Microbiota Core in Central Lung Cancer with Streptococcal Enrichment as a Possible Diagnostic Marker

CURRENT STATUS: POSTED

Salvador Bello
Hospital Universitario Miguel Servet

José J. Vengoechea
Hospital Universitario Miguel Servet

Manuel Ponce-Alonso
Hospital Universitario Ramon y Cajal

Ana L. Figueredo
Hospital Universitario Miguel Servet

Elisa Mincholé
Hospital Universitario Miguel Servet

Antonio Rezusta
Hospital Universitario Miguel Servet

Paula Gambó
Hospital Universitario Miguel Servet

Juan M. Pastor
Universidad Politécnica de Madrid

Javier Galeano
Universidad Politécnica de Madrid

Rosa del Campo
Hospital Universitario Ramon y Cajal

Email: rosacampo@yahoo.com Corresponding Author
ORCiD: https://orcid.org/0000-0003-1147-7923
SUBJECT AREAS
  General Microbiology
KEYWORDS
  microbiome, bronchoscopy, lung cancer, microaspirations, Streptococcus
Abstract

Background: Dysbiosis has been scarcely explored in the respiratory tract with cancer. We aimed to define the bacterial and fungal microbiota of the bronchi in cancer versus a healthy state, also defining the microbiota core, and their correlation with that in saliva and feces; and to detect markers for the early diagnosis of central lung cancer. For this purpose, twenty-five patients with central lung cancer and sixteen healthy controls without antimicrobial intake during the previous month were recruited. Bacterial and fungi distribution was determined by massive sequencing in bronchial biopsies. Complex computational analysis was performed to define for the first time the lung microbiota core.

Results: A greater abundance of Streptococcus, Rothia, Gemella and Lactobacillus distinguish the saliva of patients. Affected and contralateral bronchi of these patients have almost identical microbiota dominated by Streptococcus, whereas Pseudomonas was the major genera in controls. Oral and pulmonary ecosystems were significantly more similar in patients, probably due to microaspirations. Streptococcal bronchial abundance differentiates patients from controls by an ROC curve (90.9% sensitivity, 83.3% specificity, AUC=0.897). The mycobiome of controls (Candida) was significantly different from that of patients (Malassezia), with the cancer-affected bronchi similar to their saliva, but different from their contralateral bronchi.

Conclusions: Central lung cancer is highly enriched with Streptococcus, and shows significantly differences in their composition from healthy subjects. Alterations are not restricted to the tumor tissue, and seem to be the consequence of microaspirations from the oral cavity. These findings could be useful in the screening and even diagnosis of this pathology.

Background

Culture-independent techniques have revealed the composition of a stable microbiota within the distal airways [1]; and although normality criteria have not yet been established, atypical compositions linked to certain respiratory diseases such as cystic fibrosis, COPD or asthma have been detected [2].

Dysbiosis is typically detected surrounding tumors tissues [3], and this trait has been poorly explored
in the respiratory airway tissues, and using only surgical samples [4–8]. An understanding of the microbiota is necessary to decipher its possible causal role in cancer, and also to elucidate the prognosis and response to immunomodulatory therapies, because microorganisms and/or their metabolites shape the local microenvironment, influence the immune response, and impact the final battle against cancer [9]. Finally, the presence and/or abundance of specific bacteria could be used as a marker, as occurs with Streptococcus gallolyticus sbsp. gallolyticus in the colorectal cancer.

The aims of the present study were to: i) define the bacterial and fungal microbiota of central lung cancer, in relation to the corresponding contralateral bronchus and compared with healthy controls, also defining the core of those microbiotas; ii) correlate the pulmonary microbiota in lung cancer with the salivary and fecal compartments; iii) detect possible markers for the early diagnosis of central lung cancer.

Methods
Patients and samples
Twenty-five patients (24 men, mean age of 68 years) diagnosed with central lung cancer, directly visible and biopsied by bronchoscopy, were recruited (Table 1). The exclusion criteria included the intake of antibiotics, pre or probiotics, and systemic corticoids during the previous 4 weeks; the presence of acute infection; radio or chemotherapy in the last year, and immunodeficiency. Tumors were histologically classified into non-small cell lung cancer (n = 18, including 10 squamous, 4 adenocarcinoma, and 4 undifferentiated); and small cell lung cancer (n = 7). Each patient contributed with 4 samples: i)- saliva collected from a rinse with sterile distilled water, prior to the bronchoscopic procedure, ii)- biopsies of affected, and iii)- contralateral bronchi, and finally iv)- a fecal sample (provided by 18 out of the 25 patients). Bronchoscopy procedures were performed, as usually, by the nasal route (or oral if not possible) with a local instillation of lidocaine. Prior to the cancer sampling, biopsies for microbiota determination were obtained from contralateral nontumor-affected tissues.
Table 1
Demographic and clinical data of patients with lung cancer and healthy controls, differentiating between the entire control population (n = 16) and those included in the analysis of bacteria and fungi (n = 12) Statistically significant values that can differentiate control cases are highlighted in bold.

|                      | Cases (n = 25) | Controls (n = 16) | p-value | Controls (n = 12) | p-value |
|----------------------|---------------|-------------------|---------|-------------------|---------|
| **Age (years mean ± SD)** | 67.7 (12.8)    | 51 (14.4)         | .0003   | 52.1 (14.9)       | .002    |
| Male (%)             | 24 (96%)      | 8 (50%)           | .002    | 6 (50%)           | .003    |
| Active smoker (%)    | 14 (56%)      | 3 (19%)           | .02     | 1 (8.3%)          | .1      |
| Exsmoker (%)         | 11 (44%)      | 4 (25%)           | .3      | 4 (33.3%)         | .7      |
| Never smoked (%)     | 0 (0%)        | 9 (56%)           | < .001  | 7 (58.3%)         | < .001  |
| Familiar cancer (%)  | 8 (32%)       | 11 (69%)          | .04     | 8 (66.6%)         | .07     |
| Other cancer (%)     | 2 (8%)        | 3 (19%)           | .3      | 3 (25%)           | .3      |
| Hypertension (%)     | 13 (52%)      | 3 (19%)           | .05     | 3 (25%)           | .1      |
| Diabetes (%)         | 9 (36%)       | 0 (0%)            | .006    | 0 (0%)            | .01     |
| Dyslipemia (%)       | 16 (64%)      | 5 (31%)           | .08     | 5 (41.6%)         | .1      |
| Annual visit to dentist | 3 (12%)      | 10 (63%)          | .001    | 8 (66.6%)         | .001    |
| Lack of teeth (%)    | 21 (84%)      | 12 (75%)          | .6      | 9 (75%)           | .6      |
| COPD (%)             | 12 (48%)      | 0 (0%)            | .001    | 0 (0%)            | .003    |
| Mild (FEV₁ > 80%)    | 3 (25%)       | -                 | -       | -                 | -       |
| Moderate (FEV₁ 50-79%) | 8 (67%)      | -                 | -       | -                 | -       |
| Severe (FEV₁ < 50%)  | 1 (8%)        | -                 | -       | -                 | -       |
| Asthma (%)           | 0             | 0                 | 0       | 0                 | 0       |
| Interstitial Lung Disease | 0          | 0                 | 0       | 0                 | 0       |
| Respiratory function test |            |                   |         |                   |         |
| Normal (%)           | 10 (40%)      | 16 (100%)         | < .001  | 12 (100%)         | < .001  |
| Obstructive (%)      | 7 (28%)       | 0 (0%)            | .03     | 0 (0%)            | .07     |
| Mixed (%)            | 5 (20%)       | 0 (0%)            | .1      | 0 (0%)            | .1      |
| FEV mean ± SD        | 73.4 (16.6)   | 97.6 (11.4)       | < .001  | 96.6 (12.0)       | < .001  |
| FCV mean ± SD        | 89.3 (14.3)   | 103.7 (10.7)      | .001    | 101.9 (10.3)      | .01     |
| FEV/FCV mean ± SD    | 73.2 (12.9)   | 78.7 (5.3)        | .1      | 79.8 (8.4)        | .2      |

Simultaneously, 16 healthy controls were included, and each contributed the following: i)- oral microbiota and ii)- a single biopsy of their healthy bronchi. Controls without respiratory symptoms (except 2 with chronic cough) underwent bronchoscopy for non-cancer related indications (benign tracheal stenosis 9, fake haemoptysis 3, chronic cough 2, control of a previous endobronquial hamartoma resection 1, and dyspnea 1), and all of them had normal spirometries. All samples were immediately frozen at -80ºC after collection.

**Microbiota composition**
Samples were slowly defrosted at -20ºC for 24 h and 4ºC for another 24 h, to prevent bacterial death and DNA fragmentation. Total DNA was obtained by the QiaAmp kit (Qiagen) from the biopsies, from
the pellet of saliva after centrifugation, and from 200 µl aliquots of a solution of 0.5 gr of feces in 5 ml of water. Bacterial composition was determined by PCR amplification of the 16S rDNA V3-V4 region using published primers [10] in all samples, whereas the mycobiome was only analyzed in bronchial and saliva of the 16 controls and in a subset of 6 patients by amplification of the ITS-1 region [11]. PCR products were pooled equally and submitted to massive sequencing (2 × 300 bp) on a MiSeq (Illumina, San Diego, CA, USA) platform, at FISABIO (Valencia, Spain). Raw sequence data were deposited in Genbank (BioProject ID for bacterial sequencing: PRJNA586753; BioProject ID for fungal sequencing: PRJNA586768). QIIME2 software suite (2019.1 distribution) and LEfSE was used for analysis, and adequate negative sequencing controls were added in each process and run.

Bioinformatics for bacterial community characterization

Raw sequence data (FASTQ files) from 16S rDNA sequencing were demultiplexed and quality assessed using the q2-demux plugin. Then, denoising, filtering, and chimera removal were performed with DADA2 pipeline [12] (via q2-dada2 plugin), thus identifying all amplicon sequence variants (ASVs) [13] and their relative abundance in each sample. To minimize the number of spurious ASVs, those unique sequences with a total abundance lower than 7 reads across all samples were filtered out [14]. ASVs were firstly aligned and then used to construct a phylogenetic tree by following the align-to-tree-mafft-fasttree pipeline [15–16] from the q2-phylogeny plugin. ASVs were taxonomically classified by using the classify-sklearn naïve Bayes taxonomy classifier (via q2-feature-classifier plugin) [17] against the Silva 132 database [18]. Sequences not assigned to any taxa or classified as Eukaryote or Archaea, were filtered out. Diversity analysis was done using q2-diversity plugin, after samples were normalized via rarefaction (subsampled without replacement). Diversity analysis comprised alpha diversity metrics (Chao1, Shannon index, and Faith-pd [19], which measure microbiota degree of diversity) and beta diversity metrics (unweighted UniFrac [20] and weighted UniFrac [21], which measure microbiota composition differences between samples, while up-weighting differences in ASVs phylogenetic distance). Unweighted UniFrac reports differences in the presence or absences of ASVs, while weighted UniFrac also reports differences in the abundance of ASVs.

Bioinformatics for fungal community characterization

Raw sequence data from ITS1 sequencing were demultiplexed and quality assessed using the q2-demux plugin. Then, q2-itsxpress plugin [22] was used to quality filter and trim the ITS region from sequences. After that, denoising, merging, and ASVs calling were done by DADA2 pipeline (via q2-dada2 plugin), thus identifying all ASVs and their relative abundance in each sample. Very low abundance ASVs (total n < 7) were filtered out, as was done for bacteria. ASVs were taxonomically classified by using the classify-sklearn naïve Bayes taxonomy classifier, against the UNITE 7.2 database [23]. Diversity analysis was done using q2-diversity plugin, after samples were rarefied (subsampled without replacement). Diversity analysis comprised alpha diversity metrics (Chao1 and Shannon index), and beta diversity metrics Bray-Curtis distance [24], which measure non-phylogenetic microbiome composition differences between samples.

Quality and throughput of the sequencing process. 16 s rDNA sequencing in the 125 samples yielded 7,336,127 reads whereas the 4 negative controls 53, 220, 447, and 881 reads, discarding high contamination levels. After quality filtering, chimera removal, and discard of very low frequency sequences, a total of 4,897,627 reads were finally obtained. After taxonomic assignment, non-bacterial sequences were removed and 4,464,212 reads of 4434 ASVs were finally obtained. Three samples from affected bronchi (patients 23, 24, and 25) and four samples from control bronchi (controls 2, 3, 7, and 16) were not included in downstream analysis due to sequencing depth requirements (< 1000 reads per sample), and samples were rarefied to 1007 sequences to maximized the retention of samples and preserved representative diversity measures, according to alpha rarefaction plot.

The 50 samples using for mycobiome determination by ITS1 amplification and massive sequencing yielded 6,034,038 reads and the negative control performed 5,036 reads. After quality filtering, trimming, merging, and discard of very low frequency sequences, 3,535,951 reads of 515 ASVs were finally obtained. One saliva sample from a control, which read count was very low, was not included in downstream analyses due to sampling depth requirements (> 1000 reads). Rarefaction was set to 13,571 sequences per sample, since that sampling depth conserved all samples and reached a plateau according to alpha rarefaction plot inspection. The negative control was fully inspected to
identify which taxa were present.

Statistical analysis

Statistical differences in mean alpha diversity metrics between patients and controls were calculated by Kruskal-Wallis test [25]. Differences in microbiota composition between samples were assessed and plotted by performing Principal Coordinates Analysis (PCoA) based on the beta diversity metrics. Permutational multivariate analysis of variance (PERMANOVA) [26] was performed to determine which factors explained statistically significant variance in microbiota composition. All statistical tests were conducted via q2-diversity plugin from QIIME2. To determine which specific taxa explained beta diversity differences, differential abundance analyses were performed only in variables that yielded statistically significant differences in beta diversity analysis. For that purpose, linear discriminate analysis effect size (LEfSe) was used for testing taxonomic comparisons [27]. LEfSe uses the common tests for statistical significance (Kruskal-Wallis test and pairwise Wilcoxon test) with linear discriminate analysis for taxa selection. Alpha value for the factorial Kruskal-Wallis test was 0.01 and threshold on the logarithmic LDA score for discriminative taxa was set to 4.0. ROC curves were plotted by IBM SPSS Statistics software (Version 22.0. Armonk, NY: IBM Corp).

To get an overview of the complexity of the microbiome in lung cancer disease, we have designed the taxon interaction network for the different data (healthy lung, healthy saliva, affected lung, diseased contralateral lung, diseased saliva and diseased stool) Networks have nodes and links. The nodes represent the different taxa and the links represent that both taxa have appeared in the same patient sample. In this case, we have used weighted networks to represent the microbiome. The networks have been created as follows: for each patient, we select the taxa with non-zero abundances. Then, we build a complete network between these taxa. The link weight has been obtained as the product of the probability of the presence of taxa. Finally, we have added the networks of each patient in a global network. This analysis are available at https://github.com/JJ-Lab/Cancer_Lung_Microbiota website.

Results

Sample filtering

According to 16 s rDNA sequencing throughput analysis, three samples from affected bronchi
(patients 23, 24, and 25) and four samples from control bronchi (controls 2, 3, 7, and 16) were finally excluded from downstream analysis since they did not reach minimal sequencing depth requirements (set at > 1000 reads per sample).

**Alpha diversity**

Stools and saliva had similar alpha-diversity values, while bronchi were significantly more diverse according to Faith’s PD index, which considers not only the richness and relative abundance of ASVs but also its genetic distance (Fig. 1). Chao1 and Shannon diversity indices were significantly higher in patients’ bronchi than in controls’ (p < .001), while the Faith’s PD indexes were similar. The diversity of saliva was comparable between patients and controls.

**Microbiota composition**

The bacterial phyla distribution is shown in Fig. 2. Patients’ saliva presented a higher density of Firmicutes and Actinobacteria to the clear detriment of Proteobacteria, and this pattern was reproduced in the affected and contralateral bronchi. Fecal samples, only from patients, had a remarkably high proportion of Firmicutes (> 75%).

**Saliva**

Up to 213 genera were detected in saliva samples, with 13 comprising the majoritarian in both controls and patients: Streptococcus (19% and 23%, respectively), Prevotella (15 – 13%), Rothia (4–7%), Veillonella (7–8%), Neisseria (6 – 4%), Porphyromonas (6 – 4%), Haemophilus (6 – 2%), Gemella (3–5%), Fusobacterium (5 – 3%), Alloprevotella (4 – 1%), Actinomyces (2–2%), Granulicatella (1–2%), and Leptotrichia (1–2%). The remaining genera represented less than 1% of the total microbiota abundance.

The microbiota core composition of saliva from patients and controls were highly similar, although significant differences on bacterial proportions were detected by LEfSe and weighted UniFrac distance (PERMANOVA p < .005), but not by the unweighted UniFrac distance analysis (Fig. 3), emphasizing that those differences are linked to relative abundance of common taxa and no to presence or absence of specific taxa. PCoA within the major genera (limited to 90% of the total composition) (Fig. 3) demonstrated that Streptococcus, Rothia, Gemella and Lactobacillus densities are capable of distinguishing the saliva of patients from controls.
Bronchi

More than 450 bacterial genera were identified among the bronchial microbiota, with statistically significant differences in their composition or distribution among patients and controls, while the microbiota of the cancer-affected bronchus was almost identical to its contralateral counterpart (Fig. 4). Particularities depending on the histological type of cancer were not detected.

Lung cancer-related genera were Streptococcus (19% affected bronchus and 24% contralateral), Prevotella (9–9%), Blautia (5–4%), Veillonella (4–5%), Rothia (3–4%), Neisseria (2–3%), Gemella (2–2%), and Porphyromonas (2–2%). Health bronchi from control group was dominated by Pseudomonas (21%), followed by Streptococcus (8%), Actinobacter (6%), Veillonella (4%), Prevotella (3%), Delftia (3%), Janthinobacterium (3%), Escherichia-Shigella (2%), Haemophilus (2%), and Neisseria (2%). The representation of bacterial lung microbiota of both healthy controls and cancer patients using computational tools revealed a more complex ecosystem than feces (Fig. 4), also defining for the first time the core lung microbiota.

The minority bronchial microbiota (< 1%) comprised more than 400 genera, and included the predator Bdellovibrio, which was detected in 2 controls (0.24% and 0.47%) and in 2 patients (0.16% of affected, and 0.06% of contralateral bronchi). In 4 cases Pseudomonas (the natural prey of Bdellovibrio) was also present in the microbiota (38%, 14%, 5% and 12%, respectively).

Correlation between oral, fecal and bronchial microbiota

The distance between oral, fecal and bronchial microorganisms of each individual was assigned by pairwise analysis based on weighted UniFrac distance. The result of this analysis is shown in Fig. 5 and revealed that saliva and lung ecosystems were significantly more close (p < .001) in patients with lung cancer.

Streptococcal abundance characterizes the microbiota of patients

Streptococcus belongs to the Firmicutes phylum and their abundance was consistently the factor most able to distinguish the microbiota of patients from controls. This genus has a complex taxonomy and comprises numerous groups and species; we compared by phylogeny the 95 ASV obtained in at least two samples, also considering the exchange between ecosystems (Fig. 6). Feces were the most remote niche, with saliva and bronchi also distant. It is important to note that > 95% of the ASV
detected in the patients' bronchi were also in their saliva, while on the contrary only 36% of those present in saliva were also present in the lung. This fact might indicate that saliva is the main source of pulmonary Streptococcus. An ROC curve representing Streptococcus bronchial abundance distinguished patients from controls, particularly when a relative abundance > 14.6% predicted lung cancer with 90.9% sensitivity and 83.3% specificity (AUC = 0.897). Unfortunately, this result was not reproduced in saliva.

Mycobiome composition
Fungal characterization through ITS1 sequencing was determined in the 16 controls (saliva and bronchus) and a subset of 6 patients (saliva, affected and contralateral bronchi). Most of the fungal reads (n = 3,246) corresponded to Collybia and Amphinema genera from the Agaricomycetes phylum. Further inspection of the data demonstrated that those genera were absent in almost all samples, ruling out a cross-contamination event. Significant differences in the Chao1 index between the affected, contralateral, and control bronchi, and between patients’ and control’s saliva were not observed (Fig. 7). The Shannon index was significantly higher in affected bronchi from patients compared to that in bronchi from controls (p < .03), in accordance with the results obtained for bacteria.

A beta diversity analysis of fungal communities was performed by computing Bray Curtis distances, which was then used to build a PCoA plot (Fig. 7). In terms of fungal composition, affected bronchi from patients were similar to their saliva (p > .05), but different from both their contralateral bronchi (p < .006), and the bronchi (p < .001) and saliva (p < .007) from controls. This trend supports the aforementioned bacterial findings, which also suggest an interconnection between both anatomical ecosystems in lung cancer patients. A differential abundance analysis between affected bronchi from patients and from controls showed an enrichment of Malassezia in patients, whereas controls had a higher abundance of Candida (Fig. 7).

Discussion
The role of microbiota in carcinogenic processes has not yet been elucidated and will probably be different for each tumor and their localization. However, recent data clearly indicate that the
microbiota contributes to the prognosis of cancer and determines the response to treatments, particularly responses to the new immunomodulatory therapies [9, 28]. Criteria for the normal composition of lung microbiota have not yet been established, but the available data indicate that their composition in cancer patients differs considerably from that of healthy individuals [4–8]. Sampling in the respiratory system requires invasive methods, and here we have characterized for the first time, as far as we know, the respiratory microbiota surrounding central lung cancer by direct sampling of the tumor tissue by bronchoscopy. Our results reinforce the previously known particularities of the lung microbiota, which considerably differ from oral and stool microbial communities [29]. One important concern is the theoretical risk of contamination with the upper microbiota during the bronchoscopy, but our results allowed us to rule out significant contamination, as other authors had previously suggested [30–32]. Data analysis by various methodologies highlighted the particularities of the microbiota associated with cancer, but also defined the respiratory microbiota core in healthy conditions. Moreover, we considered strict inclusion criteria to avoid the possible bias of antibiotic or corticosteroid therapy, and the bacterial exchange between the anatomically separated niches such as saliva and feces. Finally, the mycobiome composition of central lung cancer was studied.

Significant differences in the lung ecosystem have been described based on health status or a lung cancer diagnosis in sputum [33], bronchoalveolar lavage [5], protected specimen brushing [6], cytological brushing [34], and surgical tissue [7, 8, 35]. The major contribution of our work is the study of the microbiome surrounding central cancer via direct sampling, but our results are not necessarily applicable to the distal airway. Interestingly, this central ecosystem was more diverse than the fecal or oral compartments. This finding contrasts, at least in part, with decreases in the microbiota biomass from upper to lower tract described in healthy people [32].

Low biodiversity is usually observed in various pathologies, including cancer, but we found significantly higher alpha-diversity values in cancer than in the control group. Other authors have published analogous [8] and opposing results [5, 6, 35]. Moreover advanced cancer stages [36] and reduced recurrence-free survival and disease-free survival [35] have been associated with higher
values of alpha diversity. Eighty-eight percent of our patients were at tumor stage III or IV, and their survival was only of 198 days in the follow up. Furthermore, other factors such as environmental exposure, residence in high-population density areas [4, 36], and pack-years of tobacco smoking, can increase the biodiversity of the lung microbiota, whereas chronic bronchitis reduces it [36]. All our patients had been smokers, while most of the controls (56%) had not (Table 1).

In terms of beta diversity, the tissues involved in cancer and the contralateral bronchus had almost identical compositions, as its have been previously published [6, 34, 35], probably reflecting the environmental influence which is not restricted to the cancer area. Although the limited size of our sample prevents us from reaching solid conclusions, no differences were detected in the composition of the bronchial microbiota as a function of the histological variants of the cancer. The abundance of Firmicutes to the clear detriment of Proteobacteria was the most noticeable result in our patients, and this result was consistent in all samples. Proteobacteria dominance in health lung microbiota, especially Pseudomonas, has been also previously corroborated [5, 6, 37]. Higher concentrations of Streptococcus, Blautia, Akkermansia, and Rothia were observed in patients, but Streptococcus was consistently the major marker linked to lung cancer. This fact has been previously reported in saliva [5], sputum [33, 38], bronchoalveolar lavage [5], lung tissue [7], and protected specimen brushing [6, 34]. The exhaustive analysis of the obtained ASVs allows us to suggest that the streptococcal variants present in lung tissue are similar to those found in saliva or feces, but the low length of the 16 s rDNA amplicon sequences (460 bp) preclude us from making robust assumptions in this regard. New studies including Streptococcus cultures and molecular characterization of the species are needed to decipher whether oral lineages are different from those found in the lungs or feces and thus establish whether there are any markers truly associated with lung cancer, as occurs with Streptococcus gallolyticus subsp. gallolyticus and colorectal carcinoma [39].

There is increasing evidence of a link between Streptococcus and lung cancer. Recently, Tsay et al. [34] detected Streptococcus and Veillonella enrichment in the lower airways with ERK and PI3K pathways upregulation - an early event that contributes to cell proliferation, survival and tissue invasion- combining microbiome and transcriptomic signatures. The major question that a remains to
be answered is whether the abundance of streptococci is a cause or consequence of the tumor process, as has been questioned in tumors from other localizations [40]. Streptococcus is a natural inhabitant of the oral cavity, which is connected to the lower respiratory tract by the larynx and trachea. The ASVs analysis allowed us to separate the oral streptococcal population from those found in the lung or gut, although the oral/lung bacterial exchange could occur via microaspirations [6, 34, 37].

Microaspiration events are common, but their frequency is significantly increased in chronic inflammatory airway diseases [37], inducing inflammation by elevation of Th17 lymphocytes, as well as expression of inflammatory cytokines (as IL-1α, IL-1β and IL-17). The alteration of the IL-23/IL-17 axis is well known in the pathogenesis of both autoimmune diseases and tumors. Recently, it has been described that Streptococcus mitis induced IL-1β, IL-6 and IL-23 transcription and Th17 responses able of releasing the potentially proinflammatory and protumoral IL-17 [41]. S. mitis also leads to neutrophil recruitment and inflammation, macrophage chemotaxis, a higher secretion of immunological inhibitory cytokine IL-10, and an increased immune checkpoint PD-L1 expression, facilitating the cancer development and expansion [41]. Lung resident γδ T cells, and their either protective roles or pro-tumorigenic functions in cancer have been recently discovered [42]. A study has provided evidence that local lung microbiota (one of the most common genera was Streptococcus) can provoke inflammation and tumor cell proliferation, via lung resident γδ T cells activation, that release IL-17, after IL-1β and IL-23 induction in myeloid cells by these local microbiota [43]. Our results demonstrated a global streptococcal enrichment in patients with cancer that affected more than just the respiratory tract, supporting the idea that microorganisms can orchestrate the balance between tumor-promoted inflammation and anti-tumor immunity depending on the specific microenvironment [43].

Streptococcal relative abundance in bronchial biopsies was a good predictor of lung cancer, but unfortunately was not reproducible in saliva. ROC curves suggested the contralateral bronchi as the best sample (90.9% sensitivity and 83.3% specificity; AUC = 0.897). Other authors found similar results in protected specimen brushing samples (87.5% of sensitivity and 55.6% specificity, AUC =
The proportional abundance of *Streptococcus* should be validated in the early stages of lung cancer with subsequent follow-up to corroborate this link. Along these lines, the intestinal enrichment of *Streptococcus* should be exhaustively explored to identify lung cancer markers in feces.

The intestinal and the respiratory ecosystems harbor a diverse and abundant microbiota, but some particularities distinguish both ecosystems. Food intake favors a higher rate of microbial reproduction increasing the total mass that is significantly reduced after defecation. On the contrary, nutritional sources for bacteria in the airway are limited to mucus and cellular debris, while clearance is carried out by the ciliary system and the immune system, particularly by macrophages. Food is an important mode of entry of foreign microorganisms into the gut ecosystem, as is inspired air. Alien microorganisms can lead to overestimation of the real diversity of the ecosystem, but we have implemented new analytic strategies to define the lung microbiota core, which had not previously been defined. Interesting findings of our work include the elevated alpha-diversity of the bronchial microbiota in comparison with saliva or feces, and the dominance of *Pseudomonas* in healthy individuals. The pulmonary presence of this genus is linked to cystic fibrosis and is the major pathogen that decreases the respiratory functionality within a pathogenic colonization. However, the lack of respiratory symptoms reduces a pathogenic role of *Pseudomonas* in healthy individuals, although more studies are needed in that line. Predator bacteria have been classically described in environmental ecosystems as scarcely detected in human samples [44–45]. Their low representation in the total ecosystem could prevent their detection by -omic strategies. This approach is an important field of research, which could represent an ecological key to modulating the microbiota composition.

The main limitation of our work is that we cannot estimate the effect of lung cancer factors, mainly tobacco (all patients had been smokers but only the half of the controls were) and COPD (12/25 patients) on the bronchial microbiota. However, it has not been established yet if tobacco has a significant influence on lung microbiota composition, and some important studies have shown contradictory results [36, 46–47].
Whereas severe COPD has been linked with significant alterations in the lung microbiota composition [48–49], mild and moderate COPD (92% of our COPD patients) has been associated with Streptococcus enrichment [50]. This finding might explain the association of mild/moderate COPD and lung cancer [51], although further studies are needed to confirm this association.

Our study has several additional limitations, including a small number of patients, all of them from the same hospital, a lack of other -omic analyses based on genetic expression, and we cannot rule out the possibility that the differences were be influenced by tobacco exposure. On the other hand, our strengths include performing the first study of microbiota combined with mycobiome of bronchial tissue obtained directly from tumor and contralateral bronchi (not adjacent to a resected tumor), as well as performing analysis of the connected ecosystems including saliva and feces.

The mycobiome results were consistent with those obtained for bacteria. The fungal community was slightly richer and more diverse in patients than in controls, although the contralateral bronchus was more similar to controls than to the affected counterpart. The mycobiome of saliva and the affected bronchus from patients matched perfectly, but differed in controls, again suggesting that in patients with cancer the bronchial microbiota is the result of a continuous exchange with that of the oral niche.

In terms of taxonomy, affected bronchi from patients had an enrichment of the Basidiomycota phylum with higher populations of Malassezia genus, whereas the enriched taxon in healthy individuals was the Ascomycota phylum and the genera Candida and Saccharomyces, as previously described [41, 52]. Although the public databases are increasing exponentially, it is important to note that a major limitation to describing the mycobiome is the lack of available taxonomic records. As far as we know, this is the first description of the lung mycobiome.

Conclusion

In summary, patients with central lung cancer have a significantly different bronchial microbiota from healthy controls, not restricted to tumor-involved tissue, and probably conditioned by continuous microaspiration events from the oral cavity, more than by the carcinogenic process. Lung cancer was associated with a considerable enrichment of Streptococcus and we propose that this feature could be used for the screening, diagnosis and prognosis of this pathology. Lung mycobiota differ considerably
in healthy individuals, and there are dissimilarities in patients between the affected and the contralateral bronchi. An innovative bioinformatics strategy used in this study has allowed us to define healthy individuals' the bronchial core microbiota, which is dominated by a non-pathogenic colonization of Pseudomonas.

**Abbreviations**

ASV: amplicon sequence variant; AUC: area under the curve; COPD: chronic obstructive pulmonary disease; FISABIO: Fundación para el Fomento de la Investigación Sanitaria y Biomédica de la Comunitat Valenciana; IL: interleukine; LDA: linear discriminant analysis; LEfSE: linear discriminant analysis effect size; PCoA: principal coordinate analysis; PERMANOVA: permutational multivariate analysis of variance; QIIME: Quantitative Insights Into Microbial Ecology; ROC: receiver operating characteristic.

**Declarations**

**Ethics approval and consent to participate**

The Ethics Committee "CEIC Aragón" approved this project in 2016 with the reference 15/2016, and all participants signed the informed consent after receiving all the information from the clinical researchers.

**Consent for publication**

Not applicable.

**Availability of data and materials**

All generated data is publically available.

**Competing interests**

RdC is the recipient of a Vertex grant IIS-2017-106179 for cystic fibrosis research. The remaining authors declare that they have no competing interest.

**Funding**

This work was partially supported by the project PI17/00115 which recipient is RdC. MPA was supported by the Programa Operativo de Empleo Juvenil, co-financed by the European Social Fund Investing in your future (ESF) and ERDF (PEJD-2018-PRE/BMD-8237), and by a Rio Hortega contract (CM19/00069) from the Instituto de Salud
Authors’ contributions
SB conceived the study. JJV, ALF, EM, and AR contributed to the patients’ selection and sample collection. JMP, JG carried out the computational analysis for core microbiota definition. MPA and RdC determined the microbiota composition. AR, PG, JMP, JG, and SB participated in the analysis and interpretation of data. SB, RdC, and MPA wrote and reviewed the manuscript, and finally all authors read and approved the final version.

Acknowledgements
The authors thank to Pilar García, Asunción Albericio, and the rest of nurse team of Interventional Pulmonology Unit, and to Yolanda Palacios, from Department of Microbiology of Miguel Servet Hospital for their contributions in the collection and conservation of the samples. The labor of Marta Cobo during sample processing is also recognized.

Authors’ information
Nothing to add.

References
1. Wu BG, Segal LN. Lung microbiota and its impact on the mucosal immune phenotype. Microbiol Spectr 2017; 5(3).
2. Pulvirenti G, Parisi GF, Giallongo A, Papale M, Manti S, Savasta S, Licari A, Marseglia GL, Leonardi S. Lower airway microbiota. Front Pediatr 2019; 7:393.
3. Chen J, Domingue JC, Sears CL. Microbiota dysbiosis in select human cancers: Evidence of association and causality. Semin Immunol 2017; 32:25-34.
4. Mao Q, Jiang F, Yin R, Wang J, Xia W, Dong G, Ma W, Yang Y, Xu L, Hu J. Interplay between the lung microbiome and lung cancer. Cancer Lett 2018; 415:40-48.
5. Wang K, Huang Y, Zhang Z, Liao J, Ding Y, Fang X, Liu L, Luo J, Kong J. A Preliminary study of microbiota diversity in saliva and bronchoalveolar lavage fluid from patients with primary bronchogenic carcinoma. Med Sci Monit 2019; 25:2819-2834.
6. Liu HX, Tao LL, Zhang J, Zhu YG, Zheng Y, Liu D, Zhou M, Ke H, Shi MM, Qu JM. Difference of
lower airway microbiome in bilateral protected specimen brush between lung cancer patients with unilateral lobar masses and control subjects. *Int J Cancer* 2018; 142:769-778.

7. Liu Y, O'Brien JL, Ajami NJ, Scheurer ME, Amirian ES, Armstrong G, Tsavachidis S, Thrift AP, Jiao L, Wong MC, Smith DP, Spitz MR, Bondy ML, Petrosino JF, Kheradmand F. Lung tissue microbial profile in lung cancer is distinct from emphysema. *Am J Cancer Res* 2018; 8:1775-1787.

8. Greathouse KL, White JR, Vargas AJ, Bliskovsky VV, Beck JA, von Muhlinen N, Polley EC, Bowman ED, Khan MA, Robles AI, Cooks T, Ryan BM, Padgett N, Dzutsev AH, Trinchieri G, Pineda MA, Bilke S, Meltzer PS, Hokenstad AN, Stickrod TM, Walther-Antonio MR, Earl JP, Mell JC, Krol JE, Balashov SV, Bhat AS, Ehrlich GD, Valm A, Deming C, Conlan S, Oh J, Segre JA, Harris CC. Interaction between the microbiome and TP53 in human lung cancer. *Genome Biol* 2018; 19:12.

9. Ma W, Mao Q, Xia W, Dong G, Yu C, Jiang F. Gut microbiota shapes the efficiency of cancer therapy. *Front Microbiol* 2019; 10:1050.

10. Gurevich A, Saveliev V, Vyahhi N, Tesler G. QUAST: quality assessment tool for genome assemblies. Bioinformatics. 2013; 29:1072-1075.

11. Toju H, Tanabe AS, Yamamoto S, Sato H. High-coverage ITS primers for the DNA-based identification of *Ascomycetes* and *Basidiomycetes* in environmental samples. PLoS One. 2012; 7(7):e40863.

12. Callahan BJ, McMurdie PJ, Rosen MJ, Han AW, Johnson AJ, Holmes SP. DADA2: High-resolution sample inference from Illumina amplicon data. Nat Methods 2016;13:581-583.

13. Callahan BJ, McMurdie PJ, Holmes SP. Exact sequence variants should replace operational taxonomic units in marker-gene data analysis. ISME J 2017;11:2639-2643.

14. Wang J, Zhang Q, Wu G, Zhang C, Zhang M, Zhao L. Minimizing spurious features in 16S rRNA gene amplicon sequencing. *PeerJ Preprints* 2018;6:e26872v1.

15. Katoh K, Misawa K, Kuma K, Miyata T. MAFFT: a novel method for rapid multiple sequence alignment based on fast Fourier transform. Nucleic Acids Res 2002;30:3059-3066.
6. Price MN, Dehal PS, Arkin AP. FastTree 2--approximately maximum-likelihood trees for large alignments. PLoS One 2010;5:e9490.

7. Bokulich NA, Kaehler BD, Rideout JR, Dillon M, Bolyen E, Knight R, Hutten GA, Gregory Caporaso J. Optimizing taxonomic classification of marker-gene amplicon sequences with QIIME 2's q2-feature-classifier plugin. Microbiome 2018;6:90.

8. Quast C, Pruesse E, Yilmaz P, Gerken J, Schweer T, Yarza P, Peplies J, Glöckner FO. The SILVA ribosomal RNA gene database project: improved data processing and web-based tools. Nucleic Acids Res 2013;41:D590-6.

9. Faith DP. Conservation evaluation and phylogenetic diversity. Biological Conservation 1992;61:1-10.

10. Lozupone C, Knight R. UniFrac: a new phylogenetic method for comparing microbial communities. Appl Environ Microbiol 2005;71:8228-8235.

11. Lozupone CA, Hamady M, Kelley ST and Knight R. Quantitative and qualitative beta diversity measures lead to different insights into factors that structure microbial communities. Appl Environ Microbiol 2007;73:1576-1585.

12. Rivers AR, Weber KC, Gardner TG, Liu S, Armstrong SD. ITSxpress: Software to rapidly trim internally transcribed spacer sequences with quality scores for marker gene analysis. F1000Res 2018;7:1418.

13. UNITE Community (2017): UNITE QIIME release. Version 01.12.2017. UNITE Community. https://doi.org/10.15156/BIO/587481.

14. Bray JR, Curtis JT. An ordination of upland forest communities of southern Wisconsin. Ecological Monographs 1957;27:325-349.

15. Kruskal W, Wallis W. Use of ranks in one-criterion variance analysis. J Amer Statist Assoc 1952;47:583-621.

16. Anderson MJ. A new method for non-parametric multivariate analysis of variance. Austral Ecol
Segata N, Izard J, Waldron L, Gevers D, Miropolsky L, Garrett WS, Huttenhower C. Metagenomic biomarker discovery and explanation. Genome Biol 2011;12:R60.

Roy S, Trinchieri G. Microbiota: a key orchestrator of cancer therapy. Nat Rev Cancer 2017; 17:271-285.

Yu G, Gail MH, Consonni D, Carugno M, Humphrys M, Pesatori AC, Caporaso NE, Goedert JJ, Ravel J, Landi MT. Characterizing human lung tissue microbiota and its relationship to epidemiological and clinical features. Genome Biol 2016; 17:163.

Dickson RP, Erb-Downward JR, Martinez FJ, Huffnagle GB. The microbiome and the respiratory tract. Annu Rev Physiol 2016; 78:481-504.

Moffatt MF, Cookson WO. The lung microbiome in health and disease. Clín Med (Lond) 2017; 17:525-52.

Charlson ES, Bittinger K, Haas AR, Fitzgerald AS, Frank I, Yadav A, Bushman FD, Collman RG. Topographical continuity of bacterial populations in the healthy human respiratory tract. Am J Respir Crit Care Med 2011; 184:957-63.

Cameron SJS, Lewis KE, Huws SA, Hegarty MJ, Lewis PD, Pachebat JA, Mur LAJ. A pilot study using metagenomic sequencing of the sputum microbiome suggests potential bacterial biomarkers for lung cancer. PLoS One 2017; 12:e0177062.

Tsai JY, Wu BG, Badri MH, Clemente JC, Shen N, Meyn P, Li Y, Yie TA, Lhakhang T, Olsen E, Murthy V, Michaud G, Sulaiman I, Tsirigos A, Heguy A, Pass H, Weiden MD, Rom WN, Sterman DH, Bonneau R, Blaser MJ, Segal LN. Airway microbiota is associated with up-regulation of the PI3K pathway in lung cancer. Am J Respir Crit Care Med 2018; 198:1188-1198.

Peters BA, Hayes RB, Goparaju C, Reid C, Pass HI, Ahn J. The microbiome in lung cancer tissue and recurrence-free survival. Cancer Epidemiol Biomarkers Prev 2019; 28:731-740.

Yu G, Gail MH, Consonni D, Carugno M, Humphrys M, Pesatori AC, Caporaso NE, Goedert JJ,
Ravel J, Landi MT. Characterizing human lung tissue microbiota and its relationship to epidemiological and clinical features. *Genome Biol* 2016; 17:163.

7. Segal LN, Clemente JC, Tsay JC, Koralov SB, Keller BC, Wu BG, Li Y, Shen N, Ghedin E, Morris A, Diaz P, Huang L, Wikoff WR, Ubeda C, Artacho A, Rom WN, Sterman DH, Collman RG, Blaser MJ, Weiden MD. Enrichment of the lung microbiome with oral taxa is associated with lung inflammation of a Th17 phenotype. *Nat Microbiol* 2016; 1:16031.

8. Hosgood HD 3rd, Sapkota AR, Rothman N, Rohan T, Hu W, Xu J, Vermeulen R, He X, White JR, Wu G, Wei F, Mongodin EF, Lan Q. The potential role of lung microbiota in lung cancer attributed to household coal burning exposures. *Environ Mol Mutagen* 2014; 55:643-51.

9. Kumar R, Herold JL, Schady D, Davis J, Kopetz S, Martinez-Moczygemba M, Murray BE, Han F, Li Y, Callaway E, Chapkin RS, Dashwood WM, Dashwood RH, Berry T, Mackenzie C, Xu Y. *Streptococcus gallolyticus gallolyticus* promotes colorectal tumor development. *PLoS Pathog* 2017; 13:e1006440.

10. Jans C, Boleij A. The road to infection: host-microbe interactions defining the pathogenicity of *Streptococcus bovis/Streptococcus equinus* complex members. *Front Microbiol* 2018; 9:603.

11. Engen SA, Schreurs O, Petersen F, Blix IJS, Baekkevold ES, Schenck K. The regulatory role of the oral commensal *Streptococcus mitis* on human monocytes. *Scand J Immunol* 2018; 87:80-87.

12. Cheng M, Hu S. Lung-resident γδ T cells and their roles in lung diseases. *Immunology* 2017; 151:375-384.

13. Jin C, Lagoudas GK, Zhao C, Bullman S, Bhutkar A, Hu B, Ameh S, Sandel D, Liang XS, Mazzilli S, Whary MT, Meyerson M, Germain R, Blainey PC, Fox JG, Jacks T. Commensal microbiota promote lung cancer development via γδ T cells. *Cell* 2019; 176:998-1013.e16.

14. Iebba V, Santangelo F, Totino V, Nicoletti M, Gagliardi A, De Biase RV, Cucchiara S, Nencioni L, Conte MP, Schippa S. Higher prevalence and abundance of *Bdellovibrio bacteriovorus* in the human gut of healthy subjects. *PLoS One* 2013; 8(4):e61608.
5. de Dios Caballero J, Vida R, Cobo M, Máiz L, Suárez L, Galeano J, Baquero F, Cantón R, del Campo R. Individual patterns of complexity in cystic fibrosis lung microbiota, including predator bacteria, over a 1-year period. *MBio* 2017; 8(5). pii: e00959-17.

6. Morris A, Beck JM, Schloss PD, Campbell TB, Crothers K, Curtis JL, Flores SC, Fontenot AP, Ghedin E, Huang L, Jablonski K, Kleerup E, Lynch SV, Sodergren E, Twigg H, Young VB, Bassis CM, Venkataraman A, Schmidt TM, Weinstock GM; Lung HIV Microbiome Project. Comparison of the respiratory microbiome in healthy nonsmokers and smokers. *Am J Respir Crit Care Med* 2013; 187:1067-1075.

7. Panzer AR, Lynch SV, Langelier C, Christie JD, McCauley K, Nelson M, Cheung CK, Benowitz NL, Cohen MJ, Calfee CS. Lung microbiota is related to smoking status and to development of acute respiratory distress syndrome in critically ill trauma patients. *Am J Respir Crit Care Med* 2018; 197:621-631.

8. Caverly LJ, Huang YJ, Sze MA. Past, present, and future research on the lung microbiome in inflammatory airway disease. *Chest* 2019; 156:376-382.

9. Budden KF, Shukla SD, Rehman SF, Bowerman KL, Keely S, Hugenholtz P, Armstrong-James DPH, Adcock IM, Chotirmall SH, Chung KF, Hansbro PM. Functional effects of the microbiota in chronic respiratory disease. *Lancet Respir Med* 2019; 7:907-920.

10. Pragman AA, Lyu T, Baller JA, Gould TJ, Kelly RF, Reilly CS, Isaacson RE, Wendt CH. The lung tissue microbiota of mild and moderate chronic obstructive pulmonary disease. *Microbiome* 2018; 6:7.

11. de Torres JP, Marin JM, Casanova C, Cote C, Carrizo S, Cordoba-Lanus E, Baz-Dávila R, Zulueta JJ, Aguirre-Jaime A, Saetta M, Cosio MG, Celli BR. Lung cancer in patients with COPD: incidence and predicting factors. *Am J Respir Crit Care Med* 2011; 184:913-919.

12. Krause R, Moissl-Eichinger C, Halwachs B, Gorkiewicz G, Berg G, Valentin T, Prattes J, Högenauer C, Zollner-Schwetz I. Mycobiome in the lower respiratory tract - a clinical
Alpha diversity. Analysis by sample location using Chao1, Shannon, and Faith’s PD indexes. Significant differences (p<.05) between groups are highlighted by asterisks.
Bacterial phyla distribution in each type of sample. The main result from this figure is the high abundance of Firmicutes in all samples from lung cancer patients, combined with lower proportions of Proteobacteria.
Figure 3
Salivary microbiota of patients and controls. A: LEfSe analysis discriminating the statistical differences between the genera abundance. To note that only >4 were LDA scores considered to highlight the significant differences. B: Salivary microbiota core, and gut microbiota core only from lung cancer patients, including only the majoritarian genera detected in all patients. The abundance of each genus is represented by the circle size and the numbers of connections were determined by their frequency in each patient. C: UniFrac beta diversity analysis of salivary samples. Unweighted UniFrac reports differences in the presence or absences of ASVs, while weighted UniFrac also reports differences in the abundance of ASVs. D: Principal Component Analysis (PCA) of the most abundant genera that constitute 90% of the total salivary microbiota.
Figure 4

Representation and comparison of the microbiota from cancer affected, contralateral and control bronchi. A: LEfSe analysis of the bronchial microbiota between patients, represented by the affected bronchi, and healthy controls. Only taxa with an LDA score >4 are shown. B: Complete and core lung microbiota from controls and lung cancer patients. The abundance of each genus is represented by the circle size and the numbers of connections were determined by their frequency in each patient. C: UniFrac beta diversity analysis of bronchi samples. Unweighted UniFrac reports differences in the presence or absences of ASVs, while weighted UniFrac also reports differences in the abundance of ASVs. D: PCA of most abundant genera that constituted 90% of the bronchial microbiota of both patients and controls.
Saliva and lung microbiota relatedness. Weighted UniFrac distance calculated between saliva and lung microbiota in control group (left) and in patients’ group (right). To notice that distance is significantly higher in control group than in patient group (p<.001), pointing to a major connection of both ecosystems in patients than in controls.
Figure 6

Streptococcal abundance. A: Heatmap correlating the abundance of Streptococcal ASVs (horizontal axis) with the samples (vertical axis). A total of 95 ASVs were considered. B: Streptococcal abundance in the different samples. C: ROC curves of the relative abundance of the Streptococcus genera to discriminate patients from controls.
Figure 7

Mycobiome results. A: LEfSe analysis of the affected bronchi from patients and the bronchi from controls. To note that only LDA scores >4 were considered in order to highlight top significant differences. B: PCoA based on Bray Curtis distances, showing the distribution of samples according to their mycobiome. C: Alpha diversity of the mycobiome detected in the different samples. Chao1 and Shannon metrics are shown separately. Significant differences (p<.05) between patients and controls are highlighted by asterisks.