Enrichment of the airway microbiome in people living with HIV with potential pathogenic bacteria despite antiretroviral therapy

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ABSTRACT

Background: Long-term antiretroviral therapy (ART) enables people living with HIV (PLW-HIV) to be healthier and live longer; though they remain at greater risk of pneumonia and chronic lung disease than the general population. Lung microbial dysbiosis has been shown to contribute to respiratory disease.

Methods: 16S-rRNA gene sequencing on the Miseq-platform and qPCR for typical respiratory pathogens were performed on sputum samples collected from 64 PLW-HIV (median blood CD4 count 676 cells/µL) and 38 HIV-negative participants.

Finding: Richness and α-diversity as well as the relative-abundance (RA) of the major taxa (RA>1%) were similar between both groups. In unweighted-Unifrac β-diversity, the samples from PLW-HIV showed greater diversity, in contrast to the HIV negative samples which clustered together. Gutt bacterial taxa such as Bilophila and members of Enterobacteriaceae as well as pathogenic respiratory taxa (Staphylococcus, Pseudomonas and Klebsiella) were significantly more frequent in PLW-HIV and almost absent in the HIV-negative group. Carriage of these taxa was correlated with the length of time between HIV diagnosis and initiation of ART (Spearman-r=0.279, p=0.028).

Interpretation: Although the core airway microbiome was indistinguishable between PLW-HIV on effective ART and HIV-negative participants, PLW-HIV’s respiratory microbiome was enriched with potential respiratory pathogens and gut bacteria. The observed differences in PLW-HIV may be due to HIV infection altering the local lung microenvironment to be more permissive to harbour pathogenic bacteria that could contribute to respiratory comorbidities. Prompt start of ART for PLW-HIV may reduce this risk.

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1. Introduction

Effective antiretroviral therapy (ART) has changed the natural history of HIV infection. People living with HIV (PLW-HIV) and using ART now have near-normal life expectancy [1]. HIV complications have also shifted from acute opportunistic infections and neoplasms to non-AIDS related comorbidities [2]. Despite the reduction in respiratory infections such as Pneumocystis jirovecii pneumonia, bacterial pneumonia and tuberculosis with effective ART [3,4], respiratory illness remains common. PLW-HIV appear at higher risk of developing chronic lung disease such as chronic obstructive pulmonary disease (COPD), pulmonary hypertension and lung cancer [5].

Recent studies have demonstrated that the microbiome has an important role in determining respiratory health and can be a key player in the pathogenesis and prognosis of inflammatory chronic lung disease [6,7]. Through the impact of HIV on innate and adaptive immunity in the lungs, the potential exists to alter the lung microenvironment and affect the homeostasis of the microbial communities within it [8].

Current evidence suggests that, without ART, HIV infection alters the airway microbiome to reduce the richness and alpha diversity (a measurement of the taxa diversity within each sample) [9] of the microbial communities in the airways, and to enrich signature bacteria such as Tropheryma whippelii and some taxa such as Prevotella and Veillonella that may contribute to lung inflammation [10-13]. Effective ART appears to reverse these changes; data from the Lung HIV Microbiome Project suggests that the lung microbiome in healthy
PLW-HIV on ART with preserved blood CD4 counts is similar to uninfected individuals [14]. The importance of the environment on the microbiome (and hence the need for comparator populations) has been shown by Iwai et al. who compared the airway microbiome between two similar cohorts of PLW-HIV admitted to hospital with pneumonia in San Francisco and Uganda. They found that the airway microbiome of each cohort was distinct - which they attributed to local environmental factors [15,16].

In this paper, using 16S rRNA gene sequencing we describe the compositional differences in the airway microbiome of a cohort of adult PLW-HIV with high blood CD4 counts on ART and HIV uninfected participants within the UK. We then add to this data by specifically evaluating the presence of three pathogenic respiratory species with targeted quantitative PCR.

