

ORIGINAL ARTICLE

Structural robustness of the gut mucosal microbiota is associated with Crohn's disease remission after surgery

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ABSTRACT

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Objectives Preventing postoperative recurrence after ileocolonic resection (ICR) for Crohn's disease (CD) is challenging. Defining the disturbances of the microbial composition and community structure after ICR and their link with early disease recurrence is crucial.

Design Microbiota composition (fingerprinting and 16S rDNA sequencing) and community structure (correlation networks of bacterial species) were assessed from ileal mucosa sampled in 20 patients undergoing ICR and 6 months later during endoscopy from above (neoterminal ileum) and below (subanastomotic colon) the surgical anastomosis.

Results ICR had a dramatic effect on gut microbial ecosystem. At surgery, CD mucosa harboured a dysbiotic microbiota with high proportions of α/β Proteobacteria and Bacilli. Six months later, half of the patients had recurrent lesions at ileocolonoscopy and presented higher numbers of Lachnospiraceae. Recurrence of endoscopic lesions was associated with enrichment in Enterococcus durans while patients in remission had increased proportions of Dorea longicatena and Bacteroides plebeius. Structural differences were striking between recurrence and remission microbiota; while the microbiota of patients with CD recurrence exhibited a loose community structure, the microbiota of patients in remission displayed communities that were robustly correlated to each other. Microbiota colonising the neoterminal ileum and subanastomotic colon 6 months after ICR only differed in patients with recurrence. **Conclusions** ICR modifies the gut microbiome. Remission after 6 months was associated with homogenous bacterial distribution around the anastomosis. Community structure and bacterial

networks highlight target species, including Faecalibacterium prausnitzii and Ruminococcus gnavus, which may allow precise modulations of the overall microbial ecosystem towards remission pattern.

INTRODUCTION

Crohn's disease (CD) is a chronic inflammatory disease of the intestine with an increasing incidence worldwide.¹ It is a multifactorial disease where host genetic impairments and environmental triggers lead to the onset and maintenance of inappropriate host immune responses at the gut mucosa interface.

More than 70 genetic loci have been linked to CD phenotype transmission,² and interestingly,

Significance of this study

What is already known on this subject?

- Recurrence of Crohn's disease (CD) after ileocolonic resection (ICR) is frequent.
- Luminal flux and gut microbiota are involved in ► this setting.

What are the new findings?

- ICR for CD strongly modifies the gut mucosal ► microbiota.
- ► Spatial homogeneity in bacterial composition around the anastomosis is associated with maintenance of endoscopic remission.
- Microbial community structure and bacterial ► species assemblages are cohesive during remission while a loose structure, together with a cluster of bacteria from the Proteobacteria phylum, is associated with endoscopic recurrence.

How might it impact on clinical practice in the foreseeable future?

- Bacterial recolonisation following ICR might ► pave the way towards remission or recurrence.
- ► Community structure and bacterial networks highlight target species that may allow precise modulations of the overall microbial ecosystem towards remission pattern.

most of them involve several key host defences (NOD2, IL23R, ATG16L1, FUT2). Environmental factors, especially smoking habits, are also involved. Results obtained from experimental mouse models³ have shed light on different mechanisms emphasising a failure in the host-microbiota crosstalk. Host-microbiota interactions sustain numerous physiological functions such as the maturation of host intestinal/immune cells (lymphoid tissue, Paneth cells) and the settlement of a commensal microbiota, thereof any alteration of this intimate relationship might modify the natural makeup of the 'healthy' microbiome. The use of immunologically impaired animal models has emphasised the importance of this relationship. For instance, TRUC mice (T-bet^{-/-} RAG2^{-/-} UC model) are able to shape an aberrant microbiota leading to



disease $^{4\ 5}$ and the transfer of their microbiota to a wild type host was shown as well to induce colitis. $^{5\ 6}$

Corticosteroids, thiopurines and monoclonal antibodies against tumour necrosis factor- α are efficient to modulate inflammatory activity, but surgery is still frequently required.⁷ Strikingly, severe inflammatory lesions recur in the neoterminal ileum in the months following surgery in more than half of the patients operated on to remove the inflamed ileum.⁸ ⁹ The crucial link between gut microbiota and CD recurrence was first suspected in this context two decades ago.¹⁰ ¹¹ In five patients with CD who had undergone resection with a diverting loop ileostomy proximal to an ileocolonic anastomosis, lesions recurred in the neoterminal ileum only once the bowel continuity was restored, which suggested that the microbiota and/or other components of the faecal stream may play a role.

The importance of the human gut microbiome in health and disease has been recognised for decades.^{12–15} In CD, it seems of great interest to investigate the mucosal microbiota that differs from faecal samples¹⁶ as it is in close contact to the disease process. In this study, we aimed at (i) assessing the impact of the ileocolonic resection (ICR) on the mucosa-associated microbiota, (ii) observing whether alterations differed between patients in remission or in recurrence after ICR, (iii) highlighting bacterial biomarkers of early recurrence of inflammatory lesions, and finally, (iv) deciphering bacterial community structure at the mucosal level in patients.

Microbial profiling and in-depth sequencing techniques were applied to analyse the gut microbiota. Since ecological constraints in the gut (chemical, immunological or trophic) are crucial in shaping the microbiota assembly and necessary for the development of a structured and functional ecosystem, we analysed the microbiota for its composition, diversity and structure in patients suffering from CD before and 6 months following operation to track out any change in microbiota composition and decipher the microbial recolonisation process following ICR.

PATIENTS AND METHODS

Patients and samples

Twenty patients with active CD were included in this study. They were part of a double-blind, randomised, placebocontrolled, 6-month clinical study,¹⁷ aiming at assessing the efficacy of the probiotic strain Lactobacillus johnsonii LA1 to decrease the endoscopic recurrence rate 6 months after surgery. CD was diagnosed on the basis of standard clinical, radiographic, endoscopic and pathological criteria. The patients had to be addressed to clinical centres for ileal or ICR. Patients who received antibiotics during the previous three months or receiving antibiotics for >2 weeks after surgery were not eligible. Other exclusion criteria included (i) total or subtotal colectomy, intestinal bypass or stricturoplasty, stomy, carcinoma resection, or abscess drainage; (ii) treatment with aminosalicylates or immunosuppressants >3 weeks following surgery; (iii) any other disease or condition that might interfere with the study assessments (as judged by the investigator); and (iv) child-bearing potential in women not using effective contraception; and pregnant or lactating women.

At time of surgery (month 0 (M0)), patients provided faecal samples before colonic cleansing. Biopsies were sampled using biopsy pinches from the fresh resected tissue at the ileal level in macroscopically ulcerated regions. Tissue specimens were also collected (4 cm² surface). Patients were examined 6 months after ICR (month 6 (M6)), provided faecal samples and an ileo-colonoscopy was performed. Biopsies were sampled from the

neoterminal ileum (above the anastomosis) and the subanastomotic colon (below the anastomosis). Endoscopic status was determined (recurrence of inflammatory lesions (REC); remission (REM)). Endoscopic recurrence was defined by a Rutgeerts score >2.¹⁸ All samples were immediately dry frozen in nitrogen or dry ice and further kept at -80° C before analysis. At 1-year follow-up, endoscopic and clinical recurrence was assessed by clinicians.

DNA preparation

Total DNA was extracted from faeces and biopsy samples as previously described.¹⁹ DNA concentration and integrity were determined spectrophotometrically and bacterial 16S rRNA gene was amplified using either primers targeting the V6-V8 region (GCclamp-U968 and L1401) for fingerprinting²⁰ or the V3-V4 region for sequencing (V3fwd: 5'-TACGGRAG GCAGCAG-3', V4rev: 5'-GGACTACCAGGGTATCTAAT-3').²¹

Fingerprinting of the dominant microbiota

Sequence-specific fingerprints were obtained with Temporal Temperature Gradient Gel Electrophoresis (TTGE) as previously described.¹⁹ TTGE profiles were analysed with Gel Compar software V.2.0 (Applied Maths, Kortrijk, Belgium). Similarity indexes (Pearson correlation method with unweighted pair group method with arithmetic mean (UPGMA) method) were calculated for each pair of profiles. Means were compared using paired Student's t test when the variances were equal, and otherwise using Wilcoxon's test.

Microbial composition assessment by high-throughput sequencing

Ten patients (ie, 46 samples: 26 mucosal samples and 20 faecal samples; see online supplementary table S1) were selected from the main cohort and their microbiota composition was further deeply analysed by 454 pyrosequencing using GS-FLX-Titanium technology following manufacturer's instructions (Roche) for mucosal samples or MiSeq (Illumina) for faecal samples. Faecal samples from one patient provided low number of sequences and were discarded. Sequences were trimmed for adaptors and PCR primers removal and binned for a minimal sequence length of 250 bases and an average minimal base quality threshold set at 25. Resulting sequences (in average 2915 for mucosal samples and 16 532 for faecal samples) were clustered at 97% of identity using OIIME²² and usearch (V.5.2.32) pipeline.²³ Operational taxonomic units (OTUs) with abundance lower than four reads were discarded. Chimera detection and removal were assessed using GOLD reference database and uchime.²³ OTUs were assigned to closest taxonomic neighbours using SEQMATCH and up to date 16S rRNA gene RDP database (release 10, update 30).²⁴ Alpha diversity (Shannon index) was analysed on the OTU table rarefied to 1000 reads.

All statistical analyses were performed using *R* program,²⁵ ade4²⁶ and *igraph* packages.²⁷ Statistics was assessed using either a Wilcoxon's test or the analysis of variance (p<0.05). If required, p values were adjusted for multiple testing effects using the Bonferroni correction. Principal component and between-class analyses were performed to map each sample based on their overall microbial composition and to assess similarities between individual's microbiota. Robustness of each clustering result was assessed using a Monte Carlo rank test (p<0.05). The network analysis was carried out with *SparCC*²⁸ (bootstrap n=500) on the most abundant OTUs (maximum abundance of a particular OTU within a sample $\geq 0.125\%$). SparCC correlation scores ≤ -0.5 or ≥ 0.5 with a p ≤ 0.001 were

kept for networks inference. The analysis was conducted as described in the *igraph* package. Network visualisation was performed using the Fruchterman–Reingold layout algorithm. Clusters were detected with *cluster_louvain()* function.²⁹ The robustness of each cluster was tested (Wilcoxon test) by comparing the number of OTU edges within a defined cluster to the number of OTU edges outside this cluster.

