Original Article

Anti-inflammatory Gut Microbial Pathways Are Decreased During Crohn’s Disease Exacerbations

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Abstract

Background and Aims: Crohn’s disease [CD] is a chronic inflammatory disorder of the gastrointestinal tract characterised by alternating periods of exacerbation and remission. We hypothesised that changes in the gut microbiome are associated with CD exacerbations, and therefore aimed to correlate multiple gut microbiome features to CD disease activity.

Methods: Faecal microbiome data generated using whole-genome metagenomic shotgun sequencing of 196 CD patients were obtained from the 1000IBD cohort [one sample per patient]. Patient disease activity status at time of sampling was determined by re-assessing clinical records 3 years after faecal sample production. Faecal samples were designated as taken ‘in an exacerbation’ or ‘in remission’. Samples taken ‘in remission’ were further categorised as ‘before the next exacerbation’ or ‘after the last exacerbation’, based on the exacerbation closest in time to the faecal production date. CD activity was correlated with gut microbial composition and predicted functional pathways via logistic regressions using MaAsLin software.

Results: In total, 105 bacterial pathways were decreased during CD exacerbation (false-discovery rate [FDR] <0.1) in comparison with the gut microbiome of patients both before and after an exacerbation. Most of these decreased pathways exert anti-inflammatory properties facilitating the biosynthesis and fermentation of various amino acids [tryptophan, methionine, and arginine], vitamins [riboflavin and thiamine], and short-chain fatty acids [SCFAs].

Conclusions: CD exacerbations are associated with a decrease in microbial genes involved in the biosynthesis of the anti-inflammatory mediators riboflavin, thiamine, and folate, and SCFAs, suggesting that increasing the intestinal abundances of these mediators might provide new treatment opportunities. These results were generated using bioinformatic analyses of cross-sectional data and need to be replicated using time-series and wet lab experiments.

Key Words: Crohn’s disease activity; faecal microbiome; whole-genome metagenomic shotgun sequencing
1. Background

Inflammatory bowel disease (IBD) is a chronic inflammatory disorder of the gastrointestinal [GI] tract that is characterised by alternating periods of exacerbation [active disease] and remission [quiescent disease].\textsuperscript{1,2} IBD comprises ulcerative colitis [UC] and Crohn’s disease [CD]; in UC, inflammation is confined to the mucosal layer of the colon, whereas in CD, inflammation can pervade every layer at any site of the GI tract.\textsuperscript{1,2} Although the factors that cause the development and onset of IBD are increasingly understood,\textsuperscript{1,4} the factors that influence the dynamics of disease activity have yet to be elucidated. A growing body of evidence, however, suggests that changes in the gut microbiome could be associated with an exacerbation of IBD.\textsuperscript{3,4} The identification of gut microbiome features associated with disease activity could thus provide novel targets for early phase intervention aimed at preventing the development of full-blown active disease or at maintaining IBD patients in a quiescent phase.

Over the past few years, a number of studies using sequencing of the 16S rRNA gene have compared the bacterial compositions in patients with active and quiescent disease, but these studies have produced conflicting results.\textsuperscript{5–7} A major limitation of 16S rRNA sequencing, however, is that it only describes which bacteria are present, not what they do. In contrast, whole-genome metagenomic sequencing can be used to predict the activity of microbial metabolic pathways, and this is information that might be more informative.\textsuperscript{8,9}

In previous analyses by our group,\textsuperscript{5,8} only a small number of associations between individual measures of disease activity—i.e. Harvey Bradshaw Index [HBI], faecal calprotectin [FC] and C-reactive protein [CRP] levels—and the gut microbiome were detected.

In this study, we aimed to better understand the role of the gut microbiome in disease activity in CD by analysing many different aspects of the gut microbiome, including microbial diversity, composition, the presence of genes involved in bacterial virulence, bacterial growth rates and microbial metabolic pathways in 196 CD patients, and comparing the results between patients with active and quiescent disease. Only one microbiome profile was available per patient, but because these patients were all clinically followed for several years after faecal sample production, we were able to calculate the number of days between stool sampling and nearest onset of disease activity. This allowed us to create a single virtual timeline of CD activity and to group patients into ‘before’, ‘during’, and ‘after’ exacerbation categories, which in turn allowed us to discover microbial features associated with CD activity.

2. Materials and Methods

2.1. Study participants: clinical characteristics

2.1.1. 1000IBD cohort and informed consent

The 1000IBD cohort consists of more than 1000 IBD patients. The cohort has been described previously\textsuperscript{19} and originates from the specialised IBD Center of the Department of Gastroenterology and Hepatology of the University Medical Center Groningen [UMCG] in Groningen, The Netherlands. The 196 CD patients included in this study, and their corresponding metagenomic sequenced stool samples, all met study inclusion criteria [Supplementary material 1, available as Supplementary data at ECCO-JCC online]. The study was approved by the Institutional Review Board of the UMCG [IRB number 2008.338].\textsuperscript{4} More information on the 1000IBD cohort can be found at [https://1000IBD.org].

2.1.2. Clinical characteristics, medication use, and dietary patterns

Extensive clinical phenotypes were documented for each CD patient in the study, including disease location, medication use at time of sampling, presence of perianal disease, previous surgery, stricturing phenotype, disease duration [indicated by the number of years since the Crohn’s disease diagnosis], CRP levels, and calprotectin levels. All these factors were measured around the time of faecal sampling. In addition, each patient filled in a Food Frequency Questionnaire [FFQ]\textsuperscript{19} at the time of faecal sampling, which recorded their food intake in the previous month. The FFQ answers were then transformed to grams/day in 25 food groups.

2.1.3. Definition of disease activity: exacerbation or remission

Electronic health records containing prospectively collected data on study participants were reviewed to determine disease activity status. Defining whether a patient with CD has an exacerbation can be difficult because individual clinical disease activity scores and biomarkers often do not correctly capture CD activity.\textsuperscript{20–22} Subjective disease activity scores rely heavily upon an increase in defaecation frequency and subjective abdominal complaints to determine disease activity. Biomarkers like faecal calprotectin [FC] show a highest sensitivity of 0.80 and specificity of 0.82 in inflammatory bowel disease [cut-off 250 μg/g],\textsuperscript{23} but FC does not detect ideal inflammation very well and is therefore less reliable in Crohn’s disease than in ulcerative colitis.\textsuperscript{23} Other biomarkers like CRP can also increase due to other causes, such as Clostridium difficile infections.\textsuperscript{24–26} To determine whether there is an exacerbation, a gastroenterologist will often take all these measures into account, as well as the outcome of a colonoscopy [if performed], the patient’s medical history, and the results of tests to exclude infectious enteritis. Therefore, in this study, a CD exacerbation was defined based on a combination of the following criteria:

i) An increase in the occurrence of IBD-associated gastrointestinal complaints that could not be attributed to other concurrent GI diseases, i.e., severe abdominal pain and cramping, increase in stool frequency, decrease in stool consistency, bloody diarrhoea, and/or malaise necessitating consecutive CD treatment adjustment or intensification.

ii) Interpretation by the treating physician based on a combination of the following criteria:

a. unplanned visits to the outpatient clinic due to typical patient complaints;

b. changes in IBD-associated biomarkers [calprotectin >200 μg/g faeces, CRP >5 mg/L] in a patient with typical complaints that could not be explained otherwise;

c. necessity of increasing dose and/or type of anti-IBD medication;

d. active inflammation seen during endoscopy;

e. active inflammation confirmed in a pathology report of gut biopsies.

2.1.4. Determination of disease activity status at the time of sampling

The dates of onset and recovery from recorded CD exacerbations were determined relative to the faecal sample production date. Faecal samples were thereby classified as taken ‘in an exacerbation’ or ‘in remission’. Samples ‘in remission’ were further categorised as ‘before the next exacerbation’ or ‘after the last exacerbation’. In addition, time to nearest disease exacerbation was recorded for all
samples, defined as ‘days until next exacerbation’ or ‘days since last exacerbation’. When there were multiple recorded exacerbations, the closest exacerbation relative to the faecal production date was used to determine the disease activity.

2.2. Stool sampling and analysis of gut microbiome
Gut microbiome taxonomic compositions and functional profiles, bacterial growth rates, and abundances of virulence factor genes were determined as previously described.18

2.2.1. Sample collection and microbial DNA isolation
Patients produced and froze [-20°C] a stool sample at home. Frozen samples were then transported on dry ice to the UMCG and stored at -80°C. Microbial DNA was extracted from faecal samples using the Qiagen Allprep DNA/RNA Mini Kit [cat # 380204] with the addition of mechanical lysis.

2.2.2. Sequencing of microbial DNA
Metagenomic shotgun sequencing of microbial DNA was performed at the Broad Institute of Harvard and MIT in Cambridge, MA, USA, using the HiSeq platform. For genomic library preparation, the Nextera XT Library preparation Kit was used.

2.2.3. Quality control and determination of microbiome parameters
Trimmomatic [v.0.32]32 was used to remove adapters and trim the ends of metagenomic reads, and samples with a read depth below 10 million reads were removed. Cleaned metagenomic reads were processed using a previously published pipeline. In brief: a. taxonomic compositions were determined by aligning reads to the reference database RefSeq NCBI database33 [accession date: June 3, 2016] using the software tools Kraken34 and Bracken35; b. functional pathways were determined using HUMAnN2 [v.0.4.0] [http://huttenhower.sph.harvard.edu/humann2] and families were grouped into pathways using multi-organism database MetaCyc36; c. abundances of genes encoding virulence factors were determined by aligning reads to the Virulence Factor Database5 using DIAMOND [version 0.8.2.10]; and d. bacterial growth rates were estimated using a previously described peak-to-trough ratio algorithm. Microbiome features present in at least 25% of samples were tested.

2.3. Statistical analysis
2.3.1. Summary statistics
We tested if the clinical characteristics, dietary patterns, Shannon Index and gene richness, and BrayCurtis distances [i.e., the inter-individual diversity or β-diversities] [Supplementary material 2, available as Supplementary data at ECCO-JCC online] differed significantly between patient groups ['in remission' vs 'in an exacerbation' and 'before the next exacerbation' vs 'after the last exacerbation']. The chi square test was used for binary phenotypes and the WilcoxonMann-Whitney U test was used for continuous phenotypes. The BenjaminiHochberg method was used to calculate the false-discovery rate [FDR] for multiple testing. An FDR <0.1 was considered statistically significant. The proportion of explained variance on β-diversity was calculated for each clinical characteristic using the ADONIS function in R [Table S2, available as Supplementary data at ECCO-JCC online].

2.3.2. Covariates
Models were constructed considering the covariates: age, sex, read depth, body mass index [BMI], disease location, use of PPI, and use of antibiotics 3 months before sampling, all of which are known to influence the gut microbiome. The dietary covariate 'pre-prepared meals' differed significantly between groups [Table 1] and was added to the models. The use of steroids differed significantly between groups [p = 0.000, chi square test], but was also correlated with disease activity [r = 0.47, p = 0.000]. Since the correlation between independent variables—collinearity—is hard to take into account into the model, we did not take steroid use into account in the linear models. However, for transparency, we have added analyses with a correction for steroid use.