2. Methods

2.1. Study design and population

The present work is part of a prospective observational cohort study conducted in the Royal Free London NHS Foundation Trust comparing the incidence of acute respiratory illness in PLW-HIV and HIV negative individuals. The study was approved by the London Hampstead Research Ethics Committee (14/LO/1409) and registered with the ISRCTN registry (ISRCTN38386321). Participants were recruited between November 2015 and January 2017. PLW-HIV were invited to participate when they attended ambulatory HIV care appointments. The only eligibility criteria were age over 18 years, consent to participate, and absence of symptoms of acute respiratory illness at study entry. HIV negative participants were recruited from Sexual Health clinics and Primary Care, using primary care records to invite potential participants with similar age, gender, and smoking status to the expected characteristics of the HIV positive population sampled. HIV negative participants had their HIV status confirmed by blood test at recruitment. All participants provided written informed consent. Participants were invited to provide a sputum sample for analysis at the baseline visit. Sputum samples that were collected from participants who could expectorate were used in the present analysis. Sputum samples were stored at –80 °C within 3 h.

Details on methods are given in the online data supplement. Briefly, sputum samples were treated with Sputasol® (Oxoid, UK) then were heated at 95 °C for 30 min. One millilitre of each sample was centrifuged at 13,000xg for 10 min; the pellets were disrupted with glass beads on Fast-Prep®-24 Instrument for 45 s. Metagenomic DNA was then extracted on the automated DiaSorin® btx extraction platform using DiaSorin® Arrow DNA extraction kit.

The bacterial loads of Pseudomonas aeruginosa [17] and Haemophilus influenzae, Moraxella catarrhalis and Streptococcus pneumoniae [18] were quantified using two quantitative-PCRs (qPCR) methods on Qigen Rotor – gene® 6000 real-time PCR machine (Corbett Research UK, Cambridgeshire, UK). The bacterial load of each tested organism was calculated in colony forming unit (CFU) per mL of sputum and a mean of technical triplicates was taken for each sample. An internal amplification control, SPUD A, was used at final concentration of 0.04 pM to test for PCR inhibition [19].

A sequence library was created by amplification of V3-V4 regions of the bacterial 16S rRNA gene through conventional PCR on the extracted metagenomic DNA using 341 forward primer and 805 reverse primer. Each sample was assigned a unique pair combination of standard Illumina® dual indexed primers (with adaptors attached). The PCR products (577bp amplicons) were cleaned up using Agencourt AMPure XP beads (Beckman Coulter, UK). The samples were pooled in an equimolar ratio at 5 nM into one library which was then checked on bioanalyzer. Sequencing was performed using Illumina MiSeq Platform using costume sequencing primers, Miseq® Reagent Kit v2 (500 cycles). Additional details on the PCRs and primers sequences are provided in the online data supplement. Sequence data are deposited in the European Nucleotide Archive (ENA), study accession number is PRJEB32399.

An extraction negative control and a no-template PCR control (water) were run alongside each batch of samples throughout the process as negative controls to allow for the evaluation of potential contamination. A laboratory prepared mock community was run as a positive control.

