Healthy Cotwins Share Gut Microbiome Signatures With Their Inflammatory Bowel Disease Twins and Unrelated Patients

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BACKGROUND & AIMS: It is currently unclear whether reported changes in the gut microbiome are cause or consequence of inflammatory bowel disease (IBD). Therefore, we studied the gut microbiome of IBD-discordant and -concordant twin pairs, which offers the unique opportunity to assess individuals at increased risk of developing IBD, namely healthy cotwins from IBD-discordant twin pairs. METHODS: Fecal samples were obtained from 99 twins (belonging to 51 twin pairs), 495 healthy age-, sex-, and body mass index–matched controls, and 99 unrelated patients with IBD. Whole-genome metagenomic shotgun sequencing was performed. Taxonomic and functional (pathways) composition was compared among healthy cotwins, IBD-twins, unrelated patients with IBD, and healthy controls with multivariable (ie, adjusted for potential confounding) generalized linear models. RESULTS: No significant differences were observed in the relative abundance of species and pathways between healthy cotwins and their IBD-twins (false discovery rate <0.10). Compared with healthy controls, 13, 19, and 18 species, and 78, 105, and 153 pathways were found to be differentially abundant in healthy cotwins, IBD-twins, and unrelated patients with IBD, respectively (false discovery rate <0.10). Of these, 8 (42.1%) of 19 and 1 (5.6%) of 18 species, and 37 (35.2%) of 105 and 30 (19.6%) of 153 pathways overlapped between healthy cotwins and IBD-twins, and healthy cotwins and unrelated patients with IBD, respectively. Many of the shared species and pathways have previously been associated with IBD.
The shared pathways include potentially inflammation-related pathways, for example, an increase in propionate degradation and L-arginine degradation pathways. **CONCLUSIONS:** The gut microbiome of healthy cotwins from IBD-discordant twin pairs displays IBD-like signatures. These IBD-like microbiome signatures might precede the onset of IBD. However, longitudinal follow-up studies are needed to infer a causal relationship.

**Keywords:** Preclinical; Prediagnostic; Crohn’s Disease; Ulcerative Colitis; Family Studies; Microbiota; Prediction; Discordant Twin Design.

Over the past decade, a consistent body of evidence has accumulated that supports the association between the gut microbiota and inflammatory bowel disease (IBD). Differences have been observed in the microbiome of Crohn’s disease (CD) and ulcerative colitis (UC) patients compared with individuals not affected by IBD in a large number of studies, most of which were based on 16S ribosomal RNA gene (16S rRNA) sequencing. It is, however, currently unclear whether these changes are a cause or a consequence of intestinal inflammation, because longitudinal microbiome studies are scarce and data from the pre-diagnostic phase of IBD are lacking. Moreover, the large interindividual heterogeneity induced by the effect of (childhood) environmental factors and host genetics on the gut microbiome hampers the elucidation of the exact contribution of the gut microbiome to the onset of IBD.

In the context of IBD, healthy cotwins with an IBD-affected twin are at increased risk of developing IBD, with reported concordance rates among monozygotic twin pairs as high as 64% for CD and 28% for UC. Studying the gut microbiome in IBD-discordant and concordant twin pairs can thus provide important insights in its role in the pathogenesis of IBD, also because of the shared (childhood) environmental and genetic factors between twins from the same twin pair.

Five previous studies, aiming to explore the gut microbiome in IBD-affected twin pairs, using fecal samples, reported differences in the gut microbiome composition in IBD-affected twins compared with their healthy cotwin. These studies were, however, performed in small numbers of IBD-discordant or -concordant twin pairs (10 or fewer), did not include an unrelated matched healthy control group, or only a small nonmatched healthy control group and were based on 16S rRNA sequencing, which does not allow for prediction of microbial functional pathways.

The goal of the present study was to elucidate the contribution of the gut microbiota in the risk of developing IBD, by exploring the gut microbiome of healthy cotwins - at increased risk of developing IBD - in comparison with the gut microbiome of their IBD-twins, unrelated patients with IBD, and those of unrelated healthy controls.

**Material and Methods**

**Study Population**

Participants from 2 cohorts were included in the present cross-sectional study.

