Title:
The human mesenteric lymph node microbiome differentiates between Crohn’s disease and ulcerative colitis.

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Abbreviations:
MLNs – mesenteric lymph nodes; IBD – inflammatory bowel disease; CD – Crohn’s disease, UC – ulcerative colitis; DNA – deoxyribonucleic acid; PCR – polymerase chain reaction; UHL – University Hospital Limerick; RNA – ribonucleic acid; PBS – phosphate buffered saline; IDT – Integrated DNA Technologies; OTU – operational taxonomic unit; GOLD – Genomic OnLine Database; RDP – Ribosomal Database Project; CI – confidence interval; Principal Coordinates Analysis (PCoA)

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Abstract

Background and Aims

Mesenteric lymph nodes are sites in which translocated bacteria incite and progress immunological responses. For this reason, understanding the microbiome of mesenteric lymph nodes in inflammatory bowel disease is important. The bacterial profile of Crohn’s disease mesenteric lymph nodes has been analysed using culture-independent methods in only one previous study. This study aimed to investigate the mesenteric lymph node microbiota from both Crohn’s disease and ulcerative colitis patients.

Methods

Mesenteric lymph nodes were collected from Crohn’s disease and ulcerative colitis patients undergoing resection. Total DNA was extracted from mesenteric lymph nodes and assessed for the presence of bacterial DNA (16S). All work was completed in a sterile environment using aseptic techniques. Samples positive for 16S DNA underwent next generation sequencing and the identity of bacterial phyla and species were determined.

Results

Crohn’s disease mesenteric lymph nodes had a distinctly different microbial profile to that observed in ulcerative colitis. The relative abundance of Firmicutes was greater in nodes from ulcerative colitis patients while Proteobacteria were more abundant in Crohn’s disease. Although species diversity was reduced in Crohn’s disease mesenteric lymph nodes, they contained greater numbers of less dominant phyla, mainly Fusobacteria.
Conclusion

This study confirms that there are distinct differences between the Crohn’s disease and ulcerative colitis mesenteric lymph node microbiomes. Such microbial differences could aid in the diagnosis of Crohn’s disease or ulcerative colitis, particular in cases of indeterminate colitis at time of resection, or help explain their mechanisms of development and progression.

Keywords

Inflammatory bowel disease; microbiome; mesenteric lymph nodes
Introduction

Inflammatory bowel diseases (IBD), which include Crohn’s disease (CD) and ulcerative colitis (UC), are thought to result from an overly aggressive immune response in genetically susceptible individuals to an environmental factor, such as gut commensals.\(^1\),\(^2\) The highest prevalence rates worldwide of CD and UC are found in Europe, with 322 and 505 cases per 100,000 persons, respectively.\(^3\)

The human gut microbiota has been investigated to a great extent, in both health and disease, in particular gastrointestinal diseases, such as IBD. The earliest of these studies investigated bacterial content in faecal matter using conventional culture methods.\(^4\)\,-\(^11\) Following this, the microbial profile of mucosal tissue was explored as it was believed to be a more accurate representation of the gut microbiota. Indeed, it has since been proven, by more recent comprehensive studies, that significant differences exist in the bacterial groups found in the mucosa and faecal matter.\(^12\),\(^13\) Culture methods were also utilised once again, to first ascertain the mucosal tissue microbial profile.\(^14\)\,-\(^17\) These studies provided a preliminary insight into the human gut microbiota, however, it has been estimated that up to 70-80% of intestinal bacteria are unculturable.\(^18\),\(^19\) For this reason, the development of culture-independent methods has significantly advanced our knowledge of gastrointestinal flora.\(^20\)\,-\(^22\) These techniques have been used to great effect in establishing a gut microbiota profile in health and inflammatory bowel disease.