2.2. Bioinformatics and statistical analysis

In bioinformatic analysis we adopted the workflow established by Microbiome helper [20] using QIIME pipeline v1.9.1. [21] where the sequences were clustered based on 97% similarity into Operational Taxonomic Units (OTU) and taxonomic classification was assigned to OTUs using open reference OTU picking against Greengenes database version 13.8. The OTU table was then rarefied per sample to 20,000. Alpha and beta diversity indices were calculated on the rarefied OTU table. The appropriate statistical significance tests were calculated using SPSS v. 23 or QIIME wrapper scripts. STAMP (v2.1.3) [22] was used to visualize the results and explore the OTUs showing significant differences in relative abundance between the two study groups using White’s non-parametric t-test [23] and all p-values were corrected using Benjamin–Hochberg False Discovery Rate (FDR) method for multiple comparisons.
Table 1
Demographic and clinical characteristics of participants.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>HIV Positive (n=64)</th>
<th>HIV Negative (n=38)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>52 (10-47)</td>
<td>53 (7-76)</td>
<td>0.59a</td>
</tr>
<tr>
<td>Sex</td>
<td>Males 55 (86%)</td>
<td>29 (76%)</td>
<td>0.218a</td>
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<tr>
<td></td>
<td>Females 9 (14%)</td>
<td>9 (24%)</td>
<td></td>
</tr>
<tr>
<td>Sex</td>
<td>Heterosexual</td>
<td>29 (76%)</td>
<td>&lt;0.001a</td>
</tr>
<tr>
<td></td>
<td>Homosexual</td>
<td>9 (24%)</td>
<td></td>
</tr>
<tr>
<td>Sex</td>
<td>Bisexual</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Ethnicity</td>
<td>Caucasian 51 (80%)</td>
<td>36 (95%)</td>
<td>0.108d</td>
</tr>
<tr>
<td></td>
<td>Black/African or Caribbean 7 (11%)</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Asian 2 (3%)</td>
<td>1 (3%)</td>
<td></td>
</tr>
<tr>
<td>Education</td>
<td>University Degree 28 (46%)</td>
<td>20 (54%)</td>
<td>0.912d</td>
</tr>
<tr>
<td></td>
<td>Secondary Education or equivalent qualifications 21 (34%)</td>
<td>10 (28%)</td>
<td></td>
</tr>
<tr>
<td>BMI</td>
<td>26.18 (4.93)</td>
<td>25.04 (2.91)</td>
<td>0.789d</td>
</tr>
<tr>
<td>Self-reported Comorbidities</td>
<td>Asthma 8 (21%)</td>
<td>5 (36%)</td>
<td>0.292d</td>
</tr>
<tr>
<td></td>
<td>COPD 2 (5%)</td>
<td>1 (7%)</td>
<td>1.000d</td>
</tr>
<tr>
<td></td>
<td>Cancer 7 (18%)</td>
<td>1 (7%)</td>
<td>0.665d</td>
</tr>
<tr>
<td></td>
<td>Heart Disease 5 (13%)</td>
<td>0 (0%)</td>
<td>0.309f</td>
</tr>
<tr>
<td></td>
<td>Diabetes 4 (10%)</td>
<td>0 (0%)</td>
<td>0.563d</td>
</tr>
<tr>
<td>Smoking</td>
<td>Current Smoker 25 (39%)</td>
<td>4 (11%)</td>
<td>0.009f</td>
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<tr>
<td></td>
<td>Ex-smoker 21 (33%)</td>
<td>16 (43%)</td>
<td></td>
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<tr>
<td></td>
<td>Never smoked 18 (28%)</td>
<td>17 (46%)</td>
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<td></td>
<td>Recreational drug use (ever) 48 (75%)</td>
<td>20 (57%)</td>
<td>0.067a</td>
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<tr>
<td></td>
<td>Recreational drug use (last 3 months) 25 (39%)</td>
<td>5 (14%)</td>
<td>0.007a</td>
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<tr>
<td></td>
<td>CD4 count (cells/μL) 676 (880-467)</td>
<td>1030 (1377-723)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Nadir CD4 (cells/μL) 225 (371-107)</td>
<td>676 (880-467)</td>
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<td></td>
<td>CD4/CD8 Ratio 0.67 (0.92-0.43)</td>
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<tr>
<td>Subjects with Blood HIV Viral load &lt; 40 copies/mL</td>
<td>54 (84%)</td>
<td>10 (16%)</td>
<td>0.016f</td>
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<td>Blood HIV Viral load &gt; 40 copies/mL Viral load (copies/mL)</td>
<td>10 (16%)</td>
<td>22 (37-71)</td>
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</tr>
<tr>
<td>Prescribed Antiretroviral therapy</td>
<td>63 (100%)</td>
<td>63 (100%)</td>
<td></td>
</tr>
<tr>
<td>Prescribed Antiretroviral therapy (years)</td>
<td>10.5 (15-5)</td>
<td>10.5 (15-5)</td>
<td></td>
</tr>
<tr>
<td>Prescribed Antiretroviral therapy over 10 years</td>
<td>32 (53%)</td>
<td>32 (53%)</td>
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<tr>
<td>FEV1 (L)</td>
<td>3.28 (3.79-2.77)</td>
<td>3.62 (4.09-2.84)</td>
<td>0.132d</td>
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<td>FEV1 z-score</td>
<td>-0.53 (-1.18-0.07)</td>
<td>0.115d</td>
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<tr>
<td></td>
<td>FVC (L)</td>
<td>4.30 (4.74-3.37)</td>
<td>0.061d</td>
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<tr>
<td></td>
<td>FVC % predicted</td>
<td>93% (99.8-81.2)</td>
<td>0.008d</td>
</tr>
<tr>
<td>Airflow Obstruction (≤C21)</td>
<td>10 (16%)</td>
<td>2 (3%)</td>
<td>0.202f</td>
</tr>
<tr>
<td>Restrictive Airflow (≤C47)</td>
<td>14 (22%)</td>
<td>3 (5%)</td>
<td>0.052f</td>
</tr>
<tr>
<td>Inhaled respiratory medication</td>
<td>10 (16%)</td>
<td>4 (11%)</td>
<td>0.500f</td>
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<tr>
<td></td>
<td>R2 Agonist ICS</td>
<td>5 (8%)</td>
<td>2 (0%)</td>
</tr>
<tr>
<td></td>
<td>4 (7%)</td>
<td>0 (0%)</td>
<td>0.293f</td>
</tr>
</tbody>
</table>