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**WHAT YOU NEED TO KNOW**

**BACKGROUND AND CONTEXT**

Changes in the composition of the gut microbiome are associated with inflammatory bowel disease (IBD), but it is presently unclear whether these changes are cause or consequence of IBD.

**NEW FINDINGS**

The gut microbiome composition of individuals at increased risk of developing IBD (i.e. healthy cotwins from IBD-discordant twin pairs) displays IBD-like signatures on a species and pathway level.

**LIMITATIONS**

This is a cross-sectional study. Future follow-up studies will help to identify those cotwins who will develop IBD in the long run, thereby allowing confirmation and further in-depth analyses of our findings.

**IMPACT**

The overlap in gut microbial features between healthy cotwins at increased risk of developing IBD and related and unrelated IBD patients suggests that these IBD-like microbiome signatures might precede the onset of IBD. This potentially opens new avenues for diagnosis and therapy in individuals with pre-symptomatic IBD.

(1) IBD-twins and healthy cotwins: “TWIN-study”. Twin pairs were included from the ongoing prospective longitudinal Dutch “Twin cohort for the study of (pre)clinical inflammatory bowel disease in the Netherlands” (TWIN) study (Netherlands Trial Register: NTR6681). Twin pairs, ≥16 years of age, either IBD-discordant or -concordant, from the Netherlands, were recruited via their treating physician, through awareness for the study on (social) media, or via the Netherlands Twin Register (NTR). Potential candidates in the NTR were identified through previously administered questionnaires or those who gave consent for record linkage were identified by linking the NTR-database with an IBD-directed search in the nationwide network and registry of histo- and cytopathology in the Netherlands (PALGA). Participants were followed longitudinally and evaluated at intervals of 6 months at the University Medical Center Utrecht. Samples of blood, urine, feces, oropharyngeal swabs, and (depending on consent) rectal biopsies were collected.

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**Abbreviations used in this paper:** 16S rRNA, 16S ribosomal RNA; BMI, body mass index; CD, Crohn’s disease; FDR, false discovery rate; IBD, inflammatory bowel disease; IBDU, inflammatory bowel disease unclassified; NTR, Netherlands Twin Register; PALGA, Nationwide network and registry of histo- and cytopathology in the Netherlands; PCoA, principal coordinate analyses; PERMANOVA, permutational multivariate analysis of variance; PPI, proton pump inhibitor; SCFA, short chain fatty acid; TWIN-study, twin cohort for the study of (pre)clinical inflammatory bowel disease in the Netherlands study; UC, ulcerative colitis; UMC, University Medical Center.

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during study visits. In addition, questionnaires were used to obtain information on demographics, family composition, diet, general health, IBD-specific characteristics, medication use, quality of life, and environmental factors.

For the present cross-sectional study, we analyzed 1 fecal sample per participant, collected between October 2017 and August 2019. Fecal samples of a twin pair were not included for whole-metagenome sequencing if only 1 twin from a pair had provided a fecal sample or when a twin had an ostomy or pouch. If the whole-metagenome shotgun sequencing results of a twin were excluded after sequencing, we kept the results of their cotwin in the data analyses. All fecal samples were collected within the same week per twin pair, except for 1 twin pair (6 months between feces collection). Both IBD-discordant (ie, one of the twins of a pair is affected by IBD) and IBD-concordant (ie, both twins of the twin pair are affected by IBD regardless of the IBD subtype) twin pairs are included in the present study. No multiples consisting of more than 2 individuals were included. Individuals from IBD-discordant twin pairs who are not diagnosed with IBD are referred to as “healthy cotwins” and individuals from IBD-discordant or IBD-concordant twin pairs who are diagnosed with IBD are referred to as “IBD-twins.”

(2) Healthy controls and unrelated patients with IBD: “Dutch Microbiome Project”. The Dutch Microbiome Project is part of LifeLines,10 a large-scale population-based cohort that prospectively included 167,729 individuals between 8 and 84 years of age and their families, who all have been resident in the 3 Northern provinces of the Netherlands. In 2015, 10,000 LifeLines participants were asked to collect a fecal sample as part of the Dutch Microbiome Project. Of these, ~9700 individuals collected 1 fecal sample between 2015 and 2018. Demographics, medical information, and medication use were collected via surveys. For the present cross-sectional study, we included age-, sex- and BMI-matched healthy participants from the Dutch Microbiome Project. Healthy individuals were matched to each twin individual (ie, healthy cotwins and IBD-twins) with a healthy control:twino ratio of 5:1. Matching was performed with the MatchIt package in R using the method “optimal”.