RESULTS

Twenty patients were included in the study. Six months after surgery, 10 patients had developed recurrence of CD lesions (REC) visible during endoscopic examination while the other half remained in endoscopic remission (REM) according to the Rutgeerts score. A total of 337 OTUs was detected in mucosal samples with an average of 88 ± 34 OTUs per sample, while 1225 OTUs constituted the faecal microbiota with an average of 327 ± 124 OTUs per sample. Bacterial phylotypes (or OTUs) were defined as groups of sequences sharing at least 97% of similarity and OTUs' representative sequences were phylogenetically affiliated to their closest relative bacterial species.

ICR for CD affects the mucosal microbiota, which is also depicted in faecal samples

ICR had a striking effect on gut microbial composition. Bacterial profiles of the dominant mucosal microbiota at 6 months after ICR (M6) differed markedly from the profiles observed at time of surgery (M0). The average TTGE pairwise similarity percentages were 55.1%±25.0% between ileal mucosa at M0 and the neoterminal ileum at M6, and 48.1% ±28.9% between ileal mucosa at M0 and subanastomotic colon at M6. The composition of mucosal microbiota, analysed by 16S rRNA gene pyrosequencing, was strongly affected by ICR (figure 1). Samples collected at M0 were significantly enriched in bacteria belonging to α-/β-Proteobacteria and Bacilli (Streptococcus), whereas M6 samples had higher proportions of Clostridia (see online supplementary table S2 and supplementary figure S1). Lachnospiraceae family displayed the largest variation between M0 and M6 with percentages ranging from 7.39%±2.79% (M0) to 31.01%±7.82% (REC) and 27.17% $\pm 4.78\%$ (REM). Top-down taxonomic analysis emphasised that Dorea and Blautia genera accounted mostly for this variation. Strikingly, Porphyromonadaceae levels were also increased in M6 samples, but only in patients with endoscopic recurrence. At OTU level, 27 bacterial species were differentially abundant between M0 and M6 sampling times (table 1; M0=17 OTUs and M6=10 OTUs).

At M0, the intestinal mucosa of patients with CD harboured high proportions of unusual, facultative anaerobic or aerobic gut bacteria such as *Streptococccus mitis*, *Undibacterium oligocarboniphilum*, *Sphingomonas melonis* and *Gemella haemolysans*. On the other hand, 6 months after ICR, mucosal samples exhibited higher levels of anaerobic bacteria belonging to Lachnospiraceae: *Clostridium nexile*, *Blautia wexlereae* and *Dorea longicatena*. Bacteroidaceae, Enterobacteriaceae and Bifidobacteriaceae were not impacted by ICR.

No difference was observed within the α -diversity between M0 and M6 samples (Shannon index, M0: 2.538 ± 0.3179 and M6: 3.030 ± 0.111 , p>0.05).

ICR also impacted faecal bacterial populations, and the average TTGE profiles similarity percentage between M0 and M6 was $41.3\% \pm 34.1\%$. The inter-individual variability was high and similarity ranged from 0% to 94.9% between M0 and M6. No significant differences were noticeable depending on the endoscopic status at 6 months (remission or recurrence).

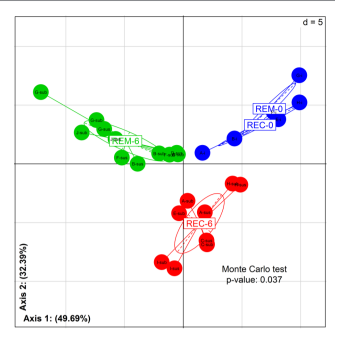


Figure 1 Impact of ileocolonic resection on mucosal microbiota: initial stage compared with 6 months later. Between class analysis (axis 1 and 2) visualising results from the principal component analysis based on the operational taxonomic unit composition in the microbiota of patients before and after surgery (M0 vs M6). Samples were clustered according to remission/recurrence (REM/REC) and sampling times (M0 and M6). Robustness of the clustering result was assessed using a Monte Carlo ranking test. Samples at M0 (REM-0/REC-0) are in blue. Samples at 6 months for patients in remission (REM-6) are in green. Samples at 6 months for patients in recurrence (REC-6) are in red. At M6: sus, neoterminal ileum; sub, subanastomotic colon.

Average TTGE profiles similarity percentages were $45.1\% \pm 23.4\%$ for patients in remission and $58.1\% \pm 30.1\%$ for patients with endoscopic recurrence.

Towards a link between bacterial dysbiosis and postoperative recurrence

The mucosa-associated microbiota was analysed in detail by 16S rRNA gene sequencing in 10 patients: 5 having persistent endoscopic remission and 5 with recurrence. No significant difference was observed between REC and REM α -diversity (Shannon index, REC: 2.907±0.513 and REM: 3.131±0.485, p>0.05). The mucosal microbiota differed markedly between these two groups of patients. Remission status was characterised by an increased number of bacteria belonging to four genera (*Bacteroides, Dorea, Ruminococcus* and *Dialister*; see online supplementary table S3) and mainly the species *D. longicatena* and *Bacteroides plebeius* (table 2). On the other hand, *Gemmiger formicilis, Enterococcus durans* and *Ruminococcus lactaris* were significantly increased in REC patients as were bacteria belonging to Bacilli class.

All but 4 of these 14 specific species (*Clostridium bolteae*, *Bacteroides dorei*, *Ruminococcus schinkii* and *Clostridium thermocellum*) were also detected in faecal samples from the patients although at lower percentages. A similar trend in being over-represented or under-represented in remission or recurrence was observed in faeces but statistical significance was not reached. We searched for bacterial biomarkers of the clinical outcome in faecal samples of patients at M0 before surgery and detected significant differences in the bacterial composition of faecal samples at M0 between patients that appeared to stay in

Table 1 D	ifferentially represented OTUs between samples colle Closest species (% of sequence similarity)	Time 0	REC6	REM6	p Value
					<u> </u>
28	Streptococcus mitis (100%)	11.41±5.6	1.63±1.0	0.44±0.2	0.009
58	Undibacterium oligocarboniphilum (100%)	4.58±2.5	0.8±0.3	0.30±0.1	0.021
68	Sphingomonas melonis (100%)	1.81±1.0	0.41±0.2	0.11±0.1	0.027
64	Gemella haemolysans (100%)	1.78±0.8	0.30±0.2	0.06±0.0	0.006
87	Clostridium lituseburense (98%)	1.70±0.9	0.00 ± 0.0	0.01±0.0	0.006
91	Thermus scotoductus (100%)	1.53±0.7	0.73±0.4	0.06±0.0	0.028
76	Mesorhizobium australicum (99%)	1.41±0.7	0.18±0.1	0.07±0.0	0.008
124	Granulicatella adiacens (100%)	1.21±0.7	0.18±0.1	0.03±0.0	0.028
250	Staphylococcus warneri (100%)	0.72±0.4	0.12±0.0	0.06±0.0	0.041
142	Granulicatella elegans (100%)	0.63±0.3	0.04±0.0	0.02±0.0	0.006
224	Bradyrhizobium yuanmingense (100%)	0.56±0.2	0.07±0.1	0.03±0.0	0.002
227	Sediminibacterium salmoneum (96%)	0.46±0.2	0.02±0.0	0.05±0.0	0.011
226	Sediminibacterium salmoneum (98%)	0.38±0.2	0.00 ± 0.0	0.02±0.0	0.032
243	Pseudomonas flavescens (100%)	0.28±0.2	0.01±0.0	0.01±0.0	0.027
256	Rhodospirillum rubrum (86%)	0.21±0.1	0.06±0.0	0±0	0.035
233	Turicibacter sanguinis (100%)	0.08±0.0	0±0	0.02±0.0	0.017
15	Eubacterium coprostanoligenes (92%)	0.03±0.0	0±0	0±0	0.033
10	Clostridium nexile (100%)	0.14±0.0	4.79±1.5	0.83±0.4	0.005
9	Blautia wexlereae (100%)	0.32±0.1	2.81±1.2	4.17±0.8	0.036
12	Dorea longicatena (100%)	0.40±0.2	0.47±0.2	3.18±0.7	0.002
53	Sutterella wadsworthensis (100%)	0.08±0.0	0.27±0.1	0.64±0.2	0.025
199	Ruminococcus schinkii (97%)	0.02±0.0	0.03±0.0	0.24±0.1	0.003
288	Bacteroides dorei (99%)	0±0	0.03±0.0	0.14±0.0	0.035
241	Desulfotomaculum guttoideum (98%)	0.04±0.0	0.00±0.0	0.13±0.0	0.01
1	Bacteroides plebeius (100%)	0.25±0.2	0±0	2.41±0.9	0.029
153	Clostridium bolteae (95%)	0±0	0±0	0.49±0.2	0.034

Shadeless and grey-shaded lines indicate respectively increased OTUs in samples either collected at time 0 or time 6. Statistics was assessed by analysis of variance. Only significant data (p<0.05) are presented.

ICR, ileocolonic resection; OTUs, operational taxonomic units; REC. recurrence; REM, remission.

remission after ICR compared with those who finally experienced endoscopic recurrence. At the OTUs level, four specific molecular species had a biomarker potential, two for remission prediction and two for recurrence prediction (figure 2A). While

Table 2Differentially represented OTUs between recurrence and
remission samples collected at 6 months after ICR

	Closest species			
OTUs	(% of sequence similarity)	REC6	REM6	p Value
20	Gemmiger formicilis (100%)	1.58±0.7	0.77±0.5	0.038
335	Enterococcus durans (100%)	0.53±0.4	0.00 ± 0.0	0.011
78	Ruminococcus lactaris (96%)	0.32±0.1	0.02±0.0	0.015
12	Dorea longicatena (100%)	0.47±0.2	3.18±0.7	0.011
1	Bacteroides plebeius (100%)	0±0	3.13±1.2	0.013
153	Clostridium bolteae (95%)	0±0	0.49±0.2	0.006
101	Bacteroides dorei (99%)	0.09±0.0	0.47±0.1	0.019
208	Butyricimonas virosa (98%)	0.00±0.0	0.36±0.2	0.05
199	Ruminococcus schinkii (97%)	0.03±0.0	0.24±0.1	0.005
97	Bilophila wadsworthia (100%)	0.01±0.0	0.15±0.1	0.05
237	Bacteroides coprocola (100%)	0±0	0.14±0.1	0.013
241	Desulfotomaculum guttoideum (98%)	0.00 ± 0.0	0.13±0.0	0.013
312	Bacteroides nordii (100%)	0±0	0.12±0.1	0.028
62	Clostridium thermocellum (88%)	0.01±0.0	0.10±0.0	0.021

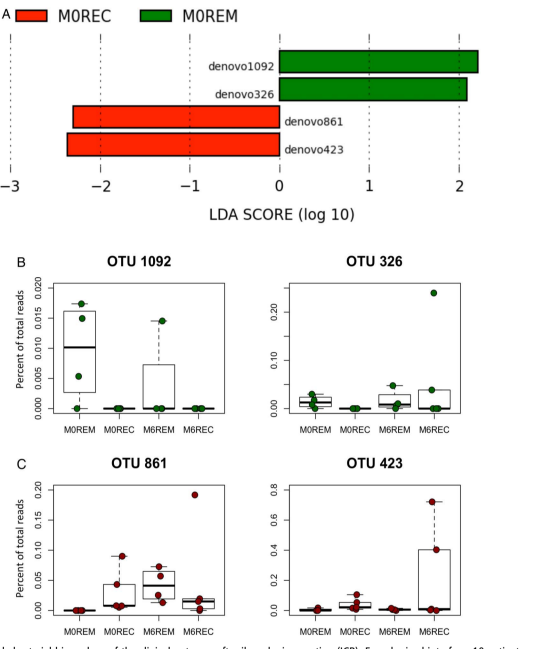
Shadeless and grey-shaded lines indicate respectively increased OTUs in patients with disease recurrence or remission. Statistics was assessed by Wilcoxon test. Only significant data (p<0.05) are presented. ICR, ileocolonic resection; OTUs, operational taxonomic units; REC. recurrence; REM,

ICR, ileocolonic resection; OTUs, operational taxonomic units; REC. recurrence; REM, remission.

detection of *Coprococcus catus* L8 and a relative of Clostridiales bacterium DJF_B152 (*Butyricicoccus* genus) in faeces of patients at M0 were significantly associated with remission at M6, the presence of *Proteus mirabilis* and of a relative of *Eubacterium rangiferina* were highlighted as biomarkers of future recurrence in patients undergoing ICR (OTUs denovo1092, 326, 423 and 861, respectively). Moreover, OTU denovo1092 and denovo326 were significantly more represented at M0 in patients being in remission at M6 than in REC patients (p=0.034 and 0.044, respectively; figure 2B).

Postoperative recurrence: from microbial dysbiosis to microbiota destructuration

Unravelling interactions among different species within ecosystems is essential to understand how microbiota responds/adapts to environmental changes. Here we describe the use of SparCC correlation-based approach to infer species association (network) for the mucosal microbiota at each clinical status time points (M0, REC, REM). Species co-occurrence and exclusion are assessed at the overall ecosystem level, and groups of highly correlated OTUs are further defined as clusters. Clusters may relate to metabolic entities involved in trophic chains or immune reactions, specific bacterial features such as aggregation, signalling, biofilm formation or specific habitat characteristics (pH, mucins). Out of 387 total OTUs, only 15 made up the mucosal microbiota community structure of the patients with CD before operation (M0, see online supplementary figure S2 and supplementary table S6). Eighty-seven per cent of these OTUs (n=13) were common to the mucosal microbiota



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Figure 2 Early bacterial biomarkers of the clinical outcome after ileocolonic resection (ICR). Faecal microbiota from 10 patients was analysed at M0 and 6 months after ICR. (A) Specific bacterial operational taxonomic units (OTUs) that may predict at M0 the clinical outcome following ICR were deciphered applying the LEfSe algorithm.⁴⁷ The LEfSe algorithm uses the non-parametric factorial Kruskal–Wallis sum-rank test to detect features with significant differential abundance with respect to the clinical outcome; biological significance is subsequently investigated using a set of pairwise tests using the (unpaired) Wilcoxon rank-sum test. As a last step, LEfSe uses linear discriminant analysis to estimate the effect size of each differentially abundant feature (ie, bacterial OTUs). (B) Boxplot of the distribution (per cent of total reads) of the four OTUs highlighted in the LEfSe analyses in the different patients' groups. M0 bacterial biomarkers associated with remission at M6 are in green. M0 bacterial biomarkers of future recurrence in patients undergoing ICR are in red. M0, samples at time of surgery; REC, recurrence; REM, remission; LDA, linear discriminant analysis.

structure inferred for the patients 6 months after ICR (see online supplementary figure S3). Within the M0 structure, no distinct and significant bacterial cluster was identified.

The mucosal microbiota structure of patients in remission after 6 months (REM) was more complex and better organised than the structure inferred for patients suffering from disease recurrence (REC) (figure 3). The REM microbiota structure was composed of 192 OTUs, whereas 102 OTUs were found in the REC structure (see online supplementary tables S4 and S5, respectively) with 82 OTUs common to both structures. The number of inter-OTUs correlations was five times higher for REM microbiota (2142) compared with REC one (436). Both REM and REC network structures were made up of five significant clusters. Each microbial cluster was well distributed in the cohort of patients, either patients in remission (see online supplementary figure S4A) or with recurrence (see online supplementary figure S4B).

Among the five REM microbial clusters, clusters I, III and IV were made up of bacteria mainly related to Firmicutes phylum (70%, 67% and 63%, respectively). Main species in cluster I

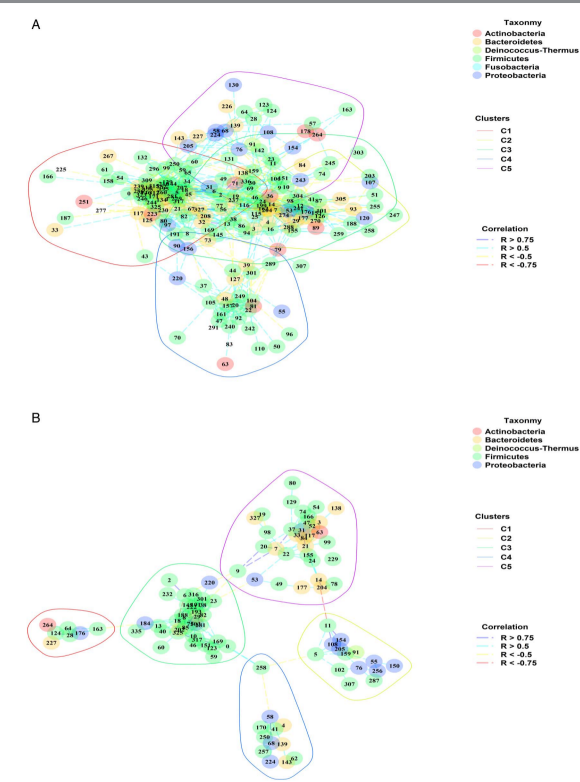


Figure 3 Correlation network organisation inferred from the microbiota composition of patients with Crohn's disease in remission and in recurrence 6 months after ileocolonic resection (ICR). (A) and (B) respectively depict the correlation network structures inferred from the microbiota composition of patients in remission or with recurrence 6 months after ICR. Each number relates to operational taxonomic unit (OTU) indexes. OTU phylogeny at phylum level is described by colours filling each OTU dot. Correlation scores are indicated by the colour of the edges linking OTU pairs. Polygons delineate each cluster (C1–C5) identified in the network. (A and B) The red, yellow, green, blue and purple polygons delineate clusters I, II, III, IV and V.

related to *Faecalibacterium prausnitzii*, *Ruminococcus bromii* and *B. plebeius*, whereas *G. formicilis* was the main driver of cluster III. Interestingly, cluster I represented a tight community, mainly constituted with OTUs from the Ruminococcaceae and Lachnospiraceae families, with each OTU sharing in average 16.45 correlations with other OTUs of the cluster. The cluster

Gut microbiota

II was taxonomically more diverse and composed of Firmicutes (49%), Bacteroidetes (34%) and Proteobacteria (11%) phyla. The most abundant OTUs related to *Dorea formicigenerans, Sutterella wadsworthensis* and *Bacteroides uniformis*. Finally, cluster V grouped OTUs affiliated to Firmicutes (42%), Proteobacteria (31%), Bacteroidetes (15%) and Actinobacteria (8%). Firmicutes population showed balanced proportions of Clostridia and Bacilli classes with OTUs related to *Clostridium, Blautia, Streptococcus, Gemella* and *Granulicatella* genera (as opposed to Clostridia class alone in cluster I).

On the other hand, the REC microbiota structure included five different clusters that were looser than in the REM microbiota structure (average 4.27 correlations/OTUs in REC vs 11.16 correlations/OTUs in REM). The cluster II was composed at 50% with Proteobacteria (α -/ β -Proteobacteria) and OTUs were affiliated to Comamonas and Pelomonas genera. The cluster III grouped Clostridium hylemonae, Ruminococcus gnavus and E. durans, a group of species specifically involved in bile acids (BAs) hydroxylation and bile salts hydrolysis. F. prausnitzii was only detected once in cluster III, and out of five correlations shared by F. prausnitzii within this cluster, four were negative. Similarly to the microbiota of patients in remission, two clusters enriched in Firmicutes were detected, one mainly containing Clostridia (cluster IV) and the other Bacilli (cluster I). Finally, the cluster V, driven mainly by Bacteroides caccae and Eubacterium halii, grouped bacteria that were well distributed in terms of phyla repartition.

Colonisation of the neoterminal ileum and subanastomotic colon after ICR

Comparison between the microbiota colonising the neoterminal ileum and the subanastomotic mucosa highlighted an average TTGE profile similarity percentage of $82.6\% \pm 21.8\%$. Interestingly, dominant mucosal microbiota in the neoterminal ileum and subanastomotic colon were significantly closer within the patients in remission at M6 than in patients with recurrence (REM: 92.2 ± 2.7 ; REC: $77.9\%\pm24.7$; p=0.02). A similar trend was observed with the pyrosequencing analysis (figure 4). Species from *Dorea* and *Brevundimonas* accounted for those explaining this difference. *D. longicatena* (REC-sub: 0.265 ± 0.247 and REC-sus: 0.636 ± 0.314) and *Brevundimonas vesicularis* (REC-sub: 0.022 ± 0.022 and REC-sus: 0.203 ± 0.096) were respectively 2.4-fold and 9.2-fold increased in the neoterminal ileum mucosa compared with the subanastomotic one in patients with recurrence.

DISCUSSION

At time of surgery, the mucosal microbiota of patients with active CD was rich in α -/ β -Proteobacteria, Bacilli and more largely in bacteria that are not usually found in a healthy gut microbiota. This is in line with previous studies.^{30 31} While not affecting the gut microbial diversity, ICR markedly modified the gut mucosal microbiota. Following ICR, a fourfold increase of Lachnospiraceae taxa was observed showing deep modifications within the gut ecosystem. Lachnospiraceae abundance has previously been noticed to fluctuate at both faecal and mucosal levels in patients with CD with active disease.³² ICR also led to an increase in Porphyromonadaceae levels in patients with endoscopic recurrence at 6 months, and this was also observed in the jejunal mucosa of a mouse model of resection.³³

It is noteworthy that microbial assemblage and recolonisation process after ICR were associated to disease relapse or maintenance of remission. Indeed, mucosal microbiota comparison of patients being either on disease remission or recurrence

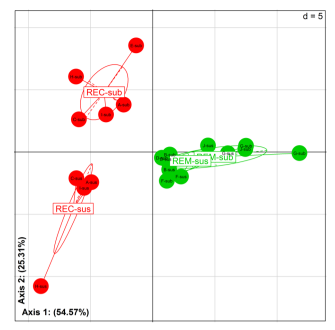


Figure 4 Microbiota recolonisation after ileocolonic resection. Between class analysis (axis 1 and 2) visualising results from the principal component analysis based on the operational taxonomic unit composition in the microbiota of patients 6 months after surgery. Samples were clustered according to remission/recurrence (REM/REC) and location around the anastomosis (sus/sub). Samples of patients in remission are in green (REM-sus and REM-sub). Samples of patients with endoscopic recurrence are in red (REC-sus and REC-sub). sus, neoterminal ileum; sub, subanastomotic colon.

6 months after ICR has shed light on different gut ecosystem composition and structures. Our results suggest that recovering an eubiotic microbiota, mainly composed of Firmicutes and Bacteroidetes, is a crucial step towards remission in patients with CD after resection. Moreover, homogeneity in bacterial composition at the anastomosis location might be associated with homeostasis and remission in these patients. Compositional differences mainly related to an increase in Bacilli abundance during disease recurrence, whereas bacteria belonging to Bacteroides, Dorea, Ruminococcus and Dialister genera were more prevalent in patients in remission. We also assessed whether specific bacterial biomarkers in faecal samples may predict, at the time of surgery, the clinical outcome of patients after ICR. Two OTUs, both being relatives of butyrateproducing bacteria (ie, C. catus and Butyricicoccus sp.), were more abundant in M0 faecal samples of patients being in remission at 6 months and undetectable at M0 in the microbiota of patients with endoscopic recurrence at M6. On the other hand, two OTUs were specific predictors of recurrence following ICR: E. rangiferina and P. mirabilis. It is noteworthy that P. mirabilis has been shown to be able to promote colitis in susceptible animals,⁴ but also to induce robust interleukin-1b secretion via the NLRP3 inflammasome, hence promoting inflammation.³⁴

In order to study the structure and thus the interactions among the bacterial components of mucosal microbiota, we built a correlation network based on the relative abundance of OTUs among the different samples. The gut microbiota structure differed significantly whether patients' disease evolved towards maintenance of remission or recurrence of lesions. A major result is the identification of a complex mucosal ecosystem made of gut commensals and associated with disease remission. At the opposite, the microbiota structure of patients displaying CD recurrence appeared less organised. In a recent study, Levy and Borenstein³⁵ applied systems biology and metabolic modelling to highlight forces driving species composition in the gut microbiome. This metabolic-based prediction of species interaction showed that habitat is the dominant structuring force of the human microbiome. Here we show that CD activity is associated with a loose and non-cohesive mucosal microbiota structure that might impair proper ecosystem functions. On the other hand, a tightly correlated and cohesive species structure was observed in patients in remission 6 months after ICR.

Whatever the clinical status of patients 6 months after ICR, we observed at least one cluster containing mostly bacteria belonging both to the phylogenetic³⁶ and the metagenomic core³⁷ (REM: cluster II and III; REC: cluster V). These clusters may constitute a functional core required for the performance of the gut ecosystem. The microbial community structure of patients in remission also presented a cluster mainly driven by F. prausnitzii and R. bromii. Both species are thought to play a beneficial role in human health, partly through the production of short-chain fatty acids (SCFAs; butyrate mostly) and were more abundant in healthy subjects than in patients with CD.³⁸ Moreover, F. prausnitzii has been associated with clinical benefit in CD and its anti-inflammatory properties have been demonstrated in animal models.^{39 40} R. bromii is a strong degrader of resistant starches and proved to be unique in stimulating starch utilisation by other bacterial species,⁴¹ which may enhance the production of SCFAs in this cluster. On the other hand, the microbial structure of patients with recurrence 6 months after ICR exhibited a cluster enriched in Proteobacteria (α-/ β-Proteobacteria) and composed of unusual bacteria that were also detected in the mucosa of patients with IBD in several previous studies.^{30 31} Interestingly, another community (cluster III) specific of REC patients was composed of bacteria sharing specific enzymatic machinery associated with the metabolism of BAs. Hence, E. durans, a component of this cluster, exhibits bile salt hydrolase activity⁴² and may deconjugate BAs. Moreover, C. hylemonae and R. gnavus, main contributors of this cluster, both possess a high BA 7 α -dehydroxylating activity, leading to the production of secondary BAs. Of note, high concentrations of secondary BAs in faeces have been linked to the pathogenesis of colon cancer and more recently to CD,⁴³ and several studies also reported increased R. gnavus proportions in the mucosa and in faeces of patients with CD. $^{44-46}$

Ecological forces and key bacterial species that support these mucosal microbial communities still remain to be determined. Yet the presented global approach may allow a better understanding of microbial modulation by environmental factors (diet, prebiotics, antibiotics, pH, oxygen) but mostly pave the way towards general microbiota ecosystem modulation on the gut by regulating specific bacteria in the structure.

CONCLUSION

We highlighted specific bacteria, with either increased or decreased proportions, associated with early recurrence of CD inflammatory lesions. Moreover, we applied robust statistical analysis to assess the structure (bacterial phylotypes clusters and correlation networks) of the mucosa-associated microbial ecosystem in these patients and highlighted an overall ecosystem disruption in patients with CD recurrence after ICR.

Looking at the level of ecosystem structure may be a critical step in the development of therapeutic interventions targeting the microbiota in CD, including antibiotics and/or faecal transplantation.

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Contributors SM and PL contributed equally. PL, PS, PP, JD and PM designed the Research. All authors contributed to data acquisition, analysis and results interpretation. SM, PL and PM wrote the manuscript. All authors revised and approved the manuscript.

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Data sharing statement All data are fully available without restriction. All relevant data are within the paper and its supporting information files. DNA sequence reads from this study are available from the Sequence Read Archive (accession numbers available after acceptance) or by contacting the corresponding author.

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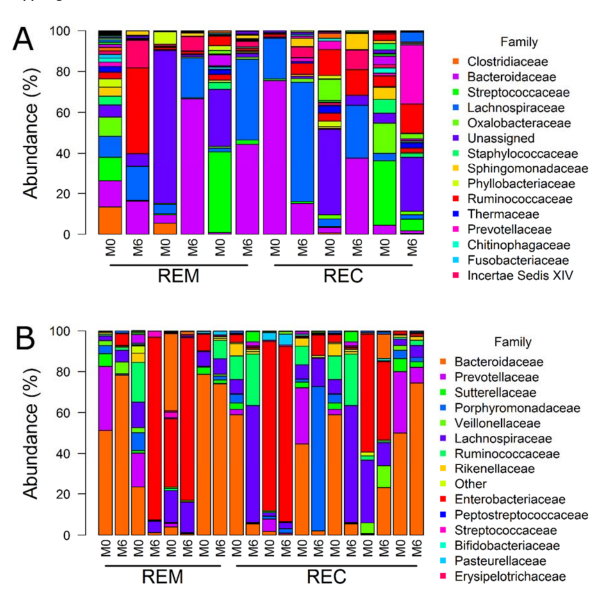
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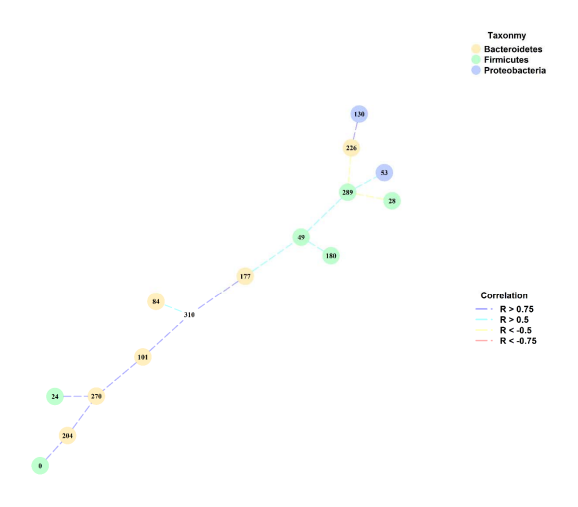


Supp. Figure 1: Distribution of bacterial families before and after ICR

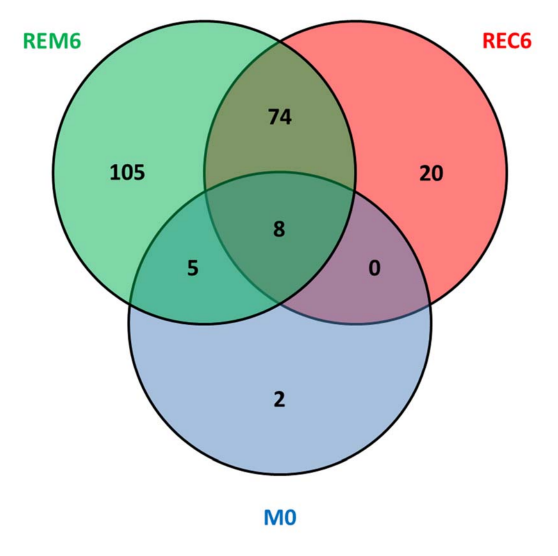
Only 15 main bacterial families are represented in the figures. Data are shown as Abundance (percent of total reads) for each family in A) Mucosal samples and B) Fecal samples.

M0: Month 0, time of surgery; M6: Month 6 after surgery; REM: Remission at M6; REC: Endoscopic recurrence at M6.

Supp. Figure 2: Structural analysis of the microbial communities detected in patients at time of surgery

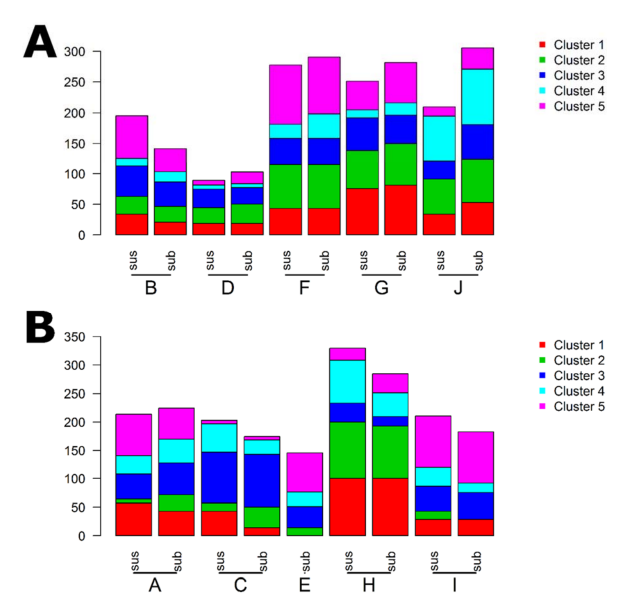


The correlation network analysis depicts the structure inferred from the mucosal microbiota composition of CD patients at ICR. Each number relates to OTU indexes. OTU phylogeny at phylum level is described by colors filling each OTU dot. Correlation scores are indicated by the color of the edges linking OTU pairs.



Supp. Figure 3: Venn diagram describing similarities between microbiota structures at OTU level

Diagram displaying the number of OTUs that are common or specific to microbiota structures inferred before and after surgery. M0: Samples at time of surgery; REC6: samples from patients with recurrence at 6 months; REM6: samples from patients in remission at 6 months.



Supp. Figure 4: Bacterial clusters distribution within each patient's microbiota 6 months after ICR in neoterminal ileum and subanastomotic colon.

The clusters abundance across samples was assessed using the occurrence of cluster specific OTUs. For each sample, the sum of OTUs occurrences found for a cluster was computed and then divided by the total number of OTUs identified in the cluster. Then, the relative clusters abundance (varying from 0 to 100%) was displayed as a cumulative bar plot.

A. Clusters distribution in the microbiota of patients in remission 6 months after ICR in both neoterminal ileum (sus) and subanastomotic colon (sub).

B. Clusters distribution in the microbiota of patients with endoscopic recurrence 6 months after ICR in both neoterminal ileum (sus) and subanastomotic colon (sub).

Patients	Age	Sex	Sample	s at MO	Sam	ples a	t M6	Endoscopic status at M6
			lleum	Feces	Sus	Sub	Feces	
В	NA	F	х	х	х	х	х	REM
D	26	F		х	х	х	х	REM
F	33	М	х	х	х	х	х	REM
G	28	F	х	х	х	х	х	REM
J	NA	М		х	х	х	х	REM
Α	NA	F	х		х	х		REC
С	31	F		х	х	х	х	REC
E	39	М	х	х	х	х	х	REC
н	27	F	х	х	х	х	х	REC
I	NA	М		х	х	х	х	REC
К	25	F		х			Х	REC

Supp. Table 1: Characteristics of patients and samples analysed by 16S rRNA gene sequencing

F: female. M: male. M0: time of surgery; M6: 6 month after surgery; Ileum: ileal biopsy sampled on to the resected tissue at M0. Sus: neoterminal ileum biopsy at M6. Sub: subanastomotic colon biopsy at M6. REM or REC: endoscopic remission (REM) or relapse (REC) six months after surgery. NA: Not available.

Taxonomic level	M0	REC6	REM6	p-valu
Bacilli	19,064±8,956	3,781±1,309	0,897±0,228	0,007
Lactobacillales	16,181±8,078	3,265±1,088	0,771±0,19	0,011
Streptococcaceae	14,196±7,14	2,415±1,011	0,617±0,168	0,011
Streptococcus	14,085±7,163	2,394±1,015	0,59±0,175	0,012
Alphaproteobacteria	8,326±2,956	2,621±1,658	0,526±0,191	0,008
Betaproteobacteria	7,583±2,699	2,152±0,782	1,926±0,828	0,017
Burkholderiales	7,56±2,686	2,083±0,737	1,926±0,828	0,016
Oxalobacteraceae	4,585±2,48	0,8±0,301	0,305±0,098	0,021
Clostridiaceae	3,242±2,214	0,042±0,042	0,008±0,004	0,039
Clostridiaceae 1	3,242±2,214	0,042±0,042	0,008±0,004	0,039
Bacillales	2,883±1,129	0,516±0,278	0,127±0,041	0,002
Staphylococcaceae	2,504±1,143	0,423±0,212	0,119±0,041	0,006
Sphingomonadales	1,862±1,009	0,434±0,208	0,116±0,056	0,028
Sphingomonadaceae	1,862±1,009	0,434±0,208	0,116±0,056	0,028
Sphingomonas	1,862±1,009	0,434±0,208	0,116±0,056	0,028
Carnobacteriaceae	1,839±0,977	0,224±0,124	0,057±0,031	0,013
Granulicatella	1,839±0,977	0,224±0,124	0,057±0,031	0,013
Gemella	1,784±0,794	0,302±0,188	0,063±0,029	0,006
Peptostreptococcaceae	1,768±0,946	0,009±0,007	0,018±0,01	0,008
Sporacetigenium	1,707±0,89	0,002±0,002	0,01±0,008	0,006
Deinococcus-Thermus	1,535±0,684	0,735±0,364	0,058±0,021	0,028
Deinococci	1,535±0,684	0,735±0,364	0,058±0,021	0,028
Thermales	1,535±0,684	0,735±0,364	0,058±0,021	0,028
Thermaceae	1,535±0,684	0,735±0,364	0,058±0,021	0,028
Thermus	1,535±0,684	0,735±0,364	0,058±0,021	0,028
Phyllobacteriaceae	1,415±0,684	0,178±0,107	0,067±0,04	0,008
Mesorhizobium	1,415±0,684	0,178±0,107	0,067±0,04	0,008
Sphingobacteria	0,934±0,503	0,096±0,06	0,084±0,049	0,02
Sphingobacteriales	0,934±0,503	0,096±0,06	0,084±0,049	0,02

Chitinophagaceae	0,844±0,457	0,082±0,058	0,077±0,049	0,021
Staphylococcus	0,719±0,433	0,121±0,047	0,056±0,021	0,041
Bradyrhizobiaceae	0,556±0,211	0,072±0,068	0,029±0,023	0,002
Bradyrhizobium	0,556±0,211	0,072±0,068	0,029±0,023	0,002
Pseudomonadaceae	0,282±0,175	0,006±0,006	0,011±0,011	0,027
Pseudomonas	0,282±0,175	0,006±0,006	0,011±0,011	0,027
Turicibacter	0,077±0,036	0±0	0,016±0,012	0,017
Lachnospiraceae	7,388±2,793	31,014±7,822	27,172±4,778	0,045
Dorea	1,081±0,438	2,445±0,726	5,996±0,943	0,001
Incertae Sedis XIV	1,008±0,262	12,127±3,771	7,617±1,228	0,026
Blautia	1,008±0,262	12,127±3,771	7,617±1,228	0,026
Porphyromonadaceae	0,349±0,207	7,803±3,416	1,125±0,254	0,035
Parabacteroides	0,272±0,174	7,284±3,479	0,641±0,211	0,046

Supp. Table 2: Differentially represented taxa between samples collected at time 0 and 6 months after ICR

Shade less and grey shade lines indicate respectively increased taxa in samples either collected at M0 or M6 in patients with endoscopic recurrence (REC6) or in remission (REM6). Statistics was assessed by ANOVA.

Таха	REC6	REM6	p-value
Bacilli	3,781±1,309	0,897±0,228	0,023
Lactobacillales	3,265±1,088	0,771±0,19	0,015
Enterococcaceae	0,597±0,361	0,009±0,008	0,011
Enterococcus	0,597±0,361	0,009±0,008	0,011
Bacteroidaceae	13,145±4,893	29,586±5,346	0,04
Bacteroides	13,145±4,893	29,586±5,346	0,04
Dorea	2,445±0,726	5,996±0,943	0,008
Ruminococcus	0,213±0,105	2,348±0,983	0,036
Dialister	0±0	0,88±0,516	0,028

Supp. Table 3: Differentially represented taxa between samples from patients in remission or with endoscopic recurrence at 6 months after ICR

Shade less and grey shade lines indicate respectively increased taxa in patients with disease recurrence (REC) or remission (REM). Statistics was assessed by Wilcoxon test.

Cluster		OTUs	Total	pos75	pos50	neg50	neg75	Phylum	Species	Similarity (%)
1	5.29e ⁻¹⁰	207	45	3	29	12	1	Firmicutes	Faecalibacterium prausnitzii	96.7
		5	38	4	21	12	1	Firmicutes	Ruminococcus bromii	99.8
		144	36	2	30	4	0	Firmicutes	Faecalibacterium prausnitzii	98.9
		18	33	0	13	16	3	Firmicutes	Clostridium hylemonae	98.5
		129	32	1	30	1	0	Firmicutes	Ruminococcus schinkii	96.7
		239	30	2	23	5	0	Bacteroidetes	Bacteroides plebeius	100
		6	30	2	22	4	1	Firmicutes	Faecalibacterium prausnitzii	98.8
		1	29	1	21	6	0	Bacteroidetes	Bacteroides plebeius	100
		45	29	4	22	2	0	Bacteroidetes	Bacteroides massiliensis	100
		152	28	0	24	4	0	Firmicutes	Faecalibacterium prausnitzii	89
		210	28	0	22	6	0	Firmicutes	Ruminococcus bromii	94.5
		171	26	0	22	4	0	Firmicutes	Coprococcus catus	95.3
		175	26	0	23	3	0	Bacteroidetes	Bacteroides dorei	95.3
		260	26	1	22	3	0	Bacteroidetes	Bacteroides massiliensis	99.6
		312	25	1	23	1	0	Bacteroidetes	Bacteroides nordii	99.7
		112	23	0	16	7	0	Firmicutes	Faecalibacterium prausnitzii	100
		146	23	0	20	3	0	Firmicutes	Clostridium viride	92.7
		315	20	0	19	1	0	Bacteroidetes	Bacteroides massiliensis	100
		286	19	0	17	2	0	Bacteroidetes	Bacteroides massiliensis	100
		134	18	0	15	3	0	Bacteroidetes	Bacteroides nordii	99.4
		195	18	0	17	1	0	Firmicutes	Faecalibacterium prausnitzii	98.8
		244	18	0	17	1	0	Firmicutes	Faecalibacterium prausnitzii	91.6
		59	18	0	11	7	0	Firmicutes	Faecalibacterium prausnitzii	98.8
		246	17	0	14	3	0	Firmicutes	Blautia wexlerae	93.9
		309	17	0	16	1	0	Firmicutes	Ruminococcus bromii	94.2
		69	17	0	10	7	0	Firmicutes	Blautia glucerasea	100
		250	16	0	6	10	0	Firmicutes	Staphylococcus warneri	100
		281	16	0	7	8	0	Firmicutes	Clostridium hylemonae	99.2
		323	16	0	15	1	0	Firmicutes	Clostridium populeti	97.5
		229	15	0	15	0	0	Firmicutes	Ruminococcus torques	97.6
		317	15	2	9	4	0	Firmicutes	Ruminococcus gnavus	99.8
		34	14	1	11	2	0	Firmicutes	Ruminococcus lactaris	96.4

		0	13	2	6	5	0	Firmicutes	Ruminococcus gnavus	100
		117	11	0	4	6	0	Bacteroidetes	Parabacteroides merdae	100
		237	11	0	3	8	0	Bacteroidetes	Bacteroides coprocola	99.8
		325	11	2	7	2	0	Firmicutes	Ruminococcus gnavus	100
		31	8	0	4	4	0	Proteobacteria	Escherichia/Shigella flexneri	99.8
		125	7	0	3	4	0	Bacteroidetes	Barnesiella intestinihominis	99
		65	7	0	6	1	0	Firmicutes	Oscillibacter valericigenes	94.5
		80	7	0	4	3	0	Firmicutes	Eubacterium ramulus	99.8
		296	6	0	4	2	0	Firmicutes	Roseburia intestinalis	99.7
		49	6	0	4	2	0	Firmicutes	Roseburia inulinivorans	99.6
		54	5	0	5	0	0	Firmicutes	Clostridium orbiscindens	95.9
		132	4	0	3	1	0	Firmicutes	Ruminococcus callidus	99.5
		99	4	0	3	1	0	Firmicutes	Clostridium asparagiforme	95.6
		158	2	0	1	1	0	Firmicutes	Megamonas funiformis	99.7
		43	2	0	1	1	0	Firmicutes	Desulfotomaculum guttoideum	97.3
		61	2	0	2	0	0	Firmicutes	Faecalibacterium prausnitzii	98.8
		166	1	0	1	0	0	Firmicutes	Blautia luti	98.6
		187	1	0	0	1	0	Firmicutes	Ruminococcus lactaris	96.4
		225	1	0	0	1	0	NA	Insolitispirillum peregrinum	82.5
		267	1	0	0	1	0	Bacteroidetes	Bacteroides thetaiotaomicron	90.1
		33	1	0	1	0	0	Bacteroidetes	Alistipes onderdonkii	97.1
2	1.26e ⁻⁰⁷	24	26	1	16	8	0	Firmicutes	Dorea formicigenerans	100
		53	26	3	15	8	0	Proteobacteria	Sutterella wadsworthensis	100
		7	25	2	13	9	0	Bacteroidetes	Bacteroides uniformis	100
		98	23	3	12	6	0	Firmicutes	Anaerostipes butyraticus	97.6
		204	22	4	14	4	0	Bacteroidetes	Bacteroides dorei	98.4
		12	21	3	12	6	0	Firmicutes	Dorea longicatena	100
		36	21	0	6	15	0	Actinobacteria	Bifidobacterium longum	98.8
		177	20	2	14	4	0	Bacteroidetes	Bacteroides dorei	98.6
		241	20	0	16	4	0	Firmicutes	Desulfotomaculum guttoideum	97.7
		162	19	0	8	10	0	Bacteroidetes	Bacteroides chinchillae	99.5
		29	19	3	14	2	0	Bacteroidetes	Parabacteroides distasonis	98.3
		101	17	3	11	3	0	Bacteroidetes	Bacteroides dorei	99

		115	16	0	6	9	0	Bacteroidetes	Bacteroides chinchillae	100
		153	16	1	12	3	0	Firmicutes	Clostridium bolteae	95.3
		14	13	0	8	5	0	Bacteroidetes	Bacteroides dorei	98.6
		16	12	1	11	0	0	Firmicutes	Ruminococcus torques	100
		41	12	2	4	6	0	Firmicutes	Desulfotomaculum guttoideum	95.9
		126	11	0	10	1	0	Firmicutes	Clostridium glycyrrhizinilyticum	97.5
		288	11	0	10	1	0	Bacteroidetes	Bacteroides dorei	98.8
		155	10	0	9	0	0	Firmicutes	Desulfotomaculum guttoideum	96.8
		176	9	0	5	4	0	Proteobacteria	Acinetobacter johnsonii	99
		243	8	0	3	5	0	Proteobacteria	Pseudomonas flavescens	100
		304	8	0	5	3	0	Firmicutes	Dorea longicatena	100
		10	7	0	4	3	0	Firmicutes	Clostridium nexile	99.8
		270	7	0	7	0	0	Bacteroidetes	Bacteroides dorei	98.8
		259	5	0	5	0	0	Firmicutes	Clostridium hylemonae	97.1
		89	5	0	3	2	0	Actinobacteria	Bifidobacterium adolescentis	100
		93	4	0	3	1	0	Bacteroidetes	Bacteroides acidifaciens	95.9
		120	3	0	2	1	0	Proteobacteria	Acinetobacter lwoffii	100
		188	3	0	0	3	0	Firmicutes	Anaerostipes caccae	99.8
		245	3	0	2	1	0	Firmicutes	Anoxybacillus rupiensis	100
		258	3	0	3	0	0	Firmicutes	Desulfotomaculum guttoideum	96.9
		305	3	0	1	2	0	Bacteroidetes	Sphingobacterium multivorum	99.7
		86	3	0	3	0	0	Firmicutes	Clostridium lactatifermentans	93.5
		247	1	0	0	1	0	Firmicutes	Lactobacillus delbrueckii	100
3	5.86e ⁻⁰⁵	2	24	1	19	4	0	Firmicutes	Eubacterium rectale	99.7
		199	18	0	12	6	0	Firmicutes	Ruminococcus schinkii	96.6
		151	17	1	9	7	0	Firmicutes	Ruminococcus obeum	100
		25	16	1	5	10	0	Fusobacteria	Fusobacterium mortiferum	98.6
		138	14	1	9	4	0	Bacteroidetes	Alistipes shahii	99.8
		19	13	0	7	6	0	Firmicutes	Ruminococcus albus	95.7
		131	12	0	7	5	0	Firmicutes	Roseburia faecis	99.7
		30	12	0	11	1	0	Firmicutes	Clostridium orbiscindens	100
		9	12	1	8	2	1	Firmicutes	Blautia wexlerae	99.8
		336	11	0	10	1	0	Firmicutes	Clostridium orbiscindens	99.8

		38	11	0	10	1	0	Firmicutes	Clostridium bolteae	99.5
		4	9	1	5	3	0	Bacteroidetes	Bacteroides fragilis	98.6
		116	8	0	7	1	0	Firmicutes	Clostridium populeti	94.5
		87	8	0	3	4	1	Firmicutes	Clostridium lituseburense	98
		106	7	0	6	1	0	Firmicutes	Eubacterium desmolans	98.2
		142	6	0	3	3	0	Firmicutes	Granulicatella elegans	100
		159	6	0	1	5	0	Firmicutes	Alicyclobacillus pohliae	99
		3	6	0	3	3	0	Bacteroidetes	Bacteroides stercoris	99
		71	6	0	6	0	0	Actinobacteria	Eggerthella lenta	99.6
		77	6	0	5	1	0	Firmicutes	Dialister invisus	100
		274	5	0	2	3	0	Bacteroidetes	Bacteroides fragilis	99.2
		255	4	0	2	2	0	Firmicutes	Clostridium disporicum	99.7
		51	4	0	3	1	0	Firmicutes	Cellulosilyticum ruminicola	92.6
		73	3	0	3	0	0	Bacteroidetes	Bacteroides eggerthii	99.5
		79	3	0	3	0	0	Actinobacteria	Collinsella intestinalis	99.5
		107	2	0	0	2	0	Proteobacteria	Paracoccus yeei	99.5
		191	2	0	1	1	0	Firmicutes	Butyricicoccus pullicaecorum	99.6
		203	2	0	1	1	0	Firmicutes	Lactococcus lactis	100
		303	1	0	0	1	0	Firmicutes	Acetanaerobacterium elongatum	92.9
		84	1	0	0	1	0	Bacteroidetes	Odoribacter splanchnicus	99.3
4	2.47e ⁻⁰⁶	20	17	1	15	1	0	Firmicutes	Gemmiger formicilis	99.8
		48	16	1	14	1	0	Bacteroidetes	Prevotella amnii	84.5
		104	14	0	11	3	0	Firmicutes	Clostridium orbiscindens	90.8
		105	14	0	13	1	0	Firmicutes	Dialister succinatiphilus	98.7
		22	14	0	12	1	0	Firmicutes	Clostridium aldenense	96.4
		127	11	0	7	4	0	Bacteroidetes	Bacteroides cellulosilyticus	99.5
		161	11	2	9	0	0	Firmicutes	Gemmiger formicilis	100
		289	11	0	5	6	0	Firmicutes	Clostridium saccharolyticum	96.4
		39	11	0	8	3	0	Bacteroidetes	Prevotella copri	99.3
		240	9	0	9	0	0	Firmicutes	Coprococcus eutactus	95.4
		81	9	0	8	1	0	Actinobacteria	Slackia piriformis	90.4
		92	9	0	9	0	0	Firmicutes	Coprococcus catus	100
		157	8	0	8	0	0	Firmicutes	Eubacterium eligens	99.8

		301	8	0	2	6	0	Firmicutes	Ruminococcus schinkii	96.8
		47	8	0	8	0	0	Firmicutes	Clostridium lactatifermentans	95.2
		156	7	0	6	1	0	Proteobacteria	Sutterella stercoricanis	99.6
		291	5	0	5	0	0	NA	Rhodospirillum rubrum	87.7
		90	5	0	5	0	0	Proteobacteria	Sutterella stercoricanis	99
		249	4	0	3	0	0	Firmicutes	Allisonella histaminiformans	98.1
		37	4	0	3	1	0	Firmicutes	Eubacterium hallii	99.7
		44	4	0	2	2	0	Firmicutes	Gemmiger formicilis	97.6
		220	3	0	3	0	0	Proteobacteria	Klebsiella pneumoniae	100
		242	3	0	3	0	0	Firmicutes	Lactobacillus taiwanensis	100
		110	1	0	1	0	0	Firmicutes	Mogibacterium pumilum	92.3
		50	1	0	1	0	0	Firmicutes	Lactobacillus rogosae	99.8
		55	1	0	1	0	0	Proteobacteria	Methylobacterium adhaesivum	98.9
		63	1	0	1	0	0	Actinobacteria	Collinsella aerofaciens	99
		70	1	0	1	0	0	Firmicutes	Sporobacter termitidis	94.1
		83	1	0	1	0	0	NA	Desulfotomaculum carboxydivorans	86.7
		96	1	0	0	1	0	Firmicutes	Gemmiger formicilis	93.8
5	1.91e ⁻⁴	60	18	0	12	6	0	Firmicutes	Clostridium ramosum	99.8
		76	18	0	15	3	0	Proteobacteria	Mesorhizobium australicum	98.8
		91	12	0	11	1	0	Deinococcus-Thermus	Thermus scotoductus	99.5
		58	10	1	9	0	0	Proteobacteria	Undibacterium oligocarboniphilum	100
		139	9	0	8	1	0	Bacteroidetes	Prevotella enoeca	83.3
		224	8	0	8	0	0	Proteobacteria	Bradyrhizobium yuanmingense	99.5
		23	8	0	6	2	0	Firmicutes	Streptococcus salivarius	100
		108	7	0	7	0	0	Proteobacteria	Pelomonas saccharophila	99
		11	7	0	6	1	0	Firmicutes	Thermanaeromonas toyohensis	85.9
		227	7	0	6	1	0	Bacteroidetes	Sediminibacterium salmoneum	96
		28	7	1	5	1	0	Firmicutes	Streptococcus mitis	100
		46	7	0	5	2	0	Firmicutes	Blautia hansenii	99.8
		68	7	1	5	1	0	Proteobacteria	Sphingomonas melonis	99.8
		64	6	1	4	1	0	Firmicutes	Gemella haemolysans	99.8
		124	5	0	4	0	0	Firmicutes	Granulicatella adiacens	100
		264	5	0	5	0	0	Actinobacteria	Actinomyces odontolyticus	100

		123	4	0	0	3	0	Firmicutes	Clostridium hylemonae	97.5
		143	4	0	3	1	0	Bacteroidetes	Lishizhenia tianjinensis	86.9
		154	4	0	3	1	0	Proteobacteria	Ralstonia insidiosa	99.7
		205	4	0	3	1	0	Proteobacteria	Comamonas aquatica	98.7
		178	3	0	2	1	0	Actinobacteria	Atopobium rimae	100
		163	2	0	2	0	0	Firmicutes	Streptococcus sinensis	98.8
		57	2	0	2	0	0	Firmicutes	Clostridium perfringens	100
		74	2	0	2	0	0	Firmicutes	Clostridium leptum	95.2
		130	1	0	1	0	0	Proteobacteria	Brevundimonas vesicularis	100
		226	1	0	1	0	0	Bacteroidetes	Sediminibacterium salmoneum	98.1
ND	ND	67	26	3	15	8	0	Bacteroidetes	Bacteroides caccae	99.8
ND	ND	230	23	0	14	8	0	Bacteroidetes	Bacteroides xylanisolvens	99.8
ND	ND	56	20	1	14	5	0	Bacteroidetes	Bacteroides nordii	99.7
ND	ND	21	17	0	12	5	0	Bacteroidetes	Bacteroides caccae	100
ND	ND	32	15	2	12	1	0	Bacteroidetes	Bacteroides thetaiotaomicron	100
ND	ND	327	15	1	11	3	0	Bacteroidetes	Bacteroides nordii	99.5
ND	ND	13	14	1	11	2	0	Firmicutes	Clostridium bolteae	100
ND	ND	223	14	0	7	7	0	Actinobacteria	Collinsella aerofaciens	100
ND	ND	8	13	0	10	3	0	Firmicutes	Blautia glucerasea	98.8
ND	ND	208	12	0	10	2	0	Bacteroidetes	Butyricimonas virosa	98.3
ND	ND	94	10	0	10	0	0	Firmicutes	Clostridium scindens	100
ND	ND	82	8	0	7	1	0	Firmicutes	Blautia hansenii	100
ND	ND	97	8	0	7	1	0	Proteobacteria	Bilophila wadsworthia	99.7
ND	ND	145	5	0	5	0	0	Firmicutes	Clostridium aldenense	98
ND	ND	169	3	0	3	0	0	Firmicutes	Ruminococcus obeum	99.8
ND	ND	277	3	0	1	2	0	NA	Bacteroides plebeius	90.7
ND	ND	251	2	0	1	1	0	Actinobacteria	Bifidobacterium breve	100
ND	ND	307	1	0	1	0	0	Firmicutes	Veillonella parvula	99.8

Supp. Table 4: OTU correlation network composition deduced from patients in remission 6 months after ICR

Cluster: Identity of the cluster; OTUs: identity of the Operational Taxonomic Unit specific of each cluster; Total: total number of significant correlations between the identified OTU and other OTUs from the cluster; pos75: number of significant positive correlations ($R \ge 0.75$) between the identified OTU and other OTUs from the cluster; pos50: number of significant positive correlations ($R \ge 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant positive correlations ($R \ge 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant positive correlations ($R \ge 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \le 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \le 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \le 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \le 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \le 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \le 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \le 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \le 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \le 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \le 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \le 0.50$) between the identified OTU a

the cluster; Similarity: similarity percentage between the OTU and the first relative species. No significant negative correlation with R≤-0.75 was observed.

Cluster	Cluster p-value	OTUs	Total	pos75	pos50	neg50	neg75	Phylum	Species	Similarity (%)
1	0.0107	28	5	2	3	0	0	Firmicutes	Streptococcus mitis	100
		163	4	0	3	1	0	Firmicutes	Streptococcus sinensis	98.8
		64	4	1	3	0	0	Firmicutes	Gemella haemolysans	99.8
		124	2	1	1	0	0	Firmicutes	Granulicatella adiacens	100
		176	2	0	2	0	0	Proteobacteria	Acinetobacter johnsonii	99
		227	1	0	1	0	0	Bacteroidetes	Sediminibacterium salmoneum	96
		264	1	0	1	0	0	Actinobacteria	Actinomyces odontolyticus	100
2	5.27e ⁻⁰⁴	205	6	2	4	0	0	Proteobacteria	Comamonas aquatica	98.7
		5	6	0	5	1	0	Firmicutes	Ruminococcus bromii	99.8
		108	5	3	2	0	0	Proteobacteria	Pelomonas saccharophila	99
		11	5	2	2	0	1	Firmicutes	Thermanaeromonas toyohensis	85.9
		159	5	0	5	0	0	Firmicutes	Alicyclobacillus pohliae	99
		154	4	2	2	0	0	Proteobacteria	Ralstonia insidiosa	99.7
		76	4	0	4	0	0	Proteobacteria	Mesorhizobium australicum	98.8
		102	3	0	3	0	0	Firmicutes	Oscillibacter valericigenes	91.9
		55	3	0	3	0	0	Proteobacteria	Methylobacterium adhaesivum	98.9
		256	2	0	2	0	0	Proteobacteria	Rhodospirillum rubrum	85.8
		91	2	1	1	0	0	Deinococcus-Thermus	Thermus scotoductus	99.5
		150	1	0	1	0	0	Proteobacteria	Methylobacterium adhaesivum	98.8
		287	1	0	1	0	0	Firmicutes	Dendrosporobacter quercicolus	81.2
		307	1	0	1	0	0	Firmicutes	Veillonella parvula	99.8
3	1.25e ⁻⁰⁶	8	13	7	6	0	0	Firmicutes	Blautia glucerasea	98.8
		18	12	4	7	0	1	Firmicutes	Clostridium hylemonae	98.5
		38	12	1	10	1	0	Firmicutes	Clostridium bolteae	99.5
		103	10	1	9	0	0	Proteobacteria	Aeromonas ichthiosmia	100
		193	10	4	6	0	0	Firmicutes	Clostridium bartlettii	99.5
		222	9	0	8	1	0	Firmicutes	Enterococcus faecalis	100
		281	9	3	6	0	0	Firmicutes	Clostridium hylemonae	99.2
		325	9	2	7	0	0	Firmicutes	Ruminococcus gnavus	100
		82	9	2	7	0	0	Firmicutes	Blautia hansenii	100
		191	8	0	7	1	0	Firmicutes	Butyricicoccus pullicaecorum	99.6
		145	7	1	6	0	0	Firmicutes	Clostridium aldenense	98

		188	6	0	6	0	0	Firmicutes	Anaerostipes caccae	99.8
		202	6	0	6	0	0	Bacteroidetes	Parabacteroides distasonis	98.6
		301	6	3	3	0	0	Firmicutes	Ruminococcus schinkii	96.8
		13	5	0	5	0	0	Firmicutes	Clostridium bolteae	100
		23	5	0	4	1	0	Firmicutes	Streptococcus salivarius	100
		6	5	1	0	3	1	Firmicutes	Faecalibacterium prausnitzii	98.8
		85	5	2	3	0	0	Firmicutes	Clostridium bartlettii	99.5
		184	4	0	3	1	0	Proteobacteria	Proteus vulgaris	99.8
		317	4	2	2	0	0	Firmicutes	Ruminococcus gnavus	99.8
		75	4	0	4	0	0	Firmicutes	Clostridium hylemonae	99.2
		0	3	1	2	0	0	Firmicutes	Ruminococcus gnavus	100
		220	3	0	1	2	0	Proteobacteria	Klebsiella pneumoniae	100
		29	3	2	1	0	0	Bacteroidetes	Parabacteroides distasonis	98.3
		316	3	0	3	0	0	Firmicutes	Clostridium bartlettii	99.7
		40	3	0	3	0	0	Firmicutes	Enterococcus durans	100
		2	1	1	0	0	0	Firmicutes	Eubacterium rectale	99.7
		232	1	0	1	0	0	Firmicutes	Phascolarctobacterium faecium	100
		335	1	0	1	0	0	Firmicutes	Enterococcus durans	100
		60	1	0	1	0	0	Firmicutes	Clostridium ramosum	99.8
ļ	7.54e ⁻³	58	7	1	4	2	0	Proteobacteria	Undibacterium oligocarboniphilum	100
		139	4	0	3	0	0	Bacteroidetes	Prevotella enoeca	83.3
		258	4	0	2	2	0	Firmicutes	Desulfotomaculum guttoideum	96.9
		68	4	1	3	0	0	Proteobacteria	Sphingomonas melonis	99.8
		250	2	0	2	0	0	Firmicutes	Staphylococcus warneri	100
		257	2	0	2	0	0	Firmicutes	Faecalibacterium prausnitzii	95
		143	1	0	1	0	0	Bacteroidetes	Lishizhenia tianjinensis	86.9
		170	1	0	1	0	0	Firmicutes	Clostridium hathewayi	99.5
		224	1	0	1	0	0	Proteobacteria	Bradyrhizobium yuanmingense	99.5
		4	1	0	1	0	0	Bacteroidetes	Bacteroides fragilis	98.6
		41	1	0	0	1	0	Firmicutes	Desulfotomaculum guttoideum	95.9
		62	1	0	0	0	0	Firmicutes	Clostridium thermocellum	87.9
5	3.91e ⁻⁰⁷	21	12	2	9	0	1	Bacteroidetes	Bacteroides caccae	100
		37	10	2	7	1	0	Firmicutes	Eubacterium hallii	99.7

		117	8	0	8	0	0	Bacteroidetes	Parabacteroides merdae	100
		52	8	0	8	0	0	Bacteroidetes	Prevotella oralis	91.1
		22	7	0	7	0	0	Firmicutes	Clostridium aldenense	96.4
		3	7	0	7	0	0	Bacteroidetes	Bacteroides stercoris	99
		84	7	1	6	0	0	Bacteroidetes	Odoribacter splanchnicus	99.3
		14	6	1	4	0	1	Bacteroidetes	Bacteroides dorei	98.6
		33	6	0	6	0	0	Bacteroidetes	Alistipes onderdonkii	97.1
		7	6	3	3	0	0	Bacteroidetes	Bacteroides uniformis	100
		155	5	0	5	0	0	Firmicutes	Desulfotomaculum guttoideum	96.8
		24	5	0	4	0	0	Firmicutes	Dorea formicigenerans	100
		47	5	0	5	0	0	Firmicutes	Clostridium lactatifermentans	95.2
		20	4	2	2	0	0	Firmicutes	Gemmiger formicilis	99.8
		63	4	0	4	0	0	Actinobacteria	Collinsella aerofaciens	99
		9	4	2	0	2	0	Firmicutes	Blautia wexlerae	99.8
		98	4	0	3	1	0	Firmicutes	Anaerostipes butyraticus	97.6
		166	3	0	3	0	0	Firmicutes	Blautia luti	98.6
		204	3	1	2	0	0	Bacteroidetes	Bacteroides dorei	98.4
		53	3	0	2	1	0	Proteobacteria	Sutterella wadsworthensis	100
		74	3	0	3	0	0	Firmicutes	Clostridium leptum	95.2
		129	2	0	2	0	0	Firmicutes	Ruminococcus schinkii	96.7
		177	2	0	2	0	0	Bacteroidetes	Bacteroides dorei	98.6
		31	2	0	0	1	1	Proteobacteria	Escherichia/Shigella flexneri	99.8
		49	2	0	2	0	0	Firmicutes	Roseburia inulinivorans	99.6
		54	2	0	2	0	0	Firmicutes	Clostridium orbiscindens	95.9
		78	2	0	2	0	0	Firmicutes	Ruminococcus lactaris	96.1
		138	1	0	1	0	0	Bacteroidetes	Alistipes shahii	99.8
		19	1	0	0	1	0	Firmicutes	Ruminococcus albus	95.7
		229	1	0	0	0	0	Firmicutes	Ruminococcus torques	97.6
		327	1	0	1	0	0	Bacteroidetes	Bacteroides nordii	99.5
		80	1	0	1	0	0	Firmicutes	Eubacterium ramulus	99.8
		99	1	0	1	0	0	Firmicutes	Clostridium asparagiforme	95.6
ND	ND	169	8	0	7	1	0	Firmicutes	Ruminococcus obeum	99.8
ND	ND	10	6	1	5	0	0	Firmicutes	Clostridium nexile	99.8

ND	ND	151	4	0	3	1	0	Firmicutes	Ruminococcus obeum	100
ND	ND	123	3	0	3	0	0	Firmicutes	Clostridium hylemonae	97.5
ND	ND	46	2	0	2	0	0	Firmicutes	Blautia hansenii	99.8
ND	ND	59	2	0	0	2	0	Firmicutes	Faecalibacterium prausnitzii	98.8

Supp. Table 5: OTU correlation network composition deduced from patients with endoscopic recurrence 6 months after ICR

Cluster: Identity of the cluster; OTUs: identity of the Operational Taxonomic Unit specific of each cluster; Total: total number of significant correlations between the identified OTU and other OTUs from the cluster; pos75: number of significant positive correlations ($R \ge 0.75$) between the identified OTU and other OTUs from the cluster; pos50: number of significant positive correlations ($R \ge 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant positive correlations ($R \ge 0.50$) between the identified OTU and other OTUs from the cluster; pos50: number of significant negative correlations ($R \ge 0.50$) between the identified OTU and other OTUs from the cluster; neg50: number of significant negative correlations ($R \le -0.5$) between the identified OTU and other OTUs from the cluster; similarity percentage between the OTU and the first relative species. No significant negative correlation with $R \le -0.75$ was observed.

OTUs	Total	pos75	pos50	neg50	neg75	Phylum	Species	Similarity (%)
0	1	1	0	0	0	Firmicutes	Ruminococcus gnavus	100
101	2	2	0	0	0	Bacteroidetes	Bacteroides dorei	99
130	1	1	0	0	0	Proteobacteria	Brevundimonas vesicularis	100
177	2	1	1	0	0	Bacteroidetes	Bacteroides dorei	98.6
180	1	0	1	0	0	Firmicutes	Clostridium spiroforme	92.7
204	2	2	0	0	0	Bacteroidetes	Bacteroides dorei	98.4
226	2	1	0	1	0	Bacteroidetes	Sediminibacterium salmoneum	98.1
24	1	1	0	0	0	Firmicutes	Dorea formicigenerans	100
270	3	3	0	0	0	Bacteroidetes	Bacteroides dorei	98.8
28	1	0	0	1	0	Firmicutes	Streptococcus mitis	100
289	4	0	2	2	0	Firmicutes	Clostridium saccharolyticum	96.4
310	2	2	0	0	0	NA	Escherichia/Shigella flexneri	92.6
49	3	0	3	0	0	Firmicutes	Roseburia inulinivorans	99.6
53	1	0	1	0	0	Proteobacteria	Sutterella wadsworthensis	100
84	1	0	1	0	0	Bacteroidetes	Odoribacter splanchnicus	99.3

Supp. Table 6: OTU correlation network composition deduced from patients at time of surgery

Cluster: Identity of the cluster; OTUs: identity of the Operational Taxonomic Unit specific of each cluster; Total: total number of significant correlations between the identified OTU and other OTUs from the cluster; pos75: number of significant positive correlations (R \geq 0.75) between the identified OTU and other OTUs from the cluster; pos50: number of significant positive correlations (R \geq 0.50) between the identified OTU and other OTUs from the cluster; pos50: number of significant positive correlations (R \geq 0.50) between the identified OTU and other OTUs from the cluster; neg50: number of significant negative correlations (R \leq -0.5) between the identified OTU and other OTUs from the cluster; similarity percentage between the OTU and the first relative species. No significant negative correlation with R \leq -0.75 was observed.