a Mean (SD)

b Median (IQR)

c n (%)

d p-value by Mann-Whitney Test

e p-value by Chi square

f p-value by Fisher Exact Test

g FEV1/FVC < 0.7

h FEV1/FVC ≥ 0.7 and FVC < 80% predicted

BMI: Body Mass Index

FEV1: Forced Expiratory volume in 1 sec

FVC: Forced Vital Capacity

ICS: Inhaled Corticosteroids

2.3. Role of funding source

None of the funders had any role in paper design, data collection, data analysis, interpretation, or writing of the paper.

3. Results

3.1. Clinical and demographic Cohort characteristics

Overall 102 participants out of 217 in the wider cohort provided sputum samples at baseline assessment. There were no significant differences between those who could and couldn’t expectorate, with respect to airflow obstruction (p = 0.34), blood CD4 count (p = 0.95), respiratory symptoms (defined by St Georges Respiratory Questionnaire (SGRQ) total score) (p = 0.16), age (p = 0.16), smoking status (p = 0.127) nor HIV status (p = 0.59) as similar proportion of PLW-HIV (45%) and controls (49%) produced sputum. Sputum samples collected from 64 PLW-HIV using ART and 38 HIV negative participants were included (Table 1). All participants were free of symptoms of acute respiratory illness (more details in the online data supplement). The median length of time between HIV diagnosis and starting ART was 1342 days (IQR 3131-448 days).

Fifty-three percent of PLW-HIV had been taking ART for over 10 years and 84% had undetectable HIV plasma load (VL) (<40 copies/mL). Ten participants (16%) had detectable VL (median 412 copies/mL, IQR 148-71). The median blood CD4 count was 676 cells/μL (IQR 880-467 cells/μL); whilst the nadir CD4 was 225 cells/μL (IQR 371-107 cells/μL) of which 44% had nadir CD4 < 200 cells/μL.
No significant differences were found between the two groups regarding age which ranged between 30 and 74 years old, sex, ethnicity, educational level, body mass index (BMI), and self-reported co-morbidities; e.g. asthma, COPD, heart disease, cancer and diabetes. However, current tobacco smoking and recent use of recreational drugs were significantly higher in PLW-HIV. The spirometry results were normal for most participants; though, the percentage of FVC predicted was significantly lower in the HIV positive group (Median 93% vs 98% in the comparator group, p < 0.01). Although no statistically significant differences were detected in terms of obstructive airflow which we defined as FEV1/FVC < 0.7, or restrictive airflow which we defined as FEV1/FVC > 0.7 and FVC < 80% predicted, airflow obstruction was three times higher among PLW-HIV.