Within the Dutch Microbiome Project, 99 participants had a self-reported diagnosis of IBD. These individuals were included in this study as “unrelated patients with IBD.”

Ethical Considerations

The TWIN-study and Dutch Microbiome Project study were conducted in accordance with the declaration of Helsinki and the Dutch Medical Research Involving Human Subjects Act. All participants provided informed consent. The TWIN-study was approved by the medical ethics committee of the UMC Utrecht. The Dutch Microbiome Project was approved by the medical ethics committee of the UMC Groningen. The PALGA search was approved by the PALGA Privacy Commission and Scientific Council. Linkage took place only for NTR participants who had provided approval for linkage to external registers.

Participant Characteristics

IBD characteristics. For the TWIN cohort, diagnoses and phenotypes were verified by review of medical records. For participants in the Dutch Microbiome Project, diagnoses and phenotypes were self-reported. IBD unclassified (IBDU) patients were grouped with UC patients in the analyses. In the TWIN-study signs of active disease was defined as a patient Harvey Bradshaw Index19 >4 in case of CD, and a patient Short Clinical Colitis Activity Index20 >4 in case of UC or IBDU, or endoscopic signs of inflammation as noted during proctoscopy in the TWIN-study or endoscopy for clinical care. If no signs of inflammation were found on proctoscopy in patients with UC, patients were classified as having quiescent disease. In the Dutch Microbiome Project, the Montreal classification, date of diagnosis, symptoms, and endoscopic data were not available.

Twin-specific characteristics. The zygosity of twins was based on self-reported zygosity or, in case of doubt, on a zygosity questionnaire, based on childhood similarity between twins. This questionnaire was developed by the NTR and has been shown to have a 95.9% accuracy in predicting DNA zygosity.16 Cohabitation was based on survey data.

Demographics. In both the TWIN-study and the Dutch Microbiome Project, data on sex, age, body mass index (BMI), smoking behavior, history of appendectomy, and history of bowel resections were collected by surveys at the moment of feces collection.

Medication use. Medication use, including antibiotics (in the past 3 months), current proton pump inhibitors (PPIs), and IBD medication, was determined by targeted medication surveys in the TWIN-study and the Dutch Microbiome Project.

Stool Sampling and Analysis of the Gut Microbiome

Fecal sample collection. In the TWIN-study, fecal samples were kept at room temperature and transported to the research facility by the participants within 31 hours of collection, and subsequently stored at −80°C (median time at room temperature until −80°C: 9.7 hours; Q1–Q3: 4.8–19.8). In the Dutch Microbiome Project, fresh fecal samples were frozen at the participants’ homes in a standardized manner at −20°C. Subsequently, frozen fecal samples were shipped on dry ice and stored at −80°C on arrival. Fecal samples from both cohorts remained frozen at −80°C until DNA extraction.

Microbial DNA extraction and sequencing. Microbial genomic DNA was isolated via the QIAamp Fast DNA Stool Mini Kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions by the same research technician in the same time period for both cohorts. The QIAcube (Qiagen) automated sample preparation system was used for DNA isolation. Whole-genome shotgun metagenomic sequencing was performed at Novogene (HK) Company Limited (Wan Chai, Hong Kong) on the Illumina (San Diego, CA) HiSeq 2000 platform to generate approximately 8 Gb of 150 base pairs, paired-end, reads per sample (mean 7.9 gigabytes, standard deviation 1.2 gigabytes, median 14.4 million reads).