2.3.3. Categorical analyses: before, during and after an exacerbation
Using the statistical software MaAsLin,9 we compared gut microbiota features [relative abundances of species and functional pathways, abundances of virulence factors, and growth rate ratios] between: i. patients in an exacerbation versus patients in remission; ii. patients before versus in an exacerbation; and iii. patients in versus after an exacerbation. The MaAsLin boosting step was turned off to ensure all independent variables were taken into account. Taxonomy and pathway relative abundances were arcsine square-root transformed. Zero inflation was considered in all tests except for growth rates. An FDR of 0.1 was used as a cut-off value for statistical significance. The function intersect in R ['base'-package] was used to find microbiome features that were either both significantly decreased or increased before and after an exacerbation, as compared with during an exacerbation.

2.3.4. Linear analyses: 6 months before and after exacerbations
To test whether the microbiome can be used to monitor CD disease activity, features were linearly associated to the time to the temporally closest CD exacerbation in days. We hypothesised that if microbiota have pathogenic significance in CD, microbiome changes precede changes in disease activity. For this, an arbitrarily chosen period of 6 months was used. This meant that patients who had an exacerbation within 6 months of sampling were included in these analyses. Associations were performed using general linear models in MaAsLin, with the parameters specified above. The R function intersect was used to identify microbiome features that, in an inverse direction, shifted in the days preceding the onset of and restored quiescent balance after an exacerbation. Scripts used to perform data analyses are available at: [https://github.com/WeersmaLabIBD/Microbiome/blob/master/Protocol_ActivityCD_Marjolein_Klaassen.md].

3. Results
3.1. Clinical characteristics
Our cohort consisted of 196 CD patients from whom a single stool sample was collected between 2012 and 2014. At time of sampling, 24 patients [12%] were having an exacerbation, 15 patients [8%] would have their next exacerbation within 6 months, and 19 patients [10%] had had their last exacerbation less than 6 months previously. In addition, 19 patients [10%] would have their next exacerbation more than 6 months after sampling, and 119 patients [70%] had had their last exacerbation more than 6 months before sampling [Table S1, available as Supplementary data as ECCO-JCC online]. The clinical characteristics, medication use, and dietary patterns of the different groups are presented in Table 1. Only PPI use, steroid use, and pre-prepared meals differed between groups, and these features were added to the models [see Methods section].
| Table 1. Clinical characteristics of CD patients corresponding to faecal samples. |
|---------------------------------------------------------------|
| **Before** | **In exacerbation** | **After exacerbation** | **Remission** | **Before during** | **During after** | **Remission** |
| Number of samples | 34 | 24 | 138 | 172 | 224 | 194 | 197 | 249 | 344 | 249 |
| Sequence read depth | 23097318 (7955650) | 2307652 (7521884) | 22252064 (7773052) | 22419149 (7773052) | 0.5503 | 0.9438 | 0.478 | 0.841 | 594557 | 1 |
| Sex (F/M) [%] | 24% (71%) | 35% (65%) | 35% (65%) | 35% (65%) | 0.7579 | 0.7171 | 0.8145 | 0.880 | 802703 | 0.97154839 | 0.931957126 |
| Age [SD] | 37.1 (11.5) | 40.4 (13.9) | 40.4 (13.9) | 40.4 (13.9) | 0.6447 | 0.372 | 0.7664 | 0.841 | 594557 | 0.988 | 3333333 | 0.91542222 | 0.909905263 | 0.57904348 |
| BMI [SD] | 25.1 (5.9) | 23.5 (3.8) | 24.8 (4.8) | 24.8 (5.0) | 0.3024 | 0.4071 | 0.3097 | 0.732 | 222587 | 0.899 | 905263 | 0.57904348 |
| Disease location | 0.307 | 0.118 | 0.116 | 0.127 | 0.6112 | 0.942 | 0.4623 | 0.841 | 594557 | 1 | 0.734071429 |
| Colon [%] | 4 (12%) | 3 (12%) | 33 (24%) | 37 (21%) | 0.3576 | 1.0 | 0.2697 | 0.732 | 222587 | 1 | 0.57904348 |
| Ileum [%] | 13 (38%) | 10 (42%) | 52 (38%) | 65 (37%) | 0.0765 | 0.2680 | 0.0783 | 0.411 | 1875 | 0.865846154 | 0.3741 |
| Behaviour [Montreal] | 19 (56%) | 11 (46%) | 78 (57%) | 97 (57%) | 0.3024 | 0.4071 | 0.3097 | 0.732 | 222587 | 0.899 | 905263 | 0.57904348 |
| Disease duration [SD] | 0.0765 | 0.2680 | 0.0783 | 0.411 | 1875 | 0.865846154 | 0.3741 |
| Age [SD] | 3.0765 | 0.2680 | 0.0783 | 0.411 | 1875 | 0.865846154 | 0.3741 |
| BMI [SD] | 2.0765 | 0.2680 | 0.0783 | 0.411 | 1875 | 0.865846154 | 0.3741 |
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| Behaviour [Montreal] | 19 (56%) | 11 (46%) | 78 (57%) | 97 (57%) | 0.3024 | 0.4071 | 0.3097 | 0.732 | 222587 | 0.899 | 905263 | 0.57904348 |
| Disease duration [SD] | 0.0765 | 0.2680 | 0.0783 | 0.411 | 1875 | 0.865846154 | 0.3741 |
3.2. Grouping based on disease activity

We performed several analyses to check whether the stool samples could be grouped into ‘before’, ‘during’, and ‘after’ an exacerbation; ‘in exacerbation’ or ‘in remission’ and ‘6 months to the next exacerbation’; and ‘6 months since the last exacerbation’, without creating bias.

First, we divided the cohort into patients with CD: a. those who would have their next exacerbation in more than 6 months [\(n = 9\)]; and b. those who would have their next exacerbation within 6 months [\(n = 25\)]; and into patients: c. who had their last exacerbation less than 6 months ago [\(n = 34\)]; and d. who had their last exacerbation more than 6 months ago [\(n = 104\)]. Next, we calculated the differences in inter-individual diversity in functional composition using Bray-Curtis distances, and found that the overall functional compositions of patients in groups a. and b. were similar [\(p = 0.8\) and \(p = 0.3\), respectively; Wilcoxon test]. This was also the case for the patients in groups c. and d. [\(p = 0.07\) and \(p = 0.8\), respectively, Wilcoxon test]. These results indicate that the time to an exacerbation, in both the groups ‘before’ and ‘after’ an exacerbation, is not of influence for the gut microbiome composition. Therefore patients with CD ‘before’ an exacerbation can be considered as a single group, as can the patients with CD ‘after’ an exacerbation.

Second, we combined [I] the patients who were in remission for longer than 6 months [groups b. and c.; \(n = 113\)] and [II] the patients who would have their next or had their last exacerbation within 6 months [groups a. and d.; \(n = 59\)]. Then we calculated the inter-individual diversity in functional composition using Bray-Curtis distances and found that the overall functional composition between patients who are in remission for longer than 6 months have similar microorganisms to those in remission for less than 6 months [PCoA 1 \(p = 0.7\), Wilcoxon test].

Third, we tested whether the gut microbiomes of patients before and after an exacerbation are similar, or in other words, whether we can group the patients who had taken a sample before their next exacerbation [groups a. and b.] with the patients who had their sample taken after an exacerbation [groups c. and d.] into remission together. To test this, we calculated the difference in inter-individual diversity in functional composition using Bray-Curtis distances, and found that the gut microbiome of patients before and after an exacerbation are similar [PCoA 1 \(p = 0.4\), PCoA 2 \(p = 0.8\), Wilcoxon test].

Last, we performed an ADONIS analysis to test the proportion of variance of the gut microbiome which can be explained by the number of days until an exacerbation, and found that days to an exacerbation did not explain a significant proportion of the variance in the gut microbiome [\(R^2 = 0.006\), FDR = 0.272, Table S2, available as Supplementary data at ECCO-JCC online].

We therefore concluded that the patients before an exacerbation, after an exacerbation, and in remission can be grouped as single groups based on their similarities in gut microbiome composition, regardless of the days until an exacerbation.

3.3. Correlations between gut microbiome features and disease activity

3.3.1. Functional but not taxonomical composition is related to CD exacerbations

The gut microbiome of patients during an exacerbation had a similar overall species composition compared with the gut microbiome of patients before [PCoA 1 \(p = 0.3\), PCoA 2 \(p = 0.09\), Wilcoxon test] and after an exacerbation [PCoA 1 \(p = 0.1\), PCoA 2 \(p = 0.8\), Wilcoxon test]. There were no significant differences in taxonomic diversity between patients in an exacerbation and in remission [\(p = 0.88\),
3.3. Monitoring CD disease activity: the microbiome in the six months before and after an exacerbation

We next investigated whether the microbial features associated with an exacerbation also showed a linear correlation with the time [specified in days] before and after an exacerbation. Analyses were confined to patients who experienced an exacerbation in the 6 months before \([n = 15]\) or after \([n = 19]\) their faecal sample was collected.

The proportion of explained variance in both the taxonomic and functional composition of the variable 'days to exacerbation' was not significant \([R = 0.004, FDR = 0.749 \text{ and } R = 0.004, FDR = 0.749, \text{ respectively}]\) \([\text{Table S2}]\). Therefore, correlations between proximity to an exacerbation and changes in the abundance of microbiome features were considered less reliable \([\text{Tables S8-S11, available as Supplementary data at ECCO-JCC online}]\).

4. Discussion

In this cohort of CD patients, we compared the gut microbiome composition of patients in an exacerbation with the microbiome of patients before and after an exacerbation. In previous analyses of this cohort, few pathways were associated with the established individual parameters used to define disease activity [faecal calprotectin levels >200 mg/kg faeces or HBI >4]. In this study, we were not able to link all of these pathways to active disease, which could potentially be explained by the fact that the use of different criteria to define active disease leads to a different grouping of patients with CD based on this disease activity. Furthermore, the effect sizes of the identified pathways could be relatively small. These differences indicate the importance of well defining active disease in the context of CD.

Moreover, a thorough review of the medical records in this study allowed us to use more than just a single marker for disease activity, since individual clinical markers (Crohn's Disease Activity Index [CDAI], HBI) and biomarkers [CRP, faecal calprotectin] are often unreliable and show poor correlation. Although combining these factors with information from the endoscopy results, medication changes, and the gastroenterologist's opinion could be considered unconventional, it does reflect how disease activity is determined in clinical practice. Therefore, we think the additional information could more reliably determine the occurrence of an exacerbation and the correlation with gut microbiome features.

We found that it is not taxonomic distributions but rather function of the microbiome that differs between active and quiescent CD patients. We found that genes encoding 105 microbial pathways were decreased during exacerbation, and these genes mostly exert anti-inflammatory properties by facilitating the biosynthesis and fermentation of various amino acids [tryptophan, methionine, and arginine], B vitamins [riboflavin and thiamine] and short-chain fatty acids [SCFAs]. Notably, all these functional pathways appeared to recover after an exacerbation.

We describe that abundances of genes involved in the fermentation of fibres to acetic acid [SCFA] and pyruvate [main precursor of SCFAs] are decreased during a CD exacerbation, whereas there were no differences in the consumption of fibre-rich foods between the CD patients in an exacerbation and CD patients in remission. This is in congruence with the previously described decreased gene abundance involved in the fermentation of fibres in active treatment-naive patients. SCFAs are the main metabolic end products of microbial fermentation of undigested complex carbohydrates in the human colon. SCFAs induce tolerogenic and anti-inflammatory enterocyte phenotypes and maintain gut integrity by being the
main energy source for enterocytes, by inducing downregulation of pro-inflammatory innate immune cells and their cytokines, and upregulation of anti-inflammatory T regulatory cells, and by increasing the transcription of mucin genes in intestinal goblet cells. A role for SCFAs in controlling inflammation has already been shown in an experimental colitis model in mice, where supplemental dietary SCFAs attenuated colonic inflammation. Decreased abundances of intestinal SCFAs could play a role in intestinal inflammation during CD exacerbations.

We also found that microbial genes encoding the biosynthesis of the B vitamins thiamine [vitamin B1], riboflavin [vitamin B2], and folate [vitamin B9] are decreased during CD exacerbations, which is interesting given established links between these vitamins and reduced inflammation. Previous studies have shown that supplementation of riboflavin and thiamine reduces the production of the pro-inflammatory cytokines TNFα, IL-1, and IL-6. Riboflavin and thiamine have also been shown to intensify the anti-inflammatory activity of the corticosteroid anti-inflammatory drug dexamethasone. Riboflavin injections in mouse models also inhibit the febrile response induced by lipopolysaccharides [LPS]. Microbe-derived riboflavin and folic acid have been described to activate mucosa-associated invariant T [MAIT] cells, and thereby control microbial infection of the gut. Taken together, all these lines of evidence indicate that decreases in the intestinal abundances of thiamine, riboflavin, and folate could be involved in sustaining a CD exacerbation.

An example of an overlapping pathway between this study and our recently published study is the decrease in the abundance of genes predicted to encode the biosynthesis of the amino acid L-arginine in CD patients compared with a general population cohort, as well as a decrease of these genes in CD patients in an exacerbation compared to those in remission. Until this present study and our previous studies, disturbances have not been described in L-arginine, nor in the other amino acid L-tryptophan, in CD disease activity [Figure 2]. However, it has been observed in mice that a decrease in tryptophan metabolism results in deficient aryl hydrocarbon receptor [AHR] activation, leading to susceptibility to colitis. Colonic inflammation was reduced in these mice when they were administered a diet supplemented with synthetic AHR ligands. Moreover, arginine is known as the precursor in the synthesis of polyamines, which are the constituents of intercellular junctions of the gut epithelium, and is therefore of importance in maintaining the integrity of the gut. Decreases in

Figure 1. Microbial function but not taxonomy is related to CD exacerbations. Principal coordinate analyses [PCoA] of Bray-Curtis distances, calculated for [I] taxonomy composition and [II] predicted functional composition. Each small-scale dot represents one faecal microbiome sample, coloured on the moment of faecal sampling being [a] during remission or in an exacerbation [I: PCoA1 p = 0.1, PCoA2 p = 0.6; II: PCoA1 p < 0.01, PCoA2 p = 0.04, Wilcoxon test]; and [b] taken before or in an exacerbation [I: PCoA1 p = 0.3, PCoA2 p = 0.09, II: PCoA1 p = 0.03, Wilcoxon Test], or after an exacerbation [I: PCoA1 p = 0.1, PCoA2 p = 0.8; II: PCoA1 p < 0.01, PCoA2 p = 0.04, Wilcoxon test], with each large-scale centroid representing the mean composition of each patient group. Significant differences were seen in predicted functional, however not in species, composition between patients with remisive and active Crohn’s disease [CD].
Figure 2. Microbial pathway relative abundances before, during and after Crohn’s disease [CD] exacerbations. Violin plots representing relative abundances of genes encoding [a] flavin, [b] thiamine, and [c] folate biosynthesis, [d] homolactic fermentation to short-chain fatty acids [SCFAs], [e] fermentation to acetate and lactate [SCFAs], [f] L-arginine, [g] L-methionine, [h] cis-vaccenate, and [i] gondoate biosynthesis, in faecal samples taken before, during, and after CD exacerbations. To approximate the zero-inflated model used in the MaAsLin analyses (resulting in false-discovery rate [FDR] values), a y-limit >0 was used.
L-tryptophan and L-arginine might play a role in intestinal inflammation during CD exacerbations. Although we were able to observe many differences in functional profiles, we did not observe significant changes in species-level taxonomic composition, virulence content, or growth dynamics. Longitudinal profiling of the variation in microbial species composition has previously shown that species variation within the microbiome is mainly dominated by inter-individual effects. It is metabolic microbial pathways rather than taxonomy that are more conserved among individuals, and pathways might therefore be more appropriate for detecting how the influence of the microbiome mediates CD disease activity during a cross-sectional comparison between active and quiescent patients. Previous studies investigating differences in taxonomic composition between active and inactive disease showed inconsistent results. Collectively, these lines of evidence might explain the lack of identification of individual species associated with active and quiescent CD.

We have compared predicted functional pathways between remissive and active CD using microbial genomic information, but transcriptomic or metabolomic data might provide better insights into the gut environment. However, a recent paper that calculated the ratio between the metagenomic [DNA] and metatranscriptomic [RNA] abundances in stool samples of IBD patients found that the functional potential [DNA] is often proportional to the metatranscriptomic expression [RNA], indicating that the functional genomic estimations are likely to represent changes in microbial genetic expression.

We found that 29% of the patients in an exacerbation had used antibiotics in the 3 months previous to sampling. Even though this was not significantly different compared with the patients before and after an exacerbation [FDR = 0.986 and FDR = 0.579, respectively], we added this factor into our linear models because of the known effects of antibiotic use on gut microbiome composition. In addition, we performed the analyses both with and without steroid use as a covariate in the model, as oral steroid use was correlated with having an exacerbation. Since this correlation between independent variables is hard to take into account in the model, we argued that the main results should be derived from the analyses without steroid use in the model. In addition, our group has previously shown that the use of steroids has no effects on the composition of the gut microbiome. Nevertheless, we display both results in this study.

In our study, we aimed to derive gut microbiome dynamics in CD from cross-sectional data. We created a virtual timeline across all participants by assessing the time since the last exacerbation and the time to the next exacerbation, by re-analysing the medical records of the CD patients a few years after sampling. When performing these analyses, we assumed that uniform patterns could still be detected even with the known inter-individual gut microbiome variation. Categorical analyses [grouping patients into before, during, and after a flare] showed 105 pathways to be decreased during a flare, whereas linear analyses [number of days to next exacerbation, number of days since last exacerbation] did not show significant results, probably due to limited sample size.

We understand that the results of our bioinformatic analyses are less reliable than true time-series experiments and that the results of this study need to be replicated by future studies. However, we do believe that our results are a valuable contribution to the growing knowledge about the gut microbiome in CD, which will enable researchers to study the proposed mechanisms, including in cross-sectional designs. Thorough review of the medical records allowed us to use more than just a single marker for disease activity, since individual clinical markers [for example HBI] and biomarkers [CRP, faecal calprotectin] are often unreliable and show poor correlation. Although combining these factors with information from the endoscopy results, medication changes, and the gastroenterologist’s opinion could be considered unconventional, it does reflect how disease activity is determined in clinical practice. Therefore, we think the additional information could more reliably determine the occurrence of an exacerbation.

In conclusion, we identified pro-inflammatory differences in gut microbiome function during CD disease activity. This work identified roles for several compounds in ameliorating disease activity, including vitamins and amino acids that are already available as dietary supplements at the local drugstore and could thus easily be tested. Our results could therefore be used as novel targets for translational studies trying to modulate the function of the gut microbiome in an anti-inflammatory manner.

**Conflict of Interest**

None declared.

**Author Contributions**

FI and RKW designed the study. FI and VC collected extensive clinical phenotype data of the patients. MAYK reviewed patients’ electronic health records. AVV designed a pipeline to determine microbial profiles from raw metagenomic reads, and processed all samples using this pipeline. MAYK, FI, and VC performed all statistical analyses. MAYK and FI wrote the manuscript. AVV, RG, JF, HMD, GD, EAMF, CW, AZ, and RKW critically reviewed the manuscript.

**Supplementary Data**

Supplementary data are available at ECCO-JCC online.

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