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RESEARCH PAPER



The role of inflammation in temporal shifts in the inflammatory bowel disease mucosal microbiome

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ABSTRACT

Studies of the human intestinal microbiome in patients with inflammatory bowel disease (IBD) consistently show that there are differences (an abnormal or unbalanced microbiome, “dysbiosis”) when compared to healthy subjects. We sought to describe changes in the microbiome in individual patients over time, and determine the clinical factors that are associated with significant alteration. Forty-two mucosal biopsies were collected from 20 patients that were spaced an average of 2.4 years apart. These were analysed using bacterial 16S rRNA gene high-throughput sequencing methods. Presence of active inflammation was determined endoscopically and histologically. Inferred metagenomics analysis was conducted using the PICRUSt package. We found that the differences in the microbiome over time in individual patients were greatest in the presence of ongoing intestinal inflammation, as determined by the Yue and Clayton theta distance between sample pairs (adjusted $p = 0.00031$). Samples from patients with previous abdominal surgery had lower alpha (within sample) diversity compared with those with no prior operations (mean Shannon index 2.083, 2.510 respectively, $p = 0.017$). There were no changes in the inferred bacterial metagenomic profile. The microbiome in IBD undergoes considerable fluctuation over time. These changes are greatest when there is histologically confirmed inflammation at both time-points.

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IBD; intestinal microbiology; inflammation; microbiome; Crohn's disease; Ulcerative colitis

Introduction

The intestinal microbiome of patients with inflammatory bowel diseases (IBD; Crohn's disease (CD) and ulcerative colitis (UC)) differs significantly from normal controls.^{1–3} As no single microorganism has been identified as the cause of IBD, the focus of current research has shifted to studying the characteristic “dysbiosis”^{4–6} (an abnormal or unbalanced microbiome) and developing therapies to correct it.^{7,8} This is based on the assumption that it is the dysbiosis that leads to the uncontrolled inflammation characteristic of IBD, either directly, or by interfering with mucosal protective mechanisms.

In general, a reduction in bacterial diversity is seen in patients with active IBD.⁹ Of the four main bacterial phyla found in the gut, Bacteroidetes, Firmicutes, Actinobacteria and Proteobacteria, increases in Bacteroidetes and a reduction in Firmicutes are commonly associated

with disease. Changes to the abundance of specific organisms have also been associated with IBD. For example, a reduction in abundance of the butyrate-producing bacterium, *Faecalibacterium prausnitzii* is commonly seen in the setting of active inflammation,^{10,11} whereas the abundance of *Escherichia coli* is commonly increased in active IBD.

While the human gut microbiome has been shown to maintain a certain degree of stability over time,^{12,13} several factors are associated with disruption of its diversity and species richness. For example, perturbations in the intestinal microbiome follow gastrointestinal infection, antibiotic use and smoking.^{14–16}

The Human Microbiome Project, which analysed the oral, gastrointestinal, skin and vaginal microbiomes of 242 healthy adults at two time points,¹⁷ demonstrated significant inter-individual differences. Despite this, most published studies focusing on the

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IBD microbiome obtained samples from a single time-point, and compared microbial communities between subjects. In addition, most studies used faecal samples to characterize the intestinal microbiome and to determine whether there was active inflammation (faecal calprotectin). Faecal specimens do not necessarily reflect the nature of the mucosa-adherent bacteria,¹⁸ nor the degree of inflammation.¹⁹ While the use of faecal calprotectin measurements is useful in the setting of screening for gastrointestinal inflammation,²⁰ it is not as accurate as endoscopic or histological assessment in patients with IBD, especially those with small bowel inflammation.^{21,22}

We analysed temporal changes in the IBD microbiome using the best methods available: mucosal samples were used to characterize the microbiome, and intestinal histology to determine whether or not there was active inflammation. Our aims were to describe the fluctuations of the microbiome of patients with established or newly diagnosed IBD over time, with a view to identify clinical and microbial factors that may be associated with remission or the development of disease activity at a later time-point.

Results

Forty-two temporally spaced biopsy or resection specimens were collected from 20 patients with IBD (16 CD, 4 UC) between 2004 and 2014 at Canberra Hospital. Specimens were collected from 19 patients who had undergone two procedures, and one patient who had undergone four separate procedures during the collection period. The mean time between procedures was 2.4 years (\pm standard deviation (SD) 1.7 years). The majority of samples were collected at time of colonoscopy (34 samples, representing 81% of the total), and eight were collected at time of intestinal resection. Inflammatory status was determined by the proceduralist (endoscopy or operation report, 42 of 42 available), and histopathology report (36 of 42 available). Overall, 60% of patient samples had active inflammation at all time-points. Table 1 describes the clinical details of the patients and samples in greater detail (see also Supplementary Table 1).

Samples from patients with previous abdominal surgery had lower alpha diversity compared with those with no prior operations (mean Shannon index 2.083, 2.510 respectively, $p = 0.017$), a finding shared

Table 1. Patient demographic data, Montreal disease phenotype classification and medication exposure. Active inflammation defined as disease activity reported endoscopically or histologically. Other medications included sirolimus and abatacept.

Number of individual patients	20
Biopsy samples, total	42
Collected at colonoscopy (%)	34 (81)
Collected at bowel resection (%)	8 (19)
Samples taken in presence of any active inflammation (% of samples)	37 (88)
Endoscopic inflammation (% of samples)	28 (67)
Histologic inflammation (% of those with result available, n = 36)	30 (71)
Active inflammation at all time-points (% of patients)	12 (60)
Active initially, remission at follow-up (% of patients)	3 (15)
Disease in remission at all time-points (% of patients)	0 (0)
Remission initially, active inflammation at follow-up (% of patients)	4 (20)
Samples taken in disease remission (% of samples)	
Endoscopic remission (% of samples)	14 (33)
Histologic remission (% of those with result available, n = 36)	4 (11)
Age at time of procedure, years (SD)	41 (11)
Active smoker at time of procedure (% of patients)	6 (30)
Duration of disease since diagnosis, years (SD)	13.6 (12.6)
Range (years)	0–42
Time between samples, mean years (SD)	2.4 (1.7)
Range (years)	0.5 – 5.9
Number of patients with Crohn's disease (% of patients)	16 (80)
Crohn's disease location classification (%)	
L1 (ileum only)	4 (25)
L2 (colon only)	2 (13)
L3 (ileum and colon)	9 (56)
L4 (upper gastrointestinal tract)	2 (13)
Crohn's disease phenotype classification (%)	
B1 (inflammatory)	5 (31)
B2 (stricturing)	9 (56)
B3 (penetrating)	2 (13)
p (perianal involvement)	5 (31)
Number of patients with ulcerative colitis (% of patients)	4 (20)
Ulcerative colitis disease extent classification (%)	
E1 (distal)	0
E2 (left sided)	1 (25%)
E3 (pan colitis)	3 (75%)
Number of previous surgeries (% of samples)	
0	26 (62)
1	11 (26)
2	1 (4)
3	4 (10)
Medications being taken at time of procedure (% of samples)	
No medications	12 (29)
Antibiotics	3 (7)
Mesalazine	4 (10)
Thiopurines	15 (36)
Methotrexate	2 (5)
Anti-TNF α	10 (24)
Prednisone	3 (7)
Other medications	3 (35)

by other studies.²³ Within the other clinical subgroups there were no significant differences in alpha diversity (Supplementary Figure 1 and Supplementary Table 2). Surgical resection was also associated with lower abundance of *Faecalibacterium prausnitzii* (mean 0.014 vs 0.248 respectively, $p = 0.001$). No significant differences were detected in other diversity indices.

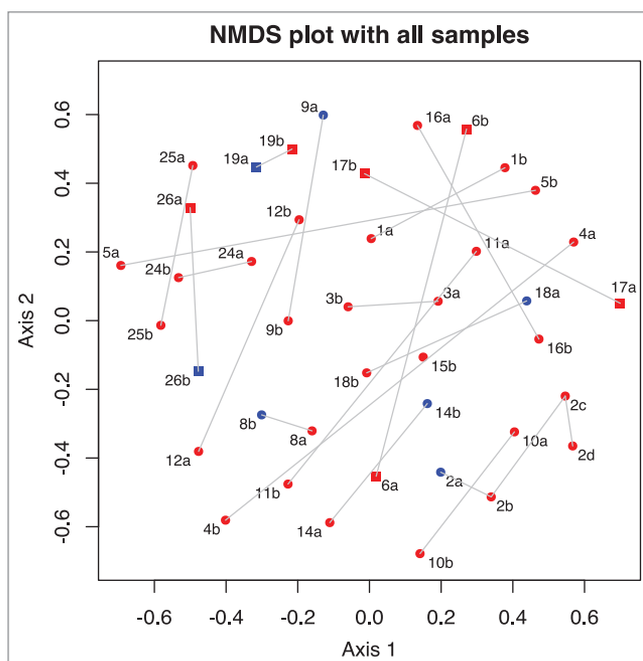


Figure 1. Non-metric multi-dimensional scaling plot of Yue and Clayton theta distances between all samples. Numbers represent different patients. *a* represents a sample collected at the first time-point, *b* represents the second time-point, *c* at third time-point and *d* the fourth. Samples taken in the presence of active inflammation (histological or endoscopic) are coloured red, those with no inflammation are coloured blue. Samples from those with Crohn's disease are circles, samples from those with ulcerative colitis are squares.

The non-metric multidimensional scaling (NMDS) plot in Fig. 1 shows that the microbiome of some patients changed markedly between procedures, whereas others did not. To find out if this was related to underlying patient characteristics, Yue and Clayton theta (Θ_{YC}) distances between different time points from the same patient were compared to clinical meta-data. A significant difference was detected between sample pairs with no active inflammation at one time-point compared to those with histologically confirmed active inflammation at both time-points (adjusted $p = 0.00031$, see Fig. 2 and Fig. 3). Statistical significance was retained even after exclusion of samples from patients with previous abdominal surgery and those with antibiotic exposure ($p = 0.0050$). A linear regression was performed to determine if time between procedures could explain the changes in the microbiome, but no significant associations with alpha diversity or with Θ_{YC} sample pair distance were detected ($p = 0.314$ and 0.227 respectively) (see also Supplementary Table 1).

Weighted Unifrac calculations demonstrated significant differences in the structures of the communities

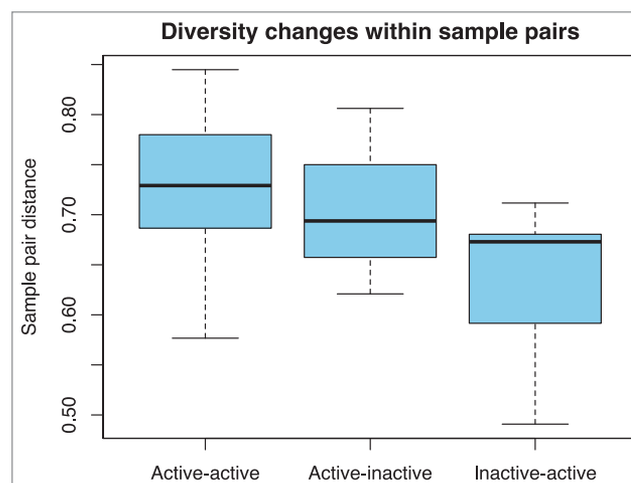


Figure 2. Differences in beta diversity distances (Yue and Clayton theta distances), pairwise, based on presence or absence of inflammation. Sample pairs (ie from the same patient) had inflammation at both time-points (Active-active); had active inflammation initially, and no inflammation at follow-up (Active-inactive); or no inflammation initially and active inflammation at follow-up (Inactive-active).

from samples with active inflammation compared to those without (score = 0.5800, $p < 0.001$). Samples collected in the presence of inflammation were associated with a lower relative abundance of bacteria from the phylum, Firmicutes, and a higher proportion of

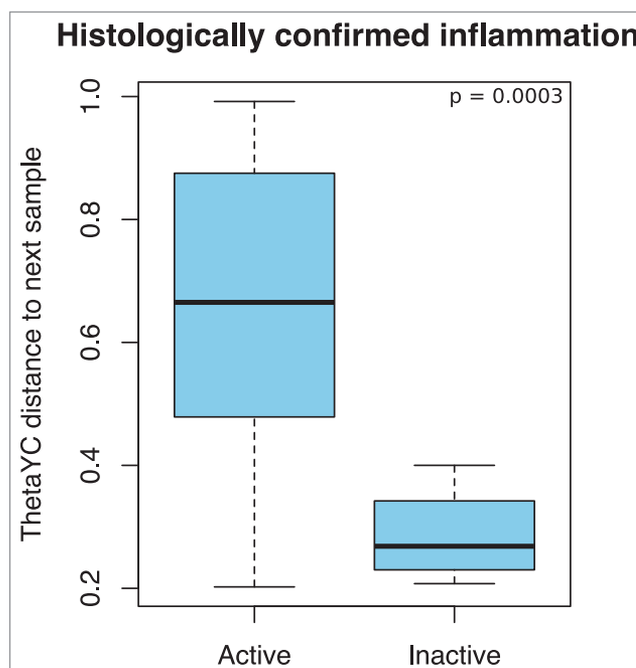


Figure 3. Beta diversity: Yue and Clayton theta distances between sample pairs. The presence (active, $n = 29$) or absence (inactive, $n = 4$) of histologically confirmed inflammation and the distance to the next sample.

Bacteroidetes compared to those with no inflammation, but these were not statistically significant (Supplementary Figure 2). OTU analysis did not reveal individual OTUs to be significantly differently represented across the two groups after correction for false discovery (Supplementary Table 3). However, the relative abundance of *Faecalibacterium prausnitzii* was noted to be lower in samples with active inflammation compared to those without (mean relative abundance 0.1931 vs 0.2817 respectively, one-way ANOVA $p = 0.010$). Consistent with previous studies, a lower relative abundance of *F. prausnitzii* was noted in patients with CD versus UC (mean 0.1268 v 0.3467 respectively, $p = 0.041$).

There were significant differences in the structure of the bacterial communities of smokers compared to non-smokers (weighted Unifrac Score 0.505499, $p = 0.011$), but no significant differences in relative abundance in individual OTUs between smokers and non-smokers once p values were corrected. There were no differences in alpha diversity or Θ_{YC} distance between sample pairs based on smoking status, even when those with UC were excluded.

Inferred metagenomics pathway analysis did not detect significant differences between the samples based on clinical metadata including prior surgery, IBD subtype, smokers, medication and antibiotic exposure and presence of inflammation (Supplementary Figure 3).

Discussion

Here we describe changes in the microbiome over an average 2.4 years in 20 patients suffering from IBD, using mucosal samples to characterize the microbiome, and endoscopic and histological analyses to assess the degree of inflammation. When both samples were obtained in the presence of inflammation, the bacterial populations were more dissimilar than if one sample had no evidence of inflammation. We believe that this reflects an inherent instability of the microbiome in patients with ongoing inflammation.

All human temporal studies of the microbiome in IBD to date used no or surrogate markers for inflammation and most used stool samples. Shaw *et al* used faecal calprotectin as a marker of inflammation and stool specimens to analyse the microbiome in a longitudinal study of 19 newly diagnosed paediatric patients.²⁴ They describe ongoing dysbiosis in patients

with high faecal calprotectin levels. Wright *et al*²³ used 16S rRNA sequencing of intestinal biopsies to analyse the microbial communities of 34 patients who had undergone ileocaecal resection. There was no decrease in alpha diversity in patients who remained in remission compared to those who developed endoscopically active disease, but they did not report changes in microbiome related to histologically confirmed inflammation. The presence of *Proteus* in post-operative ileal mucosal specimens was independently associated with post-operative recurrence (OR 13 (1.1-150), $p = 0.039$), but we only identified *Proteus* in one sample; from a patient with active Crohn's disease with no prior surgery. Will *et al*²⁵ used faecal samples to describe bacterial communities from 19 IBD patients in remission who later developed active disease. They did not find significant differences in alpha diversity between the two groups, but found that thiopurine use was associated with a significant reduction in richness and diversity based on Chao1 and Shannon diversity indices (Shannon index: 5.13 vs 6.78, in non-users and users respectively; $p < 0.01$). In our study, thiopurine use was not associated with differences in alpha diversity (mean Shannon index: 2.46 vs 2.29, in non-users and users respectively; $p = 0.322$), nor were there differences in relative abundances of specific taxa. Similarly, there were no differences in alpha diversity with respect to anti-TNF α use (mean Shannon index 2.41 vs 2.26, in those on anti-TNF α agents and those not respectively; $p = 0.30$). Martinez *et al*²⁶ used denaturing gradient gel electrophoresis (DGGE) band profiles from bacterial 16S rRNA amplicons obtained from faecal samples in IBD patients and controls. They described significant reductions in overall diversity as well as temporal instability in 16 patients with UC over one year compared with 8 healthy controls.

In the largest temporal study conducted to date, Halfvarson *et al*¹ conducted bacterial 16S rRNA high-throughput sequencing of three-monthly faecal samples to describe the changes in 109 patients with IBD over a two-year period, with a control group consisting of nine subjects. They found that the microbiome of IBD patients was more volatile, and deviated significantly from a defined "healthy plane", to which the normal controls were confined. Using faecal calprotectin levels $>150 \mu\text{g/g}$ as a marker of disease activity they found that inflammation was not directly correlated with distance to the healthy plane, but there was

some correlation between the observed dramatic fluctuations in the gut microbiome and intensified medication due to a flare of the disease. They did not identify specific bacterial taxa that varied significantly with active inflammation.

In animal models, it is not necessary to use surrogate markers and an association between inflammation and its effects on the microbiome can be demonstrated. Lupp *et al* (2007) described a clear and significant reduction in the number and diversity of colonic bacteria in the presence of host-mediated intestinal inflammation.²⁷ These changes were described in inflammatory models induced by *Citrobacter rodentium* infection, dextran sodium sulfate (DSS) administration and IL10 knockout mice.

There were several limitations of our study. The number of samples is relatively small, but we used the most accurate means of measuring both the mucosa-associated microbiome and the degree of inflammation: demonstrating statistical significance with a small sample size is indicative of the power of the effect. The absence of a control group limits comparisons to IBD subjects alone, but having a healthy control group undergoing repeated colonoscopy and mucosal biopsy is not without risk, so HREC/IRB approval was not sought. There was some heterogeneity in the nature of the specimens: 81% of samples were obtained at the time of colonoscopy, and eight samples were collected at the time of bowel resection. However, these two groups did not differ in their diversity indices even accounting for exposure to antibiotics at the time of surgery. Nor were there significant differences in bacterial composition based on the relative abundance of OTUs. Diet, which leads to substantial changes in the microbiome,^{28,29} could not be controlled, which may also have influenced results.

Patients with IBD are exposed to many different medications and an almost universal criticism of all clinical studies in IBD is a failure to account for their confounding effect. The only way to overcome this is to study inception cohorts, but even then, given that most patients suffering from IBD will go on to receive maintenance therapy, the expectation that there will be treatment-naïve controls at different time-points is unlikely to be met by any study. It is also unlikely that any patient with active disease would remain on a stable treatment regimen over time. Flares of the disease, intensification of medication and changes in the microbiome occur simultaneously: our strategy to

analyse the relationship between these factors was to use the most accurate measures of inflammation and the microbiome and to control for the effects of medication in the statistical analysis.

Our results suggest that patients with IBD are less able to maintain a stable microbial community because of uncontrolled inflammation. Our group has previously demonstrated that bowel preparation and colonoscopy does not change the microbiota in most subjects,³⁰ but noted that three patients suffering from ulcerative colitis did not have restoration of their pre-colonoscopy microbiome. Therapeutic manipulation of the microbiome, for example with faecal microbial transplantation, has had mixed results in IBD cohorts compared to those with recurrent *Clostridium difficile* colitis,^{31,32} also raising the possibility that there is instability of the microbiome in IBD patients compared to previously healthy subjects.

In summary, our study demonstrated temporal shifts in the microbiome of patients with IBD that were closely correlated to the presence or absence of inflammation. The microbiome over time was least similar in those who had ongoing histological inflammation suggesting that it was the cause rather than the consequence of “dysbiosis”. The implications are important given the efforts to develop various products to treat IBD by correcting “dysbiosis” (e.g. pre- and probiotics, therapeutic microbiota, and faecal microbial transplantation). If “dysbiosis” were the cause of these differences, the inference of our study is that there are different types of “dysbiosis” leading to inflammation in the same patient at different times.

The evidence for a microbial trigger in the development of IBD is strong, but no unique microorganism or “dysbiosis” common to all patients has been identified despite intensive analysis by many groups. Recent evidence linking cells of the monocyte-macrophage lineage to genetic susceptibility for IBD, leads us to suggest that it is the dysregulation of monocyte adaptation to the environment of the gastrointestinal mucosa that causes inflammation in patients suffering from IBD and the resultant “dysbiosis”.³³ Future studies examining the interaction between an individual’s innate immune system and specific components of the microbiome may lead to a better understanding of the mechanisms by which microbes initiate inflammation in IBD.

Materials and methods

Ethical approval was obtained from the Australian National University (Protocol: 2012/596) and ACT Health (Protocol: ETH.5.07.464). Each subject provided written informed consent.

Biopsy samples were collected from patients with known or suspected IBD attending Canberra Hospital between 2004 and 2014 at the time of colonoscopy or intestinal resection. Biopsy samples were placed in a sterile tube containing RNeasy[®] at the time of colonoscopy or resection. Samples were kept at 4°C for 24 h and transferred to -80°C for storage.

Clinical information was collected at the time of the procedure. This included disease status, date of diagnosis, Montreal classification, gender, age, smoking status, surgical history, medication use and recent exposure to antibiotics. The presence (hereafter referred to as “active”) or absence (“inactive”) of inflammation was determined by endoscopic assessment and the presence of microscopic inflammation in the histopathology report (“histologically confirmed”). Samples were only described as “inactive” if there was no evidence of inflammation endoscopically or histologically.

DNA extraction

With a few modifications, DNA extraction was performed using a Qiagen DNeasy Blood and Tissue kit (catalogue number 69504) as per the manufacturer’s instructions. Bead beating was performed in TE buffer at 30 rev/s for 3 min until visibly homogenised, using 15–20 0.1 mm beads for bacterial cell lysis, four 2.3 mm and four 1 mm beads for eukaryotic cell lysis (Daintree Scientific). The amount and quality of DNA eluted was measured using a Nanodrop ND-1000 spectrophotometer (Analytical Technologies).

16S rRNA high throughput sequencing

Amplification of variable region 4 (V4) of the 16S rRNA gene was performed using the following primers: 515F (5'-**CCA TCT CAT CCC TGC GTG TCT CCG ACT CAG BARCODE** TGT GCC AGC MGC CGC GGT AA-3') and 806R (5'-**CCT CTC TAT GGG CAG TCG GTG AT** GGA CTA CHV GGG TWT CTA AT-3') where the sequences in bold represent Adapter A and Adapter P1 of the Ion PGM Hi-Q View OT kit (Life Technologies), *BARCODE*

represents the 10 bp IonExpress barcode and the remainder represents the V4 region of the 16S rRNA gene. PCR was performed using the Platinum Taq DNA Polymerase High Fidelity kit (catalogue number 11304-011) on a BioRad T100 thermal cycler (BioRad Laboratories). 50 μ L reactions contained 5 μ L of 10x HiFi buffer, 1 μ L of 10 μ M deoxynucleotide triphosphate mix, 2 μ L of 50 μ M magnesium sulphate solution, 2.5 μ L of 10 mM IonTorrent forward primer, 2.5 μ L of 10 mM reverse primer, 0.2 μ L of Platinum Taq DNA polymerase, up to 35.8 μ L sterile distilled water, and up to 5 μ L of template DNA. Initial denaturation was performed at 94°C for 3 min. This was followed by 30 cycles of denaturation at 94°C for 15 s, annealing at 55°C for 30 s, and extension at 68°C for 1 min. Final extension was performed at 68°C for 10 min. PCR products were extracted from 1.5% agarose gels. Following this, the PCR products underwent gel purification (Wizard SV Gel and PCR Cleanup System, Promega) and bead purification (Agencourt AMPure XP, Beckman Coulter). DNA concentrations were determined with an Agilent 2100 Bioanalyzer (Agilent Technologies) on DNA 1000 chips. Equimolar amounts of PCR products were combined and used as template for the Ion PGM Hi-Q SEQ kit (400bp) reactions. The Ion PGM (Life Technologies) platform was used to sequence the products.

Sequence processing

16S rRNA gene sequences were processed and analysed using the open source program mothur v 1.36.1³⁴ and the accompanying 454 standard operating procedure.³⁵ Sequences were trimmed, using a qwindow average of 30 and window size of 50. Low-quality sequences were eliminated. Sequences were then trimmed of barcode (0 bp barcode mismatches allowed) and primer (2bp mismatches allowed) then merged. A total of 5,152,850 sequences were detected, and 881,073 were unique. Chimeras were removed using the UCHIME reference based method.³⁶ Sequences were classified using the Silva reference set (release 123)³⁷ and a cut-off of 80. A distance matrix with a cut-off of 0.15 was calculated using the dist.seqs command in mothur. Sequences were then classified into operational taxonomic units (OTUs) using a 97% cut-off value. After filtering, clustering and classifying the samples, 13,394 unique sequences were identified,

from a total 1,221,636 sequences (average 250 bp). Each sample contained an average 29,788 quality sequences (range 329–137,563), representing an average of 119 OTUs. Samples were rarefied to 1418 sequences, which excluded one pair of samples with a low number of sequence reads. Intra-sample (alpha) diversity was measured by calculating the Shannon diversity index. Comparisons were made between different clinical parameters, including active/inactive inflammation, smoking status, disease type (CD or UC), previous surgery, medication exposure and disease duration (Supplementary Table 1).

Statistical analysis

Alpha diversity was measured by calculating the Shannon index and Chao1 richness estimator using *mothur*. Non-metric multidimensional scaling (NMDS) plots were created after calculating distance matrices. An NMDS plot demonstrates the distance between samples in two dimensions. The further two points are from each other the greater the difference in their microbiomes. To determine what factors are associated with the greatest change over time, Yue and Clayton theta (Θ_{YC}) distances between sample pairs were compared within clinical groups.³⁸ This takes into account the relative abundance of shared and non-shared OTUs in the samples compared. The more dissimilar the bacterial populations within two samples are the larger the Θ_{YC} distance becomes. Analyses by OTU were performed using *mothur*, R (version 3.3.2) and *metastats*.^{39,40} The samples were grouped according to disease status (active or inactive), smoking status, previous abdominal surgery, IBD subtype (CD or UC), medication use, disease duration, sampling technique (colonoscopy or resection) and examined for differentially abundant bacterial taxa using *metastats* within *mothur*. To reduce false discovery, OTUs were only included where they represented an average relative abundance > 0.1%. Where adjusted p values were calculated, they were corrected for false discovery using the Benjamini-Hochberg method. A hypothesis was considered significant if the adjusted p value was <0.05. Comparisons between groups were conducted using one-way ANOVA tests, student's t-test and linear regression analyses, using R (*vegan* and *stats* packages) and IBM SPSS Statistics (version 24.0).

Inferred metagenomics

Rarefied sequence datafiles were aligned to the Greengenes reference (version 13.8) using 97% identity for use in the Phylogenetic Investigation of Communities by Reconstruction of Unobserved States (PICRUSt) analysis pipeline.⁴¹ Samples were first normalised by copy number, and metagenomic prediction was performed based on the Kyoto Encyclopaedia of Genes and Genomes (KEGG). Kruskal-Wallis sum-rank test and Linear Discriminant Analysis Effect Size (LEfSe) were used to determine metabolic functional pathways that were differentially abundant between groups.⁴²

Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

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