Quality control and determination of microbiome parameters. Samples with a read depth below 10 million reads were removed. From the raw metagenomic reads, the Illumina adapters were removed using kneadData (v0.5.1) toolkit, and reads were trimmed to PHRED quality 30. Trimmed reads that aligned to the human genome (GRCh37/hg19) were removed using kneadData integrated Bowtie2 tool (version 2.3.4.1), and after this, the quality of the metagenomes was tested using the FastQC toolkit (version 0.11.7). Taxonomic composition of the metagenomes was profiled by the MetaPhlAn2 tool (version 2.7.2) using the MetaPhlAn database of marker genes (version mpa v20 m200). In addition, profiling of
genes encoding microbial biochemical pathways (ie, the functional potential of the gut microbiome) was performed using the HUMAnN2 pipeline (version 0.11.1) integrated with the DIAMOND alignment tool (version 0.8.22), uniref90 protein database (version 0.1.1), and the ChocoPhlAn pan-genome database (version 0.1.1).

Non-bacteria were filtered out and the taxonomic species level was maintained. After these steps, 586 species and 576 pathways were identified across samples. For the diversity and dissimilarity analyses, no further filtering steps were used. For the regression analyses, only species and pathways that were prevalent in ≥10% of samples were included.

Design of Data Analysis

To answer our question of to which extent the gut microbiome of healthy cotwins displays a signature of an IBD-like microbiome, we performed our analyses in 2 steps. First, we compared the healthy cotwins with their IBD-twins (within-twin pairs comparison). Second, we compared the gut microbiome of healthy cotwins, IBD-twins, and unrelated patients with IBD to the microbiome of healthy controls. By means of these comparisons, we aimed to identify whether the microbiomes of healthy cotwins display a healthy signature (ie, is similar to the gut microbiome of unrelated healthy controls) or already displays an IBD signature (ie, is more similar to the microbiome of their cotwins and unrelated patients with IBD).

We analyzed the gut microbiome at the following levels: (1) α- and β-diversity; (2) (dis)similarity in gut microbiome composition among individuals based on IBD concordance, IBD phenotype, zygosity, and cohabitation; and (3) differential relative abundances of individual species and pathways including assessment of overlap in identified differentially (compared with healthy controls) abundant species and pathways among the healthy cotwins, IBD-twins, and unrelated patients with IBD.

Statistical Analyses

All statistical analyses were performed in R version 4.0.0 for macOS21 (analyses scripts can be found at: https://github.com/WeersmaLabIBD/Microbiome/blob/master/IBD_Twins_Microbiome_Utrecht_Groningen.md). Baseline characteristics are shown as numbers and proportions for categorical variables, and as means and standard deviations for continuous variables.

Microbiome diversity measures. We estimated diversity measures for both for the taxonomic (ie, species) and functional (ie, pathways) composition. The α-diversity is expressed as the Shannon Index for species and gene richness for pathways. Differences between groups were tested with the Mann-Whitney U test. The β-diversity was calculated as Bray-Curtis distance and is visually shown using principal coordinate analyses (PCoA) plots, showing PCoA1 and PCoA2 per plot. The proportion of explained variance ($R^2$) in the β-diversity was assessed based on permutational multivariate analysis of variance (PERMANOVA) using distance matrices implemented as adonis function in vegan package for R.

For the within-twin pair comparisons, we estimated the proportion of explained variance ($R^2$) in the β-diversity adjusting for twin pairs, thereby taking the clustering of twins within their twin pair into account, and stratified for zygosity for the following variables separately: IBD phenotype (no IBD, CD, or UC), IBD-activity, belonging to the same twin pair, disease location, history of bowel resection, age, sex, BMI, antibiotics use, PPI use, and read depth. For the comparison of healthy cotwins, IBD-twins, and unrelated patients with IBD with healthy controls, we estimated the explained variance in β-diversity comparing IBD-twins, healthy cotwins, healthy controls, and unrelated patients with IBD, and for the variables IBD phenotype, sex, age, BMI, antibiotics use, PPI use, and read depth.

Similarity in gut microbiome composition. To test whether the gut microbiome composition was similar between pairs of individuals on a taxonomic or functional level, the pairwise Bray-Curtis dissimilarity was calculated, ranging from 0 to 1 with lower scores indicating more similarity in the microbiome composition. We assessed the associations between dissimilarity and 4 variables: IBD concordance (ie, being discordant or concordant for IBD), IBD phenotype, zygosity, and cohabitation. To this end, dissimilarities between twins from the same twin pair were calculated and compared with the dissimilarities of 1000 random pairs of unrelated healthy controls. IBD phenotype pairs were logically formed between unrelated IBD-twins. For zygosity, we also compared the dissimilarity with random pairs of unrelated twins (ie, pairs of twins coming from 2 different twin pairs).