3.2. Bacterial communities composition

All richness and alpha-diversity indices such as number of observed OTUs or Chao1 (Fig. 1a and 1b respectively) as well as PD whole tree, Fisher alpha, Shannon and Simpson diversity indices (data not shown) were not significantly different between PLW-HIV and HIV uninfected participants. However, within the HIV positive group the number of observed OTUs, Chao1 (Fig. 1c and 1d respectively), PD whole tree and Fisher alpha (data not shown) were significantly lower in participants whose blood CD4 count was < 500 cells/µL.

In principal co-ordinate analysis (PCoA) of weighted Unifrac β-diversity index (PERMANOVA p = 0.283), the samples did not cluster by HIV status (PERMANOVA p = 0.283). In addition, the core taxa constituting the microbiome profiles in sputum at both the phylum and genus level (Fig. 2b and 2c respectively) were very similar between the HIV positive and negative groups, and no significant differences in the relative abundances (RA) of the highly abundant taxa whose RA > 1% were detected. Out of 782 OTUs that were classified into 252 taxa, only 20 were present at RA > 1%, and they were present in all samples. Nevertheless, the samples were distinguished based on the HIV infection status in unweighted UniFrac β-diversity index (PERMANOVA p = 0.002), which unlike the weighted UniFrac is more sensitive to the presence or absence of taxa regardless of the weight of their relative abundances. In PCoA of unweighted UniFrac, the controls comparator group formed a cluster while PLW-HIV the HIV positive group were more dispersed, depending on the presence of some minor taxa (Fig. 3).

Looking more closely at the microbiome profile, some low abundant taxa such as *Staphylococcus aureus* (p = 0.0016), *Pseudomonas* (p = 0.0093), genus *Lactobacillus* (p = 0.029), *Klebsiella* (p = 0.024), *Bilophila* (p < 0.0001), and family *Enterobacteriaceae* (p = 0.04) were significantly more prevalent and abundant in the PLW-HIV compared to the HIV negative comparator group (Fig. 4). *Staphylococcus aureus* (OTU 1404220) and *Pseudomonas* (OTU 144452), were present in 8% (n = 5) of PLW-HIV, and *Klebsiella* (OTU 144814) in 9% (n = 6). In contrast, these OTUs were completely absent in controls. *Bilophila* (OTU 359872) was detected in 13% (n = 8) of HIV positive and 3% (n = 1) of HIV negative group. The relative abundance of both genus *Lactobacillus* and family *Enterobacteriaceae* taxa were significantly higher in PLW-HIV – being present in 66% (n = 42) and 30% (n = 19) respectively of PLW-HIV samples, 45% (n = 17) and 18% (n = 7) in the HIV negative group. *Lactococcus* (OTU 557570) was present in 5% (n = 3) and 8% (n = 3) of the PLW-HIV and HIV negative comparators respectively, though was significantly more abundant in the former (p = 0.0068). (Fig. S1 online data supplement).

Using the Nucleotide BLAST analysis tool against the NCBI 16S ribosomal RNA database [24], the top hits with the maximum scores identified the following OTUs: 1040220, 144452, and 144814 as 99% identical to *S. aureus*, *Ps. aeruginosa*, and *Klebsiella pneumoniae* strains respectively. None of these OTUs or taxa were detected in the negative controls.
Fig. 2. Microbiome Profile of HIV Negative and Positive groups. (a) PCoA of weighted UniFrac β-diversity index showing no separation with respect to the HIV status (p=0.283, PERMANOVA). (b) Microbiome profile at the phylum level (p>0.05, Mann-Whitney non-parametric t-test). (c) Microbiome profile at the genus level. NCext: extraction negative control, NCpcr: no-template PCR negative control, Mock: laboratory prepared mock community. Group size: 38 HIV-ve group and 64 HIV +ve group.

Fig. 3. PCoA plot of unweighted UniFrac β-diversity (p=0.002, PERMANOVA). a) With respect to HIV status, Blue: HIV –ve group (n=38), orange: HIV positive (n=64). b) With respect to the presence of pathogenic taxa: S. aureus (yellow), Pseudomonas (green), Klebsiella (orange), Lactobacillus (light blue), Bilophila: (red), otherwise the free HIV-ve samples are black and the free HIV positive samples are grey.
Fig. 4. The Relative Abundance of (a) genus *Lactobacillus* (*p*=0.029) (b) *S. aureus* (*p*=0.0016) (c) genus *Pseudomonas* (*p*=0.0093) (d) genus *Klebsiella* (*p*=0.024) (e) Family *Enterobacteriaceae* (*p*=0.04) (f) genus *Bilophila* (*p* < 0.0001) in the individual sputum samples. Group size: 38 HIV-ve group (blue) and 64 HIV +ve group (orange), *p*-values by White non-parametric t-test, corrected by Benjamini-Hochberg FDR method.
The co-existence of the three respiratory pathogens was prevalent in both groups (detected in 8% of PLW-HIV and 14% in HIV comparator group, where Spn was present in 39% and Hi in 37%, with S. pneumoniae detected in 57% and 44% respectively with both organisms being bacterium excluded from the analysis. Error bars show material. Loads are presented in CFU/mL of original sputum sample for each bacterium, means were calculated for each study group. Samples which gave negative results for a given bacteria were excluded from the analysis. Error bars show ±1 SD; Group size: 38 HIV-ve group and 64 HIV +ve group.

3.3. Load and prevalence of airway bacteria using q-PCR

The prevalence of the three bacteria: S. pneumoniae (Spn), H. influenzae (Hi) and M. catarrhalis (Mc) were similar within the two study groups (Fig. 5a). In PLW-HIV S. pneumoniae and H. influenzae were detected in 57% and 44% respectively with both organisms being identified together in 25%. This compares to the HIV negative comparator group, where Spn was present in 39% and Hi in 37%, with both organisms present together in 16%. M. catarrhalis was the least prevalent in both groups (detected in 8% of PLW-HIV and 14% in HIV negatives). The co-existence of the three respiratory pathogens was similar in both groups in less than 5% (Fig. 5b).

The mean bacterial load of H. influenzae was lower in PLW-HIV compared to controls; the difference between the means of the two groups was −1.04 log_{10} (p = 0.008). Within the HIV positive group, the mean bacterial load of H. influenzae in participants with blood CD4 count <500 cells/μL was significantly lower by 0.95 log_{10} than those with a CD4 count ≥500 cells/μL (p = 0.035) (Fig. 5d). The mean bacterial load of M. catarrhalis was slightly but not significantly lower by 1.3 log_{10} in the HIV positive group compared to controls. S. pneumoniae was the most populous in both groups with a mean bacterial load (SD) of 5 (1.3) log_{10} CFU/mL which was similar within the two groups (Fig. 5c).

The prevalence of P. aeruginosa was confirmed using qPCR method. P. aeruginosa was detected in 8.7% of HIV positive group and was completely absent in the HIV negative comparator group; however, the detected loads were too low, less than the method's sensitivity (limit of quantification (LOQ) 500 CFU/mL) in most cases (data not shown).

4. Discussion

The primary aim of this study compared the airway microbiome of a UK cohort of PLW-HIV using effective ART with good blood CD4 counts, compared to an HIV negative comparator group. Our results support the findings of similar US cohort studies,[14,25] in that the core microbiome in the two populations was very similar in terms of the relative abundance of the major taxa, and the richness and α-diversity of the sputum microbial communities. However, in principal coordinate analysis (PCoA) of unweighted Unifrac β-diversity index, the samples from PLW-HIV were more dispersed, whereas those from the HIV negative comparator group were more similar and clustered together.

We find, for the first time, potentially important differences in the prevalence of some minor taxa (relative abundance <1%) that may have a role in the pathogenesis of lung disease such as Staphylococcus,
**Pseudomonas** and **Klebsiella** being detected significantly more frequently in a subset of PLW-HIV (around 8.5%), yet were almost absent in controls.

Although the 16S rRNA sequencing suggest that these pathogenic OTUs were minor taxa, the result of the mock community, run alongside the samples as a positive control for sequencing (Fig. S7 online data supplement), indicated that there might be some negative bias against the amplification of 16S rRNA gene, especially from Klebsiella and Pseudomonas. In line with our results, these pathogenic taxa are not normally detected in the healthy airway microbiome and are more associated with inflammatory chronic lung disease and acute pneumonia infections.[6,7,26]

**Lactobacillus** and **Lactococcus** were significantly more abundant in PLW-HIV - a similar observation to that reported by Twigg et al and Yang et al.[11,27,28] Other taxa which were enriched in the sputum of some PLW-HIV are regarded as gut bacteria. These include Bilophila and members of family Enterobacteriaceae. Similarly in the lung HIV Microbiome Project Lozupone et al. reported the enrichment of bronchoalveolar lavage from PLW-HIV with *Tropheryma whipplei* which is a gut bacterium and the etiological agent of Whipple’s disease and whose relative abundance was reduced following effective ART.[13]

A finding also noted by Twigg et al.[11]. The enrichment of the lung microbiome with gut bacteria may be explained by the known breakdown of the gut mucosal barrier in HIV infection leading to increased translocation of gut microbes and microbial products into the systemic circulation through which they can eventually lodge and find a favorable ecological niche in the lungs of PLW-HIV.[10,11,29].

The enrichment of the respiratory microbial communities with gut and/or potential pathogenic respiratory bacteria was in subsets of PLW-HIV, and it may also explain the dispersal of samples from PLW-HIV in the PCoA of unweighted Unifrac ß-diversity (Fig. 3). We also found a positive correlation between the presence of these taxa in the sputum and the length of time between HIV diagnosis and starting ART. However, this observation needs to be confirmed by a suitably powered future study.

Previous studies have associated respiratory microbiome enrichment with *Prevotella* and *Veillonella* and the enhanced alveolar inflammation that may contribute to chronic lung pathology in PLW-HIV.[7,27,30]. In contrast, numerous studies have linked these two genera with lung health where they were promptly replaced by members of *Proteobacteria* or *Firmicutes* in chronic lung disease.[7,31,32]. In our cohort, the relative-abundances of these two genera were not significantly different in the sputum samples of both HIV positive and negative groups. In a Canadian cohort, these two genera were less abundant in bronchoscopic specimens from PLW-HIV compared to HIV negative controls.[33]. Therefore, the role of *Prevotella* and *Veillonella* species in respiratory microbiome generally, and in PLW-HIV specifically, remain controversial. More work is needed to understand the interactions and dynamic changes of these two genera with the immune tone and ART in HIV infected population.[34].

Our primary analysis was to compare the airway microbiome between PLW-HIV and HIV uninfected subjects, however, microbiome data are complex and multidimensional. To address this many parameters and indices were applied to summarise and compare the data. The alpha and beta diversity indices were more dispersed within PLW-HIV group compared to the HIV negative comparator group. Variation in the peripheral blood CD4 count may account for the observed dispersion in PLW-HIV. To test this the HIV positive group was stratified with respect to the peripheral CD4 count. CD4 was found to be a significant covariate in unweighted Unifrac ß-diversity (p = 0.01) within this group. Furthermore, α-diversity was significantly lower in PLW-HIV with lower CD4 count. A similar observation was reported by Twigg et al.[27].

Recent epidemiological data showed that although the incidence of pneumonia has been reduced in the ART era, PLW-HIV are at greater risk of bacterial pneumonia - a major cause of morbidity and mortality in this population.[35]. *S. pneumoniae* followed by *S. aureus* and *H. influenzae* are the most frequently identified etiologic agents of pneumonia in PLW-HIV.[28] It was, therefore, important to specifically detect the prevalence and load of these typical respiratory bacterial pathogens in our cohort using a specific qPCR method as the resolution of 16S rRNA sequencing is often restricted to the genus level for most taxa and sputum contains high proportions of *Streptococcus* viridans and commensal *Haemophilus*. Quantitative PCR results demonstrated that *S. pneumoniae* followed by *H. influenzae* were the most prevalent in the PLW-HIV. This is consistent with our previous findings using the same method in a similar cohort.[36]. There were no significant differences in the loads and prevalence of *M. catarrhalis* and *S. pneumoniae* in both HIV positive and negative groups; however, *S. pneumoniae* was slightly more prevalent in the HIV positive group though this study lacked the power to prove a significant difference.

Limitations of our work include the potential risk of contamination of the sputum with microbes from the upper respiratory tract and oral cavity. Quality assessment was not performed on sputum samples to evaluate pharyngeal contamination. Nevertheless, our main finding relates to the detection of some pathogenic and gut bacterial taxa, that are not normally oral taxa, in samples only from PLW-HIV. Also, samples from other body sites were not available; therefore, we could not compare the microbial signature between upper and lower respiratory compartments and the gut. We relied on the participant’s ability to expectorate sputum; and this is likely to be simpler for people who smoke or have underlying lung disease – raising the possibility of selection bias when comparing our study population to the wider HIV community. Current tobacco smoking and recreational drug use were significantly more frequently identified in our HIV positive group which are also common in other HIV cohort studies. Both factors were found to be significant covariates in ß-diversity indices. Confounder analysis is a challenge in microbiome studies.[37,38]. Since these two variables are inherent in HIV population, adjustment by propensity score analysis matching.[39] in a relatively small sample size would result in a diminished HIV negative comparator group. Therefore, the confounding effect of smoking and recreational drug use could not be completely excluded in our data. Nevertheless, the taxa contributing to the observed significant differences in our data based on the smoking status and recreational drug use, were all oral taxa; and apart from *Lactobacillus* which was also differentially abundant between HIV positive and negative groups (Figure S5 online data supplement), none was related to our main finding where potential pathogenic taxa (*S. aureus, Pseudomonas, Klebsiella*) and gut taxa (*Enterobacteriaceae* and *Bilophila*) were enriched in terms of prevalence and/or abundance in PLW-HIV in comparison to the HIV negative comparator group. Applying regression analysis would be invalid in this case as the former two covariates and HIV status are strongly correlated. Further studies on a larger sample size would allow adjustment for the potential confounders.

In conclusion, the core microbiome in the airways of people living with HIV who are on effective ART and clinically well managed, is indistinguishable from the general population in terms of the major bacterial taxa and alpha diversity. However, HIV infection may alter the local lung microenvironment to be more permissive to harbor potential respiratory pathogenic bacteria such *Staphylococcus, Pseudomonas* and *Klebsiella*, enabling gut bacteria including *Bilophila* and members of family *Enterobacteriaceae* to occupy ecological niches in the lower respiratory tracts of PLW-HIV. The carriage of these pathogenic bacteria may contribute to respiratory complications, such as bacterial pneumonia and inflammatory chronic lung disease. The mechanism underlying the perturbation of the respiratory microbiome is unclear, though is likely to relate to the interaction between the host, HIV and the environment. Our findings emphasize the importance of protecting the lung long-term in PLW-HIV through prompt initiation of ART.
Authors' contributions

SR and JB processed the samples and did the laboratory work, SR performed the bioinformatics and data analyses, SR, TMcH, MJ, JH, ML and DS interpreted the data, JB & EP collected the samples and clinical data, ML is the principal investigator. SR wrote the first draft of the manuscript with contributions from all other authors. All authors read and approved the final version of the manuscript.

Declaration of Competing Interest

There were no conflicts of interests.

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Supplementary materials

Supplementary material associated with this article can be found in the online version at doi:10.1016/j.eclinm.2020.100413.

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