In healthy individuals, Firmicutes (phylum including Enterococaceae, Clostridiaceae, and Ruminococcaceae) and Bacteroidetes (phylum including Bacteroidaceae, Porphyromonadaceae, Prevotellaceae, Rikenellaceae and Flavobacteriaceae) phyla dominate, with lower abundance of other phyla, mainly Proteobacteria (phylum including Enterobacteriaceae and Burkholderiaceae) and Actinobacteria (phylum including Mycobacteriaceae, Micrococcaceae and Streptomycineae).\(^23\)\,-\(^25\) The IBD gut microbiota displays dysbiosis when compared to healthy individuals and there is a marked reduction in bacterial diversity, related particularly to the Firmicutes and Bacteroidetes phyla.\(^26\),\(^27\) This coincides with increase in Gammaproteobacteria (which includes Enterobacteriaceae and Pseudomonadaceae).\(^26\) The CD microbiome displays greater dysbiosis than that found in UC, specifically a greater reduction in microbial diversity, with a more altered and less stable microbiome composition.\(^28\) Inflamed mucosal tissue from CD patients contains higher levels of Bacteroidetes and Fusobacteria (includes Fusobacterium nucleatum), while Proteobacteria and Firmicutes (in particular Clostridiaceae, Enterobacteriaceae and Ruminococcaceae) are more frequently observed in inflamed UC mucosa.\(^29\),\(^30\) Despite these changes, the microbiome of UC patients has been described as one that is more similar to healthy individuals.\(^31\),\(^32\)

Although mesenteric lymph nodes (MLNs) have been associated with initiation of immunological responses to bacterial translocation,\(^33\)\,-\(^35\) the importance of the MLN microbiome in IBD remains unclear.
To our knowledge, only one study has assessed the microbial content of MLNs in CD using culture-independent methods, reporting a bacterial profile similar to that of the CD gut microbiome and no study has described the microbiome of UC patient MLNs.\textsuperscript{36} This is despite various culture-dependent studies demonstrating bacterial translocation to MLNs and mesenteric fat in IBD.\textsuperscript{37-39} Consistently, the most common organism detected in the MLNs in IBD was \textit{Escherichia coli},\textsuperscript{37,38,40} however, the full diversity of the MLN microbiota has not yet been elucidated. Addressing this gap in knowledge may prove valuable in better understanding these distinct diseases. Our group has recently highlighted the potential role of the mesentery in IBD and has demonstrated that surgical recurrence rates for ileocolic CD patients are significantly reduced when the mesentery, and MLNs, are removed during resection.\textsuperscript{41,42} As the similarity of the MLN and mucosal microbiome in CD has been comprehensively reported previously and and selective bacterial translocation from the gut to the nodes effectively refuted,\textsuperscript{36} our interest concerned the variation in bacteria in MLNs from IBD patients. Therefore, we aimed to investigate and compare the microbial profile of MLNs from CD and UC patients diagnosed pathologically using culture-independent methods for the first time.
Methods

Ethical approval

Ethical approval was obtained from the Research Ethics Committee of University Hospital Limerick (UHL).

Inclusion and exclusion criteria

All adult patients (≥18 years) undergoing resection for histopathologically diagnosed Crohn’s disease or ulcerative colitis in UHL from October 2015 to September 2016 were recruited. Patients were excluded if they had been diagnosed with colorectal cancer previously or were suspected to have colorectal cancer at the time of operation so as not to interfere with pathologic analysis.

Mesenteric lymph node harvest

Where possible, IBD patients in UHL undergo resections that include the mesentery, allowing for greater (MLN) yields. Diseased intestine, along with associated mesentery, was resected and placed immediately in a sterile surgical tray. The resected specimen was kept in the sterile surgical area until lymph nodes could be harvested. Lymph nodes were harvested from resected specimens in a sterile environment, which was outside the sterile surgical area but under the same atmospheric pressure, using aseptic techniques and sterile surgical instruments. The lymph nodes were identified and cut from the mesentery by a trained colorectal surgeon. Nodes were placed, without handling, into sterile microtubes.

Although desirable, it was not possible to utilise nodes as morphological controls (e.g., nodes halved and sent for both 16S testing and pathological analysis). All Protect Tissue Reagent (Qiagen, UK) was added to each specimen tube until tissue was fully immersed. A separate control sample of All Protect Tissue Reagent was also obtained. Samples were kept at room temperature for a maximum of 2 hours before transfer to long term storage at -80 °C.