Differentially abundant taxa and pathways. We assessed whether taxa and pathways were differentially abundant by using multivariable general linear regression models (GLM) of the Gaussian family, implemented in the GLM function in R, using the MaAsLin2 package.22 The microbiome taxa and pathway relative abundances that serve as outcomes in the GLMs were arcsine square-root transformed. Species and pathways were included in the analyses only if they were present in both cohorts and were prevalent in at least 10% of the samples.

For the comparison between IBD-twins and healthy cotwins, we used twin pair as a random effect to take within-twin pair effects (ie, clustering) into account. Furthermore, the regression analyses with 119 species and 343 pathways as outcomes were adjusted for IBD phenotype (ie, CD, UC, or no IBD), disease location, age, sex, BMI, zygosity, antibiotics use in the past 3 months, current PPI use, and sequence read depth. Second, we assessed differential abundance of taxa and pathways comparing healthy cotwins, IBD-twins, and unrelated patients with IBD with healthy controls in one model. These regression analyses with 116 individual species and 330 individual pathways as outcomes were adjusted for potentially confounding factors: IBD phenotype (ie, CD, UC, or no IBD), age, sex, BMI, antibiotics use in the past 3 months, current PPI use, and sequence read depth. Next, we assessed if overlap existed in the identified differentially abundant taxa and pathways as present in the microbiomes of healthy cotwins, IBD-twins, and unrelated patients with IBD compared with healthy controls. Last, we compared the relative abundances of 112 species and 339 pathways between IBD-twins and unrelated patients with IBD directly adjusting for the same potentially confounding factors as in the comparison with healthy controls.

Multiple testing correction. To correct for multiple testing, the Benjamini-Hochberg correction was applied to calculate the false discovery rate (FDR). An FDR <0.10 was regarded as statistically significant. This significance threshold
### Table 1. Baseline and Sample Characteristics

| The TWIN-study | The Dutch Microbiome Project |
|----------------|-----------------------------|
| **Healthy cotwins**<br>(n = 38) | **IBD-twins**<br>(n = 61) | **Healthy controls**<br>(n = 495) | **Unrelated IBD patients**<br>(n = 99) |
| **Demographics and clinical characteristics** | | | |
| Female sex | 25 (65.8) | 39 (63.9) | 312 (63.0) | 59 (59.6) |
| Age (y) | 44.7 (15.4) | 40.1 (15.4) | 42.3 (15.0) | 52.8 (10.0) |
| BMI (kg/m²) | 24.9 (3.8) | 24.2 (3.3) | 24.4 (4.0) | 26.0 (3.9) |
| Current smoker | 6 (15.8) | 10 (16.4) | 49 (9.9) | 14 (14.1) |
| History of appendectomy | 4 (10.5) | 6 (9.8) | 29 (5.9) | 11 (11.1) |
| History of bowel resection | 0 | 10 (16.4) | NA | NA |
| Current PPI use | 6 (15.8) | 9 (14.8) | 24 (4.8) | 12 (12.1) |
| Antibiotic use in past 3 mo | 3 (7.9) | 8 (13.1) | 23 (4.6) | 10 (10.1) |
| **Twin pair characteristics**<sup>a</sup> | | | |
| Zygosity | | | |
| Monozygotic | 16 (42.1) | 37 (60.7) | NA | NA |
| Dizygotic | 22 (57.9) | 24 (39.3) | NA | NA |
| Concordance for IBD | | | |
| Concordant for IBD | 0 | 24 (39.3) | NA | NA |
| Discordant for IBD | 38 (100) | 37 (60.7) | NA | NA |
| Cohabitation with cotwin during sampling | 4 (10.5) | 10 (16.4) | NA | NA |
| **IBD characteristics** | | | |
| IBD phenotype | | | |
| CD | — | 33 (54.1) | — | 28 (28.3) |
| UC | — | 26 (42.6) | — | 72 (72.7) |
| IBD unclassified | — | 2 (3.3) | — | 0 |
| No IBD | 38 (100) | — | 495 (100) | — |
| IBD duration (mo) | NA | 139 (112) | NA | NA |
| **Signs of active disease**<sup>b</sup> | | | |
| Age of diagnosis CD (Montreal classification)<sup>c</sup> | | | |
| A1 (<16 y) | NA | 3 (9.1) | NA | NA |
| A2 (17–39 y) | NA | 26 (78.8) | NA | NA |
| A3 (>40 y) | NA | 4 (12.1) | NA | NA |
| Location CD (Montreal classification)<sup>c</sup> | | | |
| L1 (ileum only) | NA | 13 (39.4) | NA | NA |
| L2 (colon only) | NA | 7 (21.2) | NA | NA |
| L3 (ileocolonic) | NA | 13 (39.4) | NA | NA |
| L4 (proximal of ileum) | NA | 4 (12.1) | NA | NA |
| Behavior CD (Montreal classification)<sup>c</sup> | | | |
| B1 (nonstricturing, nonpenetrating) | NA | 18 (54.5) | NA | NA |
| B2 (stricturing) | NA | 11 (33.3) | NA | NA |
| B3 (penetrating) | NA | 4 (12.1) | NA | NA |
| P (perianal modifier) | NA | 3 (9.1) | NA | NA |
| Location UC (Montreal classification)<sup>c</sup> | | | |
| E1 (ulcerative proctitis) | NA | 6 (23.1) | NA | NA |
| E2 (left-sided UC) | NA | 7 (26.9) | NA | NA |
| E3 (extensive UC) | NA | 11 (42.3) | NA | NA |
| Missing | NA | 2 (7.7) | NA | NA |
| Current IBD-medication use | | | |
| No IBD-medication | 38 (100) | 10 (16.4) | 495 (100) | NA |
| 5-aminosalicylic acid | — | 20 (32.8) | — | NA |
Table 1. Continued

