality filtered as described in [54,92]. We identified and removed chimeric reads by querying against the curated reference ITS1 database, as well as using the sequences as seeds in UCHIME de novo [93]. The raw sequencing results of the ITS1 analysis are deposited in the Sequence Read Archive (NCBI Bio-Project PRJNA246028; National Center for Biotechnology Information (NCBI), US National Library of Medicine, Bethesda, MD, USA) and are summarized in Additional file 5: Table S4.

For the MiSeq analysis of the ITS1 region amplified from clinical isolates of *Candida* spp., template DNA was isolated using the Epicentre Masterpure Yeast DNA Purification Kit. Primers are listed in Additional file 7: Table S2. The amplification and sequencing methods used are comparable to those reported [94].

Bacterial 16S v6 amplification and sequencing

The bacterial communities from the pre- and postantibacterial treated CF subjects were characterized by Illumina HiSeq 16S rRNA V6 amplicon sequencing (Illumina Inc., San Diego, CA, USA) as previously described by Eren et al. [95]. Briefly, we amplified the bacterial V6 region from each sample in triplicate 33 µL reactions, pooled the triplicates, and verified successful amplification on a Caliper LabChip GX (Perkin Elmer, Waltham, MA, USA). We cleaned amplicon products with a QIAQuick clean up assay (QIAGEN Inc.), and used fusion V6 primers to barcode each sample in a multiplexing strategy. Final barcoded products were quantitated with a Picogreen assay (Invitrogen/Life Technologies, Grand Island, NY, USA) and pooled in equimolar concentrations with respect to the target size. All PCR reactions contained 1× HiFi Buffer, 2 mM MgSO₄, 0.02 U/µL Platinum Taq polymerase (Invitrogen), 0.2 mM each dNTPs (ThermoFisher Scientific, Milwaukee, WI, USA), and up to 5 ng template. The final pool was size selected (Pippin Prep, SageScience, Beverly, MA, USA), quantitated (Kapa Biosystems, Woburn, MA, USA) and sequenced on a HiSeq 1000 in a 2×100 nt sequencing run. Quality filtering and chimera checks were performed as described above. The raw sequencing results of the bacterial 16S v6 analysis are deposited in the Sequence Read Archive (NCBI BioProject PRJNA256169; National Center for Biotechnology Information (NCBI), US National Library of Medicine, Bethesda, MD, USA) and are summarized in Additional file 9: Table S6.

Microbiome and mycobiome clustering and taxonomic assignment

The website http://vamps.mbl.edu provides access to individual reads, taxon assignments, and descriptions of individual clusters. To create the databases that are accessed through VAMPS researchers at the Marine Biology Laboratories at Woods Hole merged paired-end 100 bp reads into single consensus reads that completely covered the V6 region, and they assigned taxonomy with GAST [54] against a curated SILVA database [96]. To compare operational taxonomic unit (OTU) clustering performance, they employed the default UCLUST method [97] at a 97% similarity threshold with minimum cluster size of 2 using quantitative insights into microbial ecology (QIIME) (v1.5) [98].

Rarefaction analysis

To verify adequate sequencing depth of the mycobiome and the microbiome, we performed rarefaction analysis of the two datasets. For both rarefaction analyses, we used raw counts of all subject samples generated by VAMPS with the taxonomy depth selector set to "species" to create a read matrix. This table was rarefied using an R-based protocol described on the webpage "Individual Based Rarefaction using R-package" (http:// www.jennajacobs.org/R/rarefaction.html) [99]. The resulting table was transferred into GraphPad Prism 6 to plot the rarefaction curves.

Euler and Venn diagrams

To visualize the impact of antibacterial treatment on the prevalence of fungal and bacterial taxa detected in CF subjects either immediately after an exacerbation and before starting an antibacterial therapy (pre), approximately 2 weeks afterwards (post) or while hospitalized (inter), we used the Additional file 5: Tables S4 and Additional file 9: Table S6 to generate the diagrams. First, we copied the data for pre, post and if applicable inter samples into the list-fields of an online Venn diagram generator (http://www.bioinformatics.lu/venn.php). Using the R package "RVennDiagram" (http://cran.r-project.org/ web/packages/VennDiagram/index.html), we generated Euler ("draw.pairwise.venn") and Venn ("draw.triple.venn") diagrams following the instructions in the manual (http:// cran.r-project.org/web/packages/VennDiagram/VennDiagram.pdf).

Quantification of fungi, Candida species, and bacteria

In order to quantify minor members of the mycobiome, we performed analyses using either sputum DNA or preamplified DNA derived from ten rounds of PCR starting with 100 ng of sputum DNA using primers specific to a conserved region of the 18S rDNA (FungiQuant primers) [100] or *Candida* species-specific primers (Additional file 7: Table S2). We confirmed that the pre-amplification step was linear with respect to the initial sample; we performed multiple control experiments using genomic DNA isolated from different fungal species at concentrations ranging from 30 pg/µl to 300 ng/µl. The preamplification allowed for the analysis of minor members of the mycobiota. The reaction components of the first PCR were 1× Qiagen PCR buffer, 0.75 mM MgCl₂, 7% (ν/ν) DMSO, 0.2 mM dNTPs, 0.4 µM primers, 1.25 units Taq polymerase (QIAGEN Inc.), 100 ng sputum DNA, and water in a 25 µl reaction volume. Cycling conditions were 95°C 5 min; 95°C 30 s, 50°C 30 s, 72°C 45 sec for 10 cycles; and 72°C 2 min. These PCR products were purified using the MinElute PCR Purification Kit (OIAGEN Inc.), and resuspended in 10 µl of EB buffer. Quantitative PCR (qPCR) was conducted in 20 µl reaction volumes with the iO SYBR Green Supermix (Bio-Rad Laboratories, Hercules, CA, USA). Reactions contained 2 µl of the amplicon suspension from the first round PCR and 0.2 μ M of the appropriate primers. To quantify total fungal burden, we used the primers FungiQuant_RT_F and Fungi-Quant_RT_R [99]. The aforementioned species-specific qPCR primers were designed to assess Candida burdens based on gene sequences available from the Candida Genome Database (http://www.candidagenome.org/). Genes used for primer design were CAWG_05066 (C. albicans), CPAR2_301290 (C. parapsilosis), Cd36_16280 (C. dubliniensis) and CTRG_03824 (C. tropicalis) and were selected based on conservation within species and a lack of cross reactivity with bacterial, fungal, and human sequences. qPCR was performed utilizing a CFX96 real-time PCR detection system combined with a C1000 thermal cycler (Bio-Rad Laboratories). All PCRs were done in duplicate, and data were analyzed with the CFX96 System gene expression software. Melt curve analysis was performed after the PCR was complete to confirm the absence of nonspecific amplification products. Standard curves containing a known numbers of genome equivalents were used to calculate genome numbers per mg dry weight.

For total bacterial quantification, 100 ng of sputum DNA was used as the PCR reaction template. To generate a standard curve for quantification purposes we prepared 10-fold dilutions of *P. aeruginosa* genomic DNA isolated from 10^2 to 10^6 cells. The 16S rDNA genes were amplified by qPCR, using the universal bacterial primers "total bacteria_F" and "total bacteria_R" at a final concentration of 0.2 μ M [101,102] (Additional file 7: Table S2). qPCR was conducted in 20 μ l reaction volumes with the iQ SYBR Green Supermix (Bio-Rad Laboratories). qPCRs for each sputum sample were performed and analyzed on the CFX96 PCR system as described above.

Statistical analysis

Bray-Curtis dissimilarity was measured using the ecodist [103] and vegan [104] packages in R statistical software.

Significant differences between groups were established using analysis of variance with Tukey's honest significant difference post-test. Tests producing a P value less than 0.05 were deemed significant. The program Prism 6 (GraphPad, San Diego, CA, USA) was used for remaining statistical tests.

Additional files

Additional file 1: Table S1. Microbiological results from samples collected at the time of admission for treatment of exacerbation and antibacterial drugs administered.

Additional file 2: Figure S1. Analysis of amplification and sequencing method reproducibility. DNA from a sputum sample was divided, amplified, sequenced, and analyzed in parallel to evaluate our methods.

Additional file 3: Table S3. Total number of reads used in the mycobiome (fungi) and the bacterial microbiome analyses.

Additional file 4: Figure S2. Rarefaction analysis of the mycobiome and microbiome. Rarefaction curves to determine the completeness of deep sequencing of the mycobiome (A) and the microbiome (B). Repeated samples of species subsets were used to evaluate whether further sampling would likely yield additional species, as indicated by whether the curve has not yet reached a plateau. The *y*-axis indicates the number of species detected and the *x*-axis the number of sequences analyzed per sample.

Additional file 5: Table S4. Fungal sequences detected in CF sputum samples using assignments made through VAMPS. The data are presented as the percent of total reads and absolute read number.

Additional file 6: Table S5. Fungal species within CF sputum samples using the local BLAST analysis method. The data are normalized to percent of total reads.

Additional file 7: Table S2. Primer list.

Additional file 8: Figure S3. Principal component analysis based on Bray-Curtis dissimilarity. Bray-Curtis dissimilarity and principal coordinate analysis were used to measure and represent taxonomic relatedness between classes of samples in the mycobiome (fungal taxa) and the microbiome (bacterial taxa) samples. Pre-treatment samples are colored in red and are labeled A, intermediate samples collected during treatment are colored blue and are labeled B, and post-treatment samples are colored green and are labeled C. The numbers indicate the subject number.

Additional file 9: Table S6. Bacterial genera identified in CF sputum samples using VAMPS. The data are presented as the percent of total reads and absolute read number. Only the genera that account for 99% of all reads are presented.

Additional file 10: Figure S4. Comparison of the normalized richness of the mycobiome and microbiome. Comparison of the normalized mean fungal and bacterial richness of pre-treatment and post-treatment samples. The normalized mean fungal richness shows no significant difference between treatments (P > 0.1) but the normalized mean bacterial richness decreases upon antibacterial treatment (P < 0.05) (Wilcoxon matched-pairs signed rank test).

Additional file 11: Mycobiome data analysis.

Additional file 12: Figure S5. Bacterial communities within a subject before, during, and after treatment. Distribution of bacterial taxa detected in CF subjects either immediately after an exacerbation and before starting an antibacterial therapy (pre), approximately 2 weeks afterwards (post) or while hospitalized (inter) presented in Euler diagrams for the pre- and post-samples (subjects #1–6) and Venn diagrams for series with three samples (subjects #8 and 9). The numbers in parenthesis describe the total number of taxa detected in a sample; the numbers in the circles represent either the unique number of taxa in a sample or the number of shared taxa in the overlap regions.

Additional file 13: Table S7. Summary of fungal taxa found in more than one sample.

Additional file 14: Figure S6. Quantification of total fungi and specific *Candida* species in sputum of CF subjects. Quantification of the fungal burden of sputum samples from CF subjects. Fungal burden was determined by amplifying the 185 rDNA locus by qPCR. For *Candida* species-specific quantification, we used the genes CAWG_05066 (*C. albicans*), CPAR2_301290 (*C. parapsilosis*), and Cd36_16280 (*C. dublinensis*). Fungal genome copy number was determined by comparison to a standard curve generated using DNA isolated from pure cultures of *Candida albicans* as described in the methods.

Abbreviations

CF: cystic fibrosis; GAST: Global Alignment for Sequence Taxonomy; ITS1: internal transcribed spacer 1; MBL: Marine Biology Laboratories at Woods Hole; OTU: operational taxonomic unit; PCR: polymerase chain reaction; QIIME: quantitative insights into microbial ecology; qPCR: quantitative PCR; rDNA: ribosomal RNA genes; VAMPS: visualization and analysis of microbial population structures.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

AA, LAM, and EFD collected sputum samples. SDW, with important contributions from EFD, prepared sputum samples for deep sequencing analysis. SDW performed the qPCR analysis of fungi DNA, and SDW and LMF in consultation with GAO performed the qPCR analyses of bacterial DNA. THH performed the statistical analyses. SDW, HGM, SLG, and MLS performed the deep sequencing and its analysis. SDW, AS, and MLS modified and refined the fungal database for usage in VAMPS. DAH and AA contributed to study design. SDW and DAH wrote the manuscript with important contributions from coauthors, especially HLM and MLS. All authors read and gave approval to the final manuscript.

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