DNA extraction

Lymph nodes were thawed on ice for 30 minutes before processing. All work was performed in sterile conditions in a Class II biosafety cabinet using aseptic techniques and DNase free instruments and consumables. Excess All Protect Tissue Reagent was removed and nodes were washed in 1 mL sterile phosphate buffered saline (PBS; pH 7.4 [137 mM NaCl; 2.68 mM KCl; 9.94 mM Na₂HPO₄; 1.76 mM KH₂PO₄]). MLNs were weighed and samples of approximately 25 mg were used for DNA extraction. Total DNA was extracted from samples using a QIAamp cador pathogen Mini Kit (Qiagen, UK) using the manufacturer’s instructions. In brief, tissue lysis buffer (180 µL; Buffer ATL) was added to the node tissue and sample was homogenised for 1 minute using a motorised pestle. Proteinase K enzyme (20 µL) was added to this to commence enzymatic tissue digestion. Samples were incubated overnight at 56 °C with constant shaking. Following this, digested tissue (200 µL) mixed with sterile PBS (200 µL) was transferred to a Pathogen Lysis Tubes S (Qiagen, UK), pre-prepared with tissue lysis buffer (Buffer ATL) and anti-foaming reagent (Reagent DX) (100 µL total) for
mechanical disruption of hard-to-lyse bacteria by glass beads. This was completed by vigorous shaking for 10 minutes. Buffer VX (100 µL), which ensures the inactivation of nucleases when in the presence of proteinase K (100 µL) was added to the pre-treated samples (enzymatically digested and mechanically disrupted), mixed well and left to incubate at room temperature for 15 minutes. Binding reagent (Buffer ACB; 350 µL) was added to samples, mixed well and transferred to spin columns. DNA was then isolated following a series of centrifugations using QIAamp spin columns and buffer solutions. DNA was eluted in 100 µL elution buffer (Buffer AVE) (2 x 10 minute elutions) and quantity and quality were assessed using the 260/280 function on the Spectrostar Nano plate reader LVis plate function (BMG Labtech). Extracted DNA was stored at -20 °C for subsequent PCR.

Polymerase chain reaction (PCR) amplification of 16S DNA

The presence or absence of bacteria in MLNs was confirmed by PCR amplification of 16S (bacterial) DNA. All work was carried out on a dedicated clean PCR bench, in sterile conditions and using sterile and DNase free instruments and consumables. Suitable positive (extracted DNA from a combined mixture of bacterial species) and negative (no template DNA and extracted DNA from PBS false extraction) controls were included. The PCR reaction included HotStarTaq Master Mix (25 µL), 500 ng template DNA, 0.5 µM 16S rRNA Forward primer, 0.5 µM 16S rRNA Reverse primer. The total volume of the PCR reaction was 50 µL; the remaining volume made up with PCR grade water (HotStarTaq Master Mix Kit, Invitrogen). The forward and reverse primers were ready-made primers obtained from Integrated DNA Technologies (IDT) and are designed to incorporate a large portion of the 16S gene: For (5'-AGA GTT TGA TCC TGG CTC AG-3') and Rev (5'-ACG GCT ACC TTG TTA CGA CTT-3'). Reactions were performed using a Kyratec Thermal Cycler (Kyratec, Australia): heat activation step at 95 °C for 15 minutes; 30 cycles of denaturation at 94 °C for 30 seconds, annealing at 57.49 °C for 30 seconds and extension at 72 °C for 60 seconds; and the final extension step at 72 °C for 10 minutes. Amplicons were extracted using GenElute PCR Clean-Up Kit (Sigma Aldrich) as per the manufacturer’s instructions. Amplified DNA was eluted in 50 µL elution buffer (from kit) and presence, quantity and quality were assessed using the 260/280 function on the Spectrostar Nano plate reader. DNA quality was also assessed by visualisation on a 1% agarose gel. Lymph nodes were deemed as “16S negative” when: (1) no quantity of DNA was detected on the Spectrostar Nano plate reader, (2) the 260/280 ratio and graph profiles were not similar to that expected of DNA content and (3) a large quantity of the amplified product was ran on a 1% agarose gel and no DNA band was visualised. Amplified DNA was stored at -20 °C for future species identification through sequencing.

Next generation sequencing

Library preparation and sequencing were completed in the Teagasc Sequencing Centre (Cork, Ireland) as follows: DNA was quantified using the Qubit High Sensitivity Kit (Life Technologies), standardised and then used as a template for PCR. PCR primers and conditions are essentially as outlined in the Illumina 16S Metagenomic Sequencing Library preparation guide (Illumina) with the following exceptions: For the initial 16S PCR, the process was performed using Kapa Robust (Roche) in 50 µL reaction volumes, and 30 cycles were used in the PCR. The V3-V4 variable regions were amplified by the primers. These regions are the suggested target for the Illumina 16S Metagenomic
Sequencing system as previously found to be the optimum primer pair for sequencing based species diversity studies.\textsuperscript{33,44} Products were then cleaned with an appropriate volume of Ampure beads and eluted in 30 µl per sample. This was then used as the template for the index PCRs as outlined in the protocol (Illumina). Library quantification, normalisation, and pooling were as outlined in the protocol. After pooling, the sample was re-quantified by qPCR using the Kapa Library Quantification Kit for Illumina (Roche) and run on an Agilent high sensitivity chip (Agilent). Library denaturation and MiSeq sample loading were then performed according to manufacturer’s instructions (Illumina). PhiX was spiked into the final pool at 20% (v/v) and sequencing was performed using a 500 cycle V2 chemistry kit (Illumina). Previous studies using the Illumina MiSeq system and primer pairs targeting the V3-V4 variable regions have identified bacteria to species level.\textsuperscript{45,46} The V3-V4 variable regions are commonly used to investigate the gut microbiome and to detect intestinal bacterial species.\textsuperscript{45-48}

**Bioinformatics**

Illumina reads were filtered on the basis of quality (removal of low quality nucleotides at the 3’ end, in addition to bases where quality was below 20 in a trimming window) and length (removal of sequences with less than 200 bp) with PRINSEQ,\textsuperscript{49} and the paired-end reads with a minimum overlap of 20 bp were joined using Fastq-join.\textsuperscript{50} A pipeline (a series of data processing events) was generated using clean reads only (i.e. filtered reads described above), to cluster sequences (from next generation sequencing data) with 97% identity to obtain Operative Taxonomic Unit (OTUs) using closed-reference USEARCH v7.0 algorithm (Edgar RC. 2010) and remove chimeric OTUs against the Genome OnLine Database (GOLD). An OTU represents a group of similar 16S rRNA sequences that theoretically reflect shared species identity. However, while a 97% threshold is sufficient for diversity studies, similarity thresholds of greater than 98% are required to identify species definitively. The taxonomic assignment of these OTUs was obtained against the Ribosomal Database Project (RDP; \url{http://rdp.cme.msu.edu/}).\textsuperscript{51} Beta-diversity was determined using QIIME,\textsuperscript{52} and additional analysis were performed with the R package, phyloseq.\textsuperscript{53}

**Statistical analysis**

Data are presented as mean ± 95% confidence interval (CI). To determine phyla, family and species OTUs, data were statistically analysed by PERMANOVA for beta-diversity. All other statistical analyses were completed in SPSS v22 (SPSS Inc., Chicago, USA). An independent two-tailed t-test was used to compare parametric variables whilst a Mann-Whitney U test was utilized for non-parametric comparisons. Chi-square tests were used to compare nominal data. Species richness estimates were generated using Chao 1 and Abundance-based Coverage Estimator (ACE) estimators while community and species diversity were estimated using the Shannon Diversity index. A 5% level of significance was used for all statistical tests.
Results

Patient and MLN characteristics

Thirteen pathologically diagnosed IBD patients (CD: n=5, 38%; UC: n=8, 62%) were recruited (Supplementary Table 1). These underwent fifteen surgical procedures for IBD. All patients had intravenous antibiotics administered with induction of anaesthesia, approximately 15 minutes before the start of the surgical procedure, as per standard of care. MLNs were taken from locations along the small bowel mesentery and mesocolon (Figure 1A-C). No biases were observed regarding number of lymph nodes or lymph node diameter irrespective of sampling location. There was no evidence of abscesses in MLNs collected. Furthermore, histopathologic analysis of resection specimens did not identify the presence of mesenteric abscesses. In total, 25% and 24% of MLNs collected from CD and UC patients, respectively, did not contain bacterial DNA (Figure 1D, E). Demographics of patients with bacterial DNA positive nodes are provided in Table 1.

The microbiome of MLNs from CD patients is distinctly different to those from UC patients.

Bacteria from fourteen different phyla were detected, irrespective of disease cohort (Figure 2). The majority of OTUs belonged to Firmicutes and Bacteroidetes, followed by Proteobacteria and Actinobacteria. Other phyla, generally less studied and only defined in the last two decades (e.g., Parcubacteria, Verrucomicrobia) were also detected. MLNs were obtained from various locations within the mesentery from Patients 5, 6 and 7, such as the mesorectum (M8) and mesosigmoid (M9 and M10) from Patient 5, the mesorectum (M11 and M12) and right mesocolon (M13) from Patient 6, and the mesentery associated with the caecum (M16), transverse mesocolon (M17 and M18), mesosigmoid (M15 and M19) and mesorectum (M14) from Patient 7 (Figure 2). The microbial profiles of MLNs taken from the same patient were similar, irrespective of sampling location. Furthermore, the microbiomes of MLNs taken from areas of mesentery that were determined to be disease-free (i.e., no features of macroscopic inflammation) at time of harvest were similar to those nodes taken from areas of diseased mesentery (i.e. displayed features of inflammation and disease) (Patient 5, UC, disease-free nodes: M8 and diseased nodes: M9 and M10; Patient 13, CD, disease-free nodes: M32 and diseased nodes: M33; Figure 2). There was no readily apparent association between disease behaviour and bacterial profiles.

The relative abundance of Firmicutes was greater in UC MLNs than in those of CD patients (52.8% ± 6.73% vs. 18.6% ± 4.65%; p<0.0001, independent t-test) (Figure 3). In contrast, MLNs from CD patients contained greater abundance of Proteobacteria (36.1% ± 11.06% vs. 11.8% ± 3.63%; p=0.005, Mann-Whitney U test) and greater numbers of less dominant phyla, mainly Fusobacteria (7.3% ± 4.98% vs. 0.3% ± 0.1%; p<0.0001, Mann-Whitney U test) (Figure 3A, B). Similar levels of Bacteroidetes and Actinobacteria were detected in the two cohorts (Figure 3A, B). The ratio of Firmicutes to Bacteroidetes was increased in UC MLNs (1.613), but decreased in CD MLNs (0.705) (Figure 3C).
Interestingly, although Firmicutes was the most abundant phylum in UC MLNs, the most abundant bacterial genus was *Bacteroides* (phylum Bacteroidetes), followed by *Faecalibacterium* (phylum Firmicutes) \(p<0.0001\), Kruskal Wallis test). In CD MLNs, the most abundant bacterial genus was *Flavobacterium*, followed by *Bacteroides*, both members of phylum Bacteroidetes \(p<0.0001\), Kruskal Wallis test). Although *Faecalibacterium* (i.e. *Faecalibacterium prausnitzii*), an important component of the healthy gut microbiome, was abundant in UC, it was present to a lesser extent in CD \(10.9\% \pm 4.26\%\) vs. \(1.2\% \pm 1.15\%\); \(p<0.0001\), Mann-Whitney U test). Definitively, a Principal Coordinates Analysis (PCoA) of unweighted Unifrac distance, demonstrated that the bacterial composition of UC and CD MLNs were distinctly different (*Figure 4*).

**Diversity and species richness is reduced in Crohn's disease MLNs.**

Species diversity was estimated using the Shannon Diversity index. MLNs from UC patients displayed significantly greater microbial diversity than those of CD patients \(1.95 \pm 0.0035\) vs. \(0.79 \pm 0.0018\); \(p<0.0001\), Mann-Whitney U test) (*Figure 5*). We determined species richness using the Chao 1 and ACE estimators (*Supplementary Table 2*). The Chao 1 estimator provides investigators with a lower bound estimate for species richness. This indicated greater species richness in CD MLNs than those of UC \(237\) vs. \(183\) (*Supplementary Table 2*). ACE is a more comprehensive estimate of species richness, derived from the number of rare and common species present in a sample while also considering how many more undiscovered species there may be. This showed greater species richness in UC than in CD MLNs \(175-178\) vs. \(148-156\) (*Supplementary Table 2*) similar to actual raw species counts observed.
Discussion

This is the first study to investigate and report distinct differences between the microbial profiles of MLNs from UC and CD patients. Aided somewhat by the advantageous availability of resected material from UC patients, this report also represents the first time the microbiome of MLNs from UC patients has been elucidated. In comparison to the reported normal microbiome, the UC MLN microbial profile appears imbalanced in favour of elevated Firmicutes relative to Bacteroidetes. CD MLN dysbiosis is reflected by overabundance of Proteobacteria (e.g., pathogens such as Escherichia, Shigella, Helicobacter and Salmonella), a decreased Firmicutes/Bacteroidetes ratio and a reduction in species diversity and richness. The MLN bacterial profiles remained unchanged irrespective of extent of disease or sampling location. Notably, MLNs from UC patients who underwent completion proctectomy at least 6 months following their total colectomy were free of bacterial DNA (Patients 10, 12 and 14, Supplementary Table 1). During this time, patients had an end ileostomy in situ. This suggests that, in these patients, diversion of the faecal stream allowed clearance of bacterial DNA from MLNs by the host immune system.

It has been reported previously that the UC and CD faecal and mucosal microbiota are distinct from one another. Our results now confirm that this variation extends to the MLN microbiome. This variation differentiates clearly between the diseases, representing new diagnostic potential and may also offer targets for innovative therapeutic developments. Approximately 10-15% of IBD cases are recorded as indeterminate colitis, meaning a definitive diagnosis of CD or UC cannot be made from the resection specimen, or from biopsies at colonoscopy. We believe that the distinctive UC and CD MLN microbiota could enable definitive diagnoses at resection, potentially revolutionising prophylactic treatment decisions, patient aftercare and survival.

In healthy individuals, the gut microbiota is comprised mainly of Firmicutes and Bacteroidetes (approximately 50% and 40%, respectively), thus the ratio of Firmicutes to Bacteroidetes is often used as an indicator of gut microbiota balance. In this study, the relative abundance of Firmicutes was greater than Bacteroidetes in MLNs from UC patients. Inversely, Bacteroidetes had a greater relative abundance than Firmicutes in CD MLNs. The ratio of Firmicutes to Bacteroidetes in the MLN microbiome from both CD and UC patients has not been characterised previously. However, it has been studied to some extent in IBD patient-derived faeces. Most of those studies reported an overall decrease in the ratio relative to healthy controls. Conversely, there have also been reports of increases in the ratio. This disparity could be explained by geographical, dietary or treatment factors. In addition, such studies often reported their results based on UC and CD patients combined, which could confound results as there is presumption that both diseases share similar microbiomes. One previous study has described a decrease in Firmicutes relative to Bacteroidetes in UC and CD mucosal tissue when studied separately. This differs from the profile that we have observed in UC MLNs. This further differentiates UC and CD and emphasises the need, when analyzing the microbiome of MLNs, to segregate IBD-derived samples.
Mechanistically, the gut microbiota reportedly restricts translocation of pathogenic bacteria to the MLN. This may be compromised in CD but not UC, as the MLN microbiota of CD patients display an over-abundance of Proteobacteria (known to contain numerous pathogenic species). Likewise, the clearance of harmful bacteria from MLNs could also be reduced. The MLNs are sites in which immunological responses can commence and proliferate. It may be that CD MLNs are host to bacterial types which trigger more aggressive responses than those of UC MLNs, such as members of the Proteobacteria phylum. Conversely, Faecalibacterium, which mediates anti-inflammatory effects, is present in CD MLNs in low proportions compared to UC. Is it possible that the pathologic features of CD, which differ greatly to that of UC with regard to the increased level of mesenteric involvement, are a consequence of the CD MLN microbiota? It is reasonable to argue that future work could usefully investigate the role of various members of Proteobacteria or Faecalibacterium, or indeed the ratios of one to the other, in the MLNs and in immunological responses in both CD and UC. In conclusion, the distinctive UC and CD MLN microbiota provide us with a novel diagnostic tool for defining indeterminate colitis, an opportunity to understand the mechanisms mediating each disease, and the prospect of improving patient outcomes.
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Author Contributions:

M.G.K. collected and processed mesenteric lymph nodes, performed experiments and was involved in study design, clinical data collection, data analysis, and drafting of manuscript. J.C.C. was involved in study design, sample collection and drafting of manuscript. P.A.K. aided in laboratory analysis and drafting of manuscript. P.D.C. completed sample library prep and next generation sequencing. R.C.R. completed bioinformatics analysis. K.McD. was involved in study design and drafting of manuscript. C.P.D. initiated study concept and design, data analysis and drafting of manuscript.
Disclosures:

The authors declare no competing personal or financial interests and have nothing to disclose.
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Figure Legends

Figure 1: Mesenteric lymph node (MLN) mapping. (A) Digital image of the small and large bowel with associated continuous mesentery. Source locations of MLNs are mapped on the mesentery. (B) Proportions of MLNs taken from each location of UC mesentery. The majority of lymph nodes were taken from the mesorectum (42.9%). (C) Proportions of MLNs taken from each location of Crohn’s disease (CD) mesentery. The majority of lymph nodes were taken from the ileocolic region (87.5%). (D) Proportions of 16S PCR positive MLNs taken from each location of ulcerative colitis (UC) mesentery. (E) Proportions of 16S PCR positive MLNs taken from each location of CD mesentery.

Figure 2: Relative abundance of predominant bacterial phyla in mesenteric lymph nodes (MLNs) from inflammatory bowel disease (IBD) patients. There was a distinct difference in the profile of phyla from MLNs of Crohn’s disease (CD) and ulcerative colitis (UC) patients. S = sample number, M = merge of samples of same node.

Figure 3: Abundance of phyla in pooled mesenteric lymph nodes (MLNs) of Crohn’s disease (CD) and ulcerative colitis (UC) patients. (A) Quantities (abundance %) of major bacterial phyla, Firmicutes, Bacteroides, Proteobacteria and Actinobacteria in pooled MLNs of UC and CD patients. (B) Statistical analysis of phyla abundance with 95% confidence interval (CI). MLNs of UC patients had higher proportions of Firmicutes (Mean ± 95% CI: 52.8% ± 6.73% vs. 18.6% ± 4.65%; p<0.0001, independent t-test) while MLNs of CD patients had higher proportions of Proteobacteria (36.1% ± 11.06% vs. 11.8% ± 3.63%; p=0.005, Mann-Whitney U test) and other type bacteria, comprising mainly Fusobacteria (7.3% ± 4.98% vs. 0.3% ± 0.15%; p<0.0001, Mann-Whitney U test). Independent t-tests were used to compare normally distributed (parametric) data, while Mann-Whitney U tests were used to compare non-parametric data. (C) The ratio of Firmicutes to Bacteroidetes in MLNs from UC (1.613) and CD (0.705) patients.

Figure 4: Principal coordinate analysis (PCoA) of Unifrac difference. The chart is based on unweighted unifrac distances and displays variation between cohort samples.

Figure 5: Microbial diversity of mesenteric lymph nodes (MLNs) from Crohn’s disease (CD) and ulcerative colitis (UC) patients. Shannon Diversity Indices for MLNs of UC and CD patients defined using data from 16S sequencing data at species level. Data are presented as mean ± 95% confidence intervals (CI). MLNs from UC patients demonstrated significantly greater microbial diversity than those of CD patients (1.95 ± 0.0035 vs. 0.79 ± 0.0018; p<0.0001, Mann-Whitney U test). As data were found to be non-parametric (not-normally distributed), Mann-Whitney U tests were used to compare data.
**Table 1:** Demographics of patients with PCR positive lymph nodes. All patients received antibiotics at induction of anaesthesia.

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<th>Age at dx</th>
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Figure 2
Figure 3

A

B

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<th>CD</th>
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<td>Firmicutes</td>
<td>52.8% (46.07 - 59.54%)</td>
<td>18.6% (13.95 - 23.18%)</td>
<td>0.000 (t-test)</td>
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<td>Bacteroidetes</td>
<td>30.6% (23.52 - 37.74%)</td>
<td>32.0% (25.74 - 38.25%)</td>
<td>0.756 (MW-U)</td>
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<tr>
<td>Proteobacteria</td>
<td>11.8% (8.17 - 15.34%)</td>
<td>36.1% (25.04 - 47.12%)</td>
<td>0.005 (MW-U)</td>
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<td>Actinobacteria</td>
<td>4.6% (2.68 - 6.43%)</td>
<td>6.1% (4.04 - 8.11%)</td>
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<td>Others</td>
<td>0.3% (0.11 - 0.40%)</td>
<td>7.3% (2.32 - 12.27%)</td>
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C

UC  /  = 1.613

CD  /  = 0.705
Figure 4
Figure 5

Shannon Diversity Index

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<td>Crohn’s disease</td>
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