|                  | The TWIN-study | The Dutch Microbiome Project |
|------------------|----------------|-----------------------------|
| Corticosteroids  | NA             | 3 (4.9)                     |
| Methotrexate     | NA             | 1 (1.6)                     |
| Ciclosporin      | NA             | 0                           |
| Thiopurine       | NA             | 24 (39.3)                   |
| Anti-tumor necrosis factor-α | NA | 12 (19.7) |
| Vedolizumab (anti-integrin α4β7) | NA | 2 (3.3) |
| Ustekinumab (anti-interleukin12/23) | NA | 0 |
| Tofacitinib (pan-janus kinase inhibitor) | NA | 0 |

Fecal sample and sequence characteristics

Bristol stool scale\(^d\)

| Type 1 or 2 | 7 (18.4) | 10 (16.4) | 47 (9.5) | 6 (6.1) |
| Type 3 or 4 | 26 (68.4) | 23 (37.7) | 349 (70.5) | 53 (53.5) |
| Type 5, 6, or 7 | 5 (13.2) | 28 (45.9) | 68 (13.7) | 34 (34.3) |
| Missing      | 0 | 0 | 31 (6.3) | 6 (6.1) |

Number of bowel movements per wk 7.4 (2.71) 12.5 (11.5) 8.3 (4.17) Missing: n = 31 n = 6

Number of sequence reads 16,108,523 (4,134,472) 15,271,177 (3,989,233) 24,057,674 (3,552,760) 25,111,866 (4,072,961)

NOTE. Continuous variables are depicted as mean (standard deviation), and categorical variables as number (proportion) unless indicated otherwise.

n, number of participants; NA, not applicable.

\(a\)All twin-related characteristics are here depicted per individual, but logically apply to both of the twin pair.

\(b\)Signs of active disease was defined as Harvey Bradshaw Index19 >4 in case of CD, and a Short Clinical Colitis Activity Index20 >4 in case of UC or IBD unclassified, or endoscopic signs of inflammation as noted during either proctoscopy performed as part of the TWIN-study protocol or during a colonoscopy which was performed as part of clinical care.

\(c\)All proportions for Montreal classification characteristics are only calculated for those participants to whom it applies.

\(d\)The Bristol stool scale ranges from 1 to 9, with 1 being separate hard lumps, and 9 entirely liquid consistency without solid pieces. In the healthy controls and unrelated patients with IBD this reflects the mean over a week.

\(e\)These variables are not assessed in the Dutch Microbiome Project.

Figure 1. No differences were detected in the gut microbiome composition (beta-diversity) between healthy cotwins and IBD-twins, but both gut microbiomes differ from healthy controls. PCoA plots for the (A) taxonomic (species) and (B) functional (pathways) composition of the gut microbiome. Each small dot represents 1 fecal sample. The larger centroids depict the center per group (ie, healthy cotwins, IBD-twins, healthy controls, and unrelated patients with IBD). On the x- and y-axes the first and second principal coordinate and proportion of explained variance are displayed. Based on the FDR from univariable PERMANOVA analyses performed on the whole Bray-Curtis distance matrix (Supplementary Material 4), the healthy cotwins and IBD-twins are not significantly different from each other on a taxonomic and functional level, whereas the gut microbiome composition was statistically significantly different comparing healthy cotwins, IBD-twins, and unrelated patients with IBD with healthy controls. **, FDR < 0.05; ns, FDR > 0.10.
was chosen because of the explorative setup of our study, to capture both common associations and those with low effect sizes.

Results

Participant and Sample Characteristics

In total, 99 twins from 51 twin pairs (61/99 IBD-twins, and 38/99 healthy cotwins) from the TWIN-study, and 495 age-, sex-, and BMI-matched unrelated healthy controls and 99 unrelated patients with IBD from the Dutch Microbiome Project were included in the present study (Table 1 and Supplementary Materials 1 and 2). Fifty-three of the 99 twins were part of a monozygotic twin pair and 46 were part of a dizygotic twin pair. Twelve (23.5%) of the 51 twin pairs were IBD-concordant and 39 (76.5%) IBD-discordant. Seven (13.7%) of the 51 twin pairs were living together at the time of fecal sampling (Table 1 and Supplementary Material 2).

Healthy controls from the Dutch Microbiome Project had a comparable age, sex, and BMI, but smoked less frequently as compared with all twins, whereas the unrelated patients with IBD were older and had a higher BMI compared with the TWIN cohort. A history of appendectomy and antibiotic or PPI use was encountered more frequently in twins and unrelated patients with IBD than in healthy controls. IBD-twins were compared with the unrelated patients with IBD more often diagnosed with CD (54.1% vs 28.3%, Table 1). The sequence read depth was higher in healthy controls and unrelated patients with IBD as compared with the TWIN participants (Table 1), the linear regression analyses were therefore adjusted for sequence read depth.

Healthy Cotwins and IBD-Twins Are Alike in Microbiome Diversity and Differ From Healthy Controls

There were no significant differences in $\alpha$-diversity, that is, the per-participant diversity, on taxonomic and functional level between the healthy cotwins and the IBD-twins (FDR = 0.14, FDR = 0.62, respectively, Supplementary Material 3). Furthermore, no statistically significant differences in the Shannon Index were observed between healthy
cotwins, IBD-twins, unrelated patients with IBD, and healthy controls. The gene richness was statistically significantly higher in healthy cotwins compared with healthy controls, and IBD-twins compared with healthy controls (FDR = 0.007, FDR = 0.007, respectively, Supplementary Material 3).
No statistically significant differences were found in the gut microbiome of the healthy cotwins compared with the IBD-twins for the overall taxonomic (PERMANOVA of Bray-Curtis β-diversity: $R^2 = 0.015$, $FDR = 0.12$) and functional (PERMANOVA: $R^2 = 0.011$, $FDR = 0.36$) composition (Figure 1, Supplementary Material 4). The overall taxonomic and functional composition was different for healthy cotwins, IBD-twins and unrelated patients with IBD compared with healthy controls (PERMANOVA: taxonomic: $R^2 = 0.004$, 0.012, 0.010; $FDR = 0.012$, 0.0002, 0.0002; functional: $R^2 = 0.013$, 0.012, 0.020; $FDR = 0.0002$, 0.0002, 0.0002 respectively, Figure 1, Supplementary Material 4). Overall, no differences between the gut microbiome composition of healthy cotwins and IBD-twins were detected, whereas both groups’ microbiome composition differed from healthy controls.

Association Between IBD Concordance, IBD Phenotype, Zygosity, and Co-housing and the Composition of Gut Microbiome

The associations of IBD concordance, IBD phenotype, zygosity, and co-housing during sampling with the interindividual heterogeneity in gut microbiome composition were assessed by using the Bray-Curtis dissimilarity by comparing 2 individuals. In line with our observations for the similarity in microbiome diversity between healthy cotwins and IBD-twins, we found no differences in the dissimilarity between IBD-discordant and IBD-concordant twin pairs, both at a taxonomic and functional levels (Figure 2A). When looking at unrelated random pairs of healthy controls and unrelated random pairs of twins, we observed an increase in dissimilarity from healthy, healthy cotwins, to UC, to CD, underscoring the heterogeneity in changes in the gut microbiome composition in IBD (Figure 2B). The gut microbiomes of monozygotic or dizygotic twin pairs or twin pairs co-housing during sampling was not more similar than those of random unrelated healthy controls (Figure 2C and D).

Healthy Cotwins, IBD-Twins, and Unrelated IBD Patients Differ in Relative Abundance of Species and Pathways From Healthy Controls

To identify whether the gut microbial features of the healthy cotwins are more alike to those displayed in an IBD microbiome (ie, IBD-twins and unrelated patients with IBD) or more alike to those displayed in healthy controls, we performed multivariable linear regression analyses adjusted for IBD subtype (CD, UC, and no IBD), age, sex, BMI, antibiotics and PPI use, and sequencing depth. In the within-twin pairs analyses, additionally adjusted for zygosity and disease location, and with twin pair as random effects, none of the 119 species and 343 predicted bacterial pathways tested were differentially abundant between the healthy cotwins and the IBD-twins (FDR > 0.1, Supplementary Material 5). However, compared with the healthy controls, 19 species and 105 pathways were differentially abundant within the gut microbiome of IBD-twins, and 18 species and 153 pathways in unrelated patients with IBD (Figure 3, and Supplementary Materials 6 and 7, FDR < 0.1). The relative abundance of 13 species and 78 pathways differed significantly between microbiomes of healthy cotwins and healthy controls (Figure 3, Supplementary Material 8).

The number of species and pathways that were differentially abundant when we compared IBD-twins and unrelated patients with IBD directly, 7 species and 34 pathways (Supplementary Material 9 ), was less than when we compared both groups with healthy controls. In combination with the overlap of 7 species and 70 pathways that were differentially abundant compared with healthy controls, this indicates that the gut microbiome composition of IBD-twins and unrelated patients with IBD was comparable.

Healthy Cotwins Share Differentially Abundant Taxa and Pathways With IBD-Twins and Unrelated IBD Patients Compared With Healthy Controls

Of the 19 species and 105 pathways that were differentially abundant between IBD-twins compared with healthy controls, 8 of these species (42.1%) and 37 pathways (35.2%) were also differentially abundant between the healthy cotwins compared with healthy controls. Moreover, of the 18 species and 153 pathways that were differentially abundant between the unrelated patients with IBD and the healthy controls, 1 of these species (5.6%) and 30 pathways (19.6%) were also differentially abundant between the healthy cotwins compared with the healthy controls (Figure 3 and Supplementary Material 10).

Among these, an increase in the relative abundance of potentially pathogenic species *Escherichia unclassified*, *Gordonibacter pamelaeae*, and *Eggerthella unclassified*, among others, was observed in the gut microbiomes of healthy cotwins and IBD-twins or unrelated patients with IBD, as compared with the healthy controls (Figures 3B and 4A–C). *Faecalibacterium prausnitzii* was only statistically
Figure 4. Relative abundance of a selection of species. The relative abundance of IBD-associated species, (A) *Escherichia unclassified*, (B) *Eggerthella unclassified*, (C) *Gordonibacter pamelaeae* is increased among healthy cotwins and shared differentially abundant between IBD-twins and/or unrelated patients with IBD compared with healthy controls. The relative abundance of (D) *Faecalibacterium prausnitzii*, often inversely associated with IBD, is decreased in unrelated patients with IBD and IBD-twins, but not in healthy cotwins compared with healthy controls. The FDR values are based on multivariable general linear model analyses adjusted for age, sex, BMI, antibiotics use, PