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Original Article

Intestinal function and transit associate with gut microbiota dysbiosis in cystic fibrosis

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ABSTRACT

Background: Most people with cystic fibrosis (pwCF) suffer from gastrointestinal symptoms and are at risk of gut complications. Gut microbiota dysbiosis is apparent within the CF population across all age groups, with evidence linking dysbiosis to intestinal inflammation and other markers of health. This pilot study aimed to investigate the potential relationships between the gut microbiota and gastrointestinal physiology, transit, and health.

Study design: Faecal samples from 10 pwCF and matched controls were subject to 16S rRNA sequencing. Results were combined with clinical metadata and MRI metrics of gut function to investigate relationships.

Results: pwCF had significantly reduced microbiota diversity compared to controls. Microbiota compositions were significantly different, suggesting remodelling of core and rarer satellite taxa in CF. Dissimilarity between groups was driven by a variety of taxa, including *Escherichia coli, Bacteroides spp., Clostridium spp.,* and *Faecalibacterium prausnitzii.* The core taxa were explained primarily by CF disease, whilst the satellite taxa were associated with pulmonary antibiotic usage, CF disease, and gut function metrics. Species-specific ordination biplots revealed relationships between taxa and the clinical or MRI-based variables observed.

Conclusions: Alterations in gut function and transit resultant of CF disease are associated with the gut microbiota composition, notably the satellite taxa. Delayed transit in the small intestine might allow for the expansion of satellite taxa resulting in potential downstream consequences for core community function in the colon.

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1. Introduction

Cystic fibrosis (CF) associated respiratory infections are the major cause of disease morbidity and mortality. However, a number of gastrointestinal (GI) problems may also arise, limiting the quality of life, including meconium ileus at birth, distal intestinal obstruction syndrome, small intestinal bacterial overgrowth (SIBO),

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increased risk of malignancy, and intestinal inflammation [1,2]. It is therefore unsurprising that people with CF experience persistent GI symptoms [3,4] with "how can we relieve gastrointestinal symptoms in people with CF?" a top priority question for research [5].

Microbial dysbiosis at the site of the GI tract in CF patients has been described, with changes evident from birth through to adulthood [6–10]. Moreover, the extent of this divergence from healthy microbiota, initially due to loss of cystic fibrosis transmembrane conductance regulator (CFTR) function [11], is further compounded by routine treatment with broad spectrum antibiotics [10]. The reshaping of the gut microbiota may have functional consequences that could further impact on patients. These include the reduction

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Table 1				
Clincial	characteristics	of study	participants.	

Study I.D	Sex	Age (Years)	Group	Pancreatic Status	Calprotectin (µg/g)	FEV1%	BMI	Antibiotic Usage				
								Р	А	М	β	S
365	М	12	CF	PI	4.22	87	16.18	-	-	-	+	_
431	М	12	HC	PS	2.44	-	17.95	-	-	-	-	-
128	Μ	13	CF	PI	27.59	97	17.72	+	-	-	-	-
296*	М	13	HC	PS	-	-	23.44	-	-	-	-	-
643	М	13	CF	PI	9.77	90	21.83	-	-	+	-	-
159	М	13	HC	PS	2.72	-	23.49	-	-	-	-	-
297	М	15	CF	PI	27.61	126	20.83	-	-	-	-	-
947*	М	15	HC	PS	-	-	20.94	-	-	-	-	-
617	F	15	CF	PI	21.15	72	18.42	-	-	+	+	+
964	F	15	HC	PS	12.71	-	19.15	-	-	-	-	-
167	М	19	CF	PI	7.37	99	20.63	-	-	-	-	-
673	М	19	HC	PS	0.94	-	20.34	-	-	-	-	-
279	F	19	CF	PI	27.32	66	20.87	-	-	+	-	-
205	F	19	HC	PS	3.84	-	31.91	-	-	-	-	-
596	F	21	CF	PI	14.05	61	21.91	-	-	+	-	-
152	F	21	HC	PS	4.22	-	21.26	-	-	-	-	-
610*	М	23	CF	PI	-	66	18.64	-	+	+	-	-
548	М	24	HC	PS	3.56	-	24.49	-	-	-	-	-
619*	F	27	CF	PI	-	60	19.27	-	-	-	-	-
501	F	27	HC	PS	7.19	-	28.66	-	-	-	-	-
259	М	30	CF	PI	28.30	61	20.21	-	-	+	+	-
986	М	29	HC	PS	4.96	-	22.64	-	-	-	-	-
681	F	36	CF	PI	11.79	88	21.71	+	-	+	-	-
749	F	35	HC	PS	3.00	-	19.57	-	-	-	-	-

Subjects marked with an asterisk* indicate those who failed to produce a stool sample for subsequent metagenomic and metabolomic analysis and thus were excluded from downstream analyses. All participants with CF had the gene mutation p.Phe508del/p.Phe508del, with pancreatic insufficiency but no CF-related diabetes. For antibiotic usage, '+' indicates routine administration of the given antibiotic class prior to sampling. Abbreviations: FEV1 – Percent predicted forced expiratory volume in 1 second, BMI – Body mass index, P – Polymyxin, A – Aminoglycoside, M – Macrolide, $\beta - \beta$ -lactam, S – Sulfonamide. Asterisks denote participants who did not provide any stool samples upon visitation, and thus were excluded from downstream microbiota analysis.

of taxa associated with the production of short-chain fatty acids (SCFAs) which play key roles in modulating local inflammatory responses and promoting gut epithelial barrier integrity [12–14]. Furthermore, studies of microbiota dysbiosis in CF have demonstrated its relationship with intestinal inflammation, intestinal lesions, and increased gene expression relating to intestinal cancers [15–18]. Whilst many of these clinical parameters have ties to gut microbiota changes, they remain understudied exclusively past childhood despite advances in less invasive approaches to investigate CF gut physiology and function [19]. Our group has recently published on the use of magnetic resonance imaging (MRI) to assess gut transit time, along with other parameters, in adolescents and adults [20].

In this pilot study, we linked those MRI physiology metrics and clinical metadata directly to high-throughput amplicon sequencing data identifying constituent members of the gut microbiota, to explore the relationships between microbial dysbiosis, intestinal function and clinical state.

2. Materials and methods

2.1. Study participants and design

Twelve people with CF, homozygous for p.Phe508del along with 12 healthy controls, matched by age and gender, were recruited from Nottingham University Hospitals NHS Trust. Participants were asked to provide stool samples when attending for MRI scanning, with the study design and MRI protocols described previously [20]. A patient clinical features were also recorded upon visitation (Table 1), including a three-day food diary preceding sample collection (Table S1). Further descriptive statistics of the study population can be found in the Supplementary Materials, including MRI metrics (Table S2), and summary statistics on diet (Table S3-S6). Faecal samples were only obtained from ten individuals in each group. Written informed consent, or parental consent and as-

sent for paediatric participants, was obtained from all participants. Study approval was obtained from the West Midlands Coventry and Warwickshire Research Ethics Committee (18/WM/0242). All stool samples obtained were immediately stored at -80 °C prior to DNA extraction to reduce changes before downstream community analysis [21].

2.2. Targeted amplicon sequencing

DNA from dead or damaged cells, as well as extracellular DNA was excluded from analysis via cross-linking with propidium monoazide (PMA) prior to DNA extraction, as previously described [22]. Next, cellular pellets resuspended in PBS were loaded into the ZYMO Quick-DNA fecal/Soil Microbe Miniprep Kit (Cambridge Bioscience, Cambridge, UK) as per manufacturer's instructions, with the following amendments: ZR BashingBead Lysis Tubes were replaced with standard 1.5 mL Eppendorf tubes loaded with ZYMO Beads for mechanical homogenisation with the use of a Retsch Mixer Mill MM 400 (Retsch, Haan, Germany). Samples were homogenised for 2 min at 17.5/s frequency. Following DNA extraction, approximately 20 ng of template DNA was then amplified using Q5 high-fidelity DNA polymerase (New England Biolabs, Hitchin, UK) using a paired-end sequencing approach targeting the bacterial 16S rRNA gene region (V4-V5). Primers and PCR conditions can be found in the Supplementary Materials. Pooled barcoded amplicon libraries were sequenced on the Illumina MiSeq platform (V3 Chemistry).

2.3. Sequence processing and analysis

Sequence processing and data analysis were initially carried out in R (Version 4.0.1), utilising the package DADA2 [23]. The full protocol is detailed in the Supplementary Materials. Raw sequence data reported in this study has been deposited in the European Nucleotide Archive under the study accession number PRJEB44071.

2.4. Faecal calprotectin

Stool was extracted for downstream assays using the ScheBo® Master Quick-Prep (ScheBo Biotech, Giessen, Germany), according to the manufacturer instructions. Faecal calprotectin was analysed using the Bühlmann fCAL ELISA (Bühlmann Laboratories Aktienge-sellschaft, Schonenbuch, Switzerland), according to the manufacturer's protocol.

2.5. Statistical analysis

Regression analysis, including calculated coefficients of determination (r^2), degrees of freedom (df), *F*-statistic and significance values (*P*) were calculated using XLSTAT v2021.1.1 (Addinsoft, Paris, France). Fisher's alpha index of diversity and the Bray-Curtis index of similarity were calculated using PAST v3.21 [24]. Significant differences in microbiota diversity were determined using Kruskal-Wallis performed using XLSTAT. Analysis of similarities (ANOSIM) with Bonferroni correction was used to test for significance in microbiota composition and was performed in PAST. Similarity of percentages (SIMPER) analysis, to determine which taxa contributed most to compositional differences between groups, was performed in PAST.

Redundancy analysis (RDA), was performed in CANOCO v5 [25]. Following the determination of clinical variables significantly explanatory for microbiome composition, RDA biplots with these variables were plotted in PAST v3.21. Statistical significance for all tests was deemed at the $p \le 0.05$ level. Supplementary information, including metadata, are available at figshare.com under https://doi.org/10.6084/m9.figshare.15073797.v1 and https://doi.org/10.6084/m9.figshare.15073899.v1.

3. Results

To investigate the contributions of common and rare bacterial taxa in the gut microbiota of individuals within and between study cohorts [26, 27], taxa were partitioned into either common and abundant core taxa or rarer and infrequent satellite taxa, based upon their prevalence and relative abundance across samples within each cohort (Fig. 1). Within the healthy control group, 30 taxa were core constituting 60.5% of the total abundance, with the remainder accounted for by 386 satellite taxa. In the CF group, 22 core taxa represented 34.7% of the abundance, with 323 satellite taxa constituting the remainder. Core taxa are listed in Table S7. The whole, core, and satellite microbiota demonstrated similar patterns in diversity, whereby there was significantly reduced diversity in the CF group (Fig. 2A, Table S8).

Within-group core microbiota similarity was higher within the healthy control group, with a mean similarity (\pm SD) of 0.60 \pm 0.08 compared to 0.40 \pm 0.11 for the CF group (Fig. 2B). As expected, satellite taxa similarity within groups was much lower than for the core but was also significantly reduced in CF compared to controls, at 0.35 \pm 0.08 and 0.21 \pm 0.09 for the healthy control and CF group respectively. ANOSIM testing determined the whole microbiota, core, and satellite taxa of the CF group were significantly different in composition compared to healthy controls (Fig. 2B, Table S9). SIMPER analysis was implemented to reveal which taxa were responsible for driving this dissimilarity (Table 2). Of the taxa contributing to > 50% of the differences between healthy control and CF groups, those within the genus Bacteroides were represented most. Escherichia coli contributed most towards the differences between groups, despite satellite status, followed by Bacteroides sp. (OTU 3), Clostridium sp. (OTU 5), Faecalibacterium prausnitzii, and Bacteroides fragilis.

Redundancy analysis (RDA) was used to relate variability in microbiota composition to associated MRI metrics and clinical fac-



Fig. 1. Distribution and abundance of bacterial taxa across different sample groups. (A) Healthy control. (B) Cystic fibrosis. Given is the percentage number of patient stool samples each bacterial taxon was observed to be distributed across, plotted against the mean percentage abundance across those samples. Core taxa are defined as those that fall within the upper quartile of distribution (orange circles), and satellite taxa (grey circles) defined as those that do not, separated by the vertical line at 75% distribution and labelled respectively. Distribution-abundance relationship regression statistics: (a) $r^2 = 0.50$, $F_{1,414} = 407.3$, P < 0.0001; (b) $r^2 = 0.29$, $F_{1,343} = 137.3$, P < 0.0001. Core taxa are listed in Table S7.

tors (Table 3). Pulmonary antibiotics and CF disease significantly explained the most variance across the whole and satellite microbiota. Measurements of intestinal transit and function contributed to the whole microbiota variance, albeit to a lesser extent, with variation in OCTT and SWBC also contributing to satellite taxa variance alongside faecal calprotectin levels. In the core taxa analysis, the presence of CF disease was the dominant factor in significantly explaining the compositional variability, followed by sex and body mass index (BMI).

A species redundancy analysis biplot (RDA) was constructed to investigate how significant clinical variables from the whole microbiota direct ordination approach explained the relative abundance of taxa from the SIMPER analysis (Fig. 3). Certain taxa grouped away from many of the significant clinical variables shown in a similar manner. This effect was most pronounced for *F. prausnitzii, Eubacterium rectale* and *Ruminoccocus bromii*. A combination of clinical factors, including CF disease, increased fasting colonic volume, increased SBWC and prolonged OCTT, explained the variance observed in relative *E. coli* abundance, whilst a more modest effect was observed towards *Streptococcus* sp. (OTU 18), *Dialister invisus, Clostridium perfringens* and *Romboutsia timonensis*. Species of *Bacteroides*, which was the most common genus within the topcontributing SIMPER analysis, were explained by the clinical variables to high variability. JID: JCF

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Fig. 2. Microbiome diversity and similarity compared across healthy controls and cystic fibrosis samples. Whole microbiota (black plots) and partitioned data into core (orange plots) and satellite taxa (grey plots) are given. (**A**) Differences in Fisher's alpha index of diversity between healthy controls and cystic fibrosis samples. Black circles indicate individual patient data. Error bars represent 1.5 times inter-quartile range (IQR). Asterisks between groups denote a significant difference in diversity following use of Kruskal-Wallis tests (P < 0.001). Summary statistics are provided in Table S8. (**B**) Microbiome variation measured within and between sampling groups, utilising the Bray-Curtis index of similarity. Error bars represent standard deviation of the mean. Asterisks indicate significant differences between sampling groups following the use of one-way ANOSIM testing (P < 0.001). Summary statistics are provided in Table S9.

4. Discussion

In this pilot study, we investigated the relationships between clinical factors, MRI markers of GI function and the composition of faecal bacterial microbiota. Demonstrated previously for CF lung and Crohn's gut microbiota [26,27], we have shown here that it is possible to partition the CF gut microbiota into core and satellite taxa to investigate potential community functions and relationships, with the notion that the core constituents contribute to the majority of functionality exhibited by the community [22,26]. As to be expected, the core taxa made up most of the abundance within the healthy control group. Whilst many taxa were also commonly represented in the CF group, the latter was dominated in abundance by the satellite taxa. Our findings of reduced diversity across the whole, core, and satellite microbiota are in agreement with previous findings described within the CF gut [7,8,10]. Along with reduced within group similarity in CF compared to healthy controls across all microbiota partitions, this suggests a perturbed community harbouring greater instability, less subsequent resilience, and inherent challenges to the colonisation and establishment of normal commensals. CF associated factors such as varied antibiotic usage will contribute to this reduced similarity, further augmented by the wide age range of pwCF within this study and variation across lifestyle factors. The combination of the aforementioned may elicit stochastic community disruption and increased inter-individual variation as observed across other mammalian microbiomes [28].

At the surface, a reduction in the number of taxa labelled as core within the CF group hinted at perturbation and restructuring, further evidenced by the occurrence of taxa exclusively core to this group. This included species of *Streptococcus*. *Pseudomonas*. Veillonella, and Enterococcus, all of which were significantly more abundant in the CF group (Table S7), and of which are implicated in both CF lung and gut microbiomes [8,11,18,26,29,30]. The concept of the "gut-lung axis" in CF arises from the direct translocation of the respiratory microbiota from sputum swallowing to the gut [31], but also the emergence of species in the gut prior to the respiratory environment [30]. This apparent bidirectionality is further supported by the administration of oral probiotics to decrease pulmonary exacerbations in CF [32]. Aside from sputum swallowing, the increase in Streptococcus and Veillonella here could reflect an increased availability of simple carbohydrates from the observed dysmotility of the gut [20]. Streptococci are well equipped with numerous genes for rapid carbohydrate degradation in an environment usually fluctuating in substrate availability, with fermentation-derived lactic acid supporting the expansion of Veil*lonella* species in the small intestine [33].

E. coli contributed most to the dissimilarity between healthy and CF groups despite maintaining satellite status throughout both the healthy and CF groups, seemingly resultant of the wide age range of our study participants, of which the higher relative abundances were observed in the younger adolescent patients (Table 2). In childhood studies, a significantly higher relative abundance of *Proteobacteria* is often reported in relation to dysbiosis, with *E. coli* abundance associating with poor growth outcomes and intestinal inflammation [34–36]. Other notable taxa contributing to the dissimilarity observed between groups encompassed a variety of key species associated with SCFA production in the colon. This included

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Table 2

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Similarity of percentage (SIMPER) analysis of microbiota dissimilarity (Bray-Curtis) between Healthy Control (HC) and Cystic Fibrosis (CF) stool samples.

	%Relative abundance					
Таха	Mean HC Mean CF		Av. Dissimilarity	% Contribution	Cumulative %	
Escherichia coli	1.84	9.54	4.72	6.39	6.39	
Bacteroides 3	3.84	4.69	3.36	4.55	10.94	
Clostridium 5	0.77	6.44	3.09	4.18	15.13	
Faecalibacterium prausnitzii	8.56	2.95	2.99	4.05	19.18	
Bacteroides fragilis	1.02	5.29	2.75	3.73	22.90	
Bacteroides dorei	3 <u>.</u> 32	4.31	2.52	3.42	26.32	
Eubacterium rectale	5.03	1.35	2.18	2.95	29.27	
Romboutsia timonensis	1.24	3.95	2.15	2.91	32.18	
Bacteroides uniformis	2.72	4.09	1.62	2.20	34.38	
Dialister invisus	1.00	3.45	1.61	2.19	36.57	
Bacteroides vulgatus	2 <u>.</u> 37	2.14	1.56	2.11	38.68	
Ruminococcus bromii	2.69	0.42	1.24	1.68	40.36	
Alistipes putredinis	2.08	0.06	1.02	1.38	41.74	
Bacteroides coprocola	1.56	0.92	1.01	1.37	43.11	
Fusicatenibacter saccharivorans	2.62	0.8	1.00	1.36	44.47	
Streptococcus 18	0.26	1.95	0.88	1.19	45.66	
Blautia luti	2.93	2.31	0.86	1.16	46.82	
Oscillibacter ruminantium	1.90	0.27	0.84	1.14	47.96	
Clostridium perfringens	0.00	1.58	0.79	1.07	49.03	
Parabacteroides distasonis	1.38	1.85	0.77	1.05	50.08	

Taxa identified as core are highlighted in orange, whereas satellite taxa are highlighted in grey. Mean relative abundance (%) is also provided for each group. Percentage contribution is the mean contribution divided by the mean dissimilarity across samples (73.79%). Cumulative percent does not equal 100% as the list is not exhaustive. Given the sequencing length of 16S gene regions, taxon identification should be considered putative.

Table 3

Redundancy analysis to explain percent variation in whole microbiota, core taxa and satellite taxa between all subjects from significant clinical variables measured.

	Microbiota			Core taxa			Satellite taxa			
	Var. Exp (%)	pseudo-F	P (adj)	Var. Exp (%)	pseudo-F	P (adj)	Var. Exp (%)	pseudo-F	P (adj)	
Antibiotics	21.5	5.4	0.002				27.1	7.3	0.002	
BMI				7.0	2.0	0.042				
Calprotectin							5.9	1.8	0.050	
CF Disease	10.9	2.2	0.002	28.9	7.3	0.002	10.3	2.1	0.006	
Colon Fasting Vol.	7.5	2.0	0.016							
OCTT	7.4	2.1	0.012				6.7	1.9	0.046	
SBWC	5.6	1.7	0.048				7.2	2.4	0.048	
Sex				7.9	2.1	0.010				
Total	52.9			43.8			57.2			

Var. Exp (%) represents the percentage of the microbiota variation explained by a given parameter within the redundancy analysis model. P(adj) is the adjusted significance value following false discovery rate correction. Antibiotics is the presence/absence of recurrent antibiotic regimes for a given patient. BMI – Body mass index, Colon Fasting Vol – Colon volume at baseline corrected for body surface area, OCTT – Oro-caecal transit time, Antibiotics, SBWC – Small bowel water content corrected for body surface area.

F. prausnitizii and *E. rectale*, both of which were significantly decreased in abundance within the CF group, but also *R. bromii* and *B. luti*. These taxa have all been previously reported to decrease in the CF gut [8,29,37] alongside other inflammatory conditions [38]. There were also notable contributions to the dissimilarity between groups by *Clostridium* sp. (OTU 5) (significant difference in relative abundance) and *D. invisus* (not significant). *Clostridium* OTU 5 aligned exclusively with cluster I members at the 97% threshold, of whom demonstrate the capacity to generate lactate, acetate, propionate, and butyrate via carbohydrate fermentation [39], whilst *D. invisus* is an intermediary fermenter capable of both acetate

and propionate production. This may lend support to the theory that alternate species can retain some functional redundancy in the presence of perturbation to the local community in the CF gut [40].

Variance across the whole microbiota and satellite taxa was significantly explained by the use of antibiotics (Table 3), of which most pwCF are administered on a routine basis to supress lung infection [41]. The occurrence of both OCTT and SBWC accounting for significant explanation in both the whole microbiota and satellite, but not core taxa analysis, underpins the strong impact of gut physiology and transit on the microbiota in CF. Faecal calprotectin also explained the variance across the satellite taxa, and has been



Fig. 3. Redundancy analysis species biplots for whole microbiota. The 20 taxa contributing most to the dissimilarity (cumulatively > 50%) between healthy and cystic fibrosis groups from the SIMPER analysis (Table 2) are shown independently of the total number of ASVs identified (345). Orange circles represent core taxa within the CF group, whilst grey circles denote satellite taxa. Biplot lines depict clinical variables that significantly account for the total variation in taxa relative abundance within the whole microbiota analysis at the $p \le 0.05$ level as seen in Table 3, with species plots indicating the strength of explanation provided by the given clinical variables. 'OCT' – Oro-caecal transit time, Antibiotics, 'SBWC' – Small bowel water content corrected for body surface area, Colon Fasting Volume corrected for body surface area, CF disease. For example, biplot lines depicting the MRI metrics represent how changes in the metric influences the relative abundance of each of the taxa shown. Those taxa shown in the same direction of the metric label are considered to have a higher value than those taxa that are not. The percentage of microbiome variation explained by each axis is given in parentheses.

associated with increased abundances of *Escherichia*, *Streptococcus*, *Staphylococcus* and *Veillonella*, of which contained satellite species significantly increased in our CF group [29,42]. *Acidaminococcus* sp. have also associated with increased faecal calprotectin levels [18], with *Acidaminococcus intestinii* another constituent of the CF satellite microbiota that was not present in healthy controls (data not shown). The core taxa was only largely explained by the presence of CF disease itself, perhaps relating to the direct disruption of CFTR function which alone can influence changes in the microbiome [42].

Perhaps unsurprisingly, the species ordination biplots of the taxa from SIMPER analysis demonstrated clustering of the key SCFA producers mentioned previously away from the significant disease-associated clinical factors, with antibiotic usage and transit metrics previously shown to reduce the abundance of such taxa [15,43]. Similarly affected were taxa from genera that are associated with better outcomes in other similarly pro-inflammatory intestinal environments, such as Crohn's disease or ulcerative colitis, including *Oscillibacter and Fusicanterbacter* [44,45].

C. perfringens has been associated with disease exacerbation in ulcerative colitis [38], SIBO in the CF mouse small intestine [46] and increased deconjugation of bile salts leading to further fat malabsorption by the host [47]. Here it was completely absent from our healthy control group, whilst in the CF group was found to associate with a variety of CF-induced clinical factors as well as OCTT. Also strongly associating with OCTT and impacted substantially more, was *E. coli*. Increased bacterial load relates to slower transit within the CF mouse small intestine [47]. Concurrently with the observed increase in SWBC reported prior [20], this in theory allows for the expansion of such facultative anaerobes in the small intestine that could potentially affect downstream community dynamics and functional profiles in the colon, given that PMA treatment was utilised to select for viable living taxa from faecal sampling.

Although dietary profiles were similar between groups (Tables S3–6) and did not contribute to significant variation in the microbiota, increased fat intake to meet energy requirements is a staple of the CF diet [48]. The infant gut metagenome demonstrates enrichment of fatty acid degradation genes [34] whilst CF-derived *E. coli* strains exhibit improved utilisation of exogenous glycerol as a growth source [49]. Finally, the genus *Bacteroides*, which has been reported to both increase and decrease within CF disease across different age groups [8,11,15], displayed high variability within the species ordination biplot (Fig. 3), perhaps resultant of the varying antimicrobial susceptibility within the genus [50].

We acknowledge the small sample size of this pilot study limits the power of specific analyses, with the absence of withingroup direct ordination approaches which would have allowed for investigation of CF group antibiotic usage and extra clinical factors such as lung function. However, the principle strength of this JID: JCF

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study is the valuable insight into the relationships between microbiota composition and intestinal physiology and function in CF. Future studies should encompass larger cohorts in a longitudinal fashion with the combination of both lung and faecal microbiota data to elucidate such relationships better, including the impact of pulmonary antibiotic usage on the gut microbiota, and the aptly termed gut-lung axis. Evaluation of associations between the microbiota, physiology and the immune response would also improve our understanding of the mechanisms contributing to GI health in CF. Given their possible beneficial effect on intestinal inflammation [51], the impact of CFTR modulator therapy will provide further insights.

5. Conclusion

This cross-sectional pilot study has identified relationships between markers of clinical status, gastrointestinal function and bacterial dysbiosis in the CF population. By partitioning the community into core and satellite taxa, we were able to reveal the relative contributions of CF-associated lifestyle factors and elements of intestinal function to these subcommunity compositions, and how specific taxa were affected by these clinical factors. Further, as the first study to combine high-throughput gene amplicon sequencing with non-invasive MRI to assess underlying gut pathologies, we demonstrate the potential for future collaborations between gastroenterology and microbiology with larger cohort recruitment to investigate these relationships between gut function and the microbiome further.

Author contributions

CvdG, AS, GM, and RJM conceived the study. RJM, HG, LH, and MMW performed sample processing and analysis. RJM, DR, and CvdG performed the data and statistical analysis. CN, GM, and AS were responsible for sample collection, clinical care records and documentation. RJM, CN, GM and CvdG verified the underlying data. RJM, DR, and CvdG were responsible for the creation of the original draft of the manuscript. RJM, CN, GM, DR, AS, and CvdG contributed to the development of the final manuscript. CvdG is the guarantor of this work. All authors read and approved the final manuscript.

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Declaration of Competing Interest

RJM, HG, LH, DM, and CvdG declare support from the CF Trust. CN and GM report grants and speaker honorarium from Vertex, outside the submitted work. ARS reports grants from Vertex, as well as speaker honoraria and expenses from Teva and Novartis and personal fees from Vertex, outside the submitted work. In addition, ARS has a patent issued "Alkyl quinolones as biomarkers of *Pseudomonas aeruginosa* infection and uses thereof".

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.jcf.2021.11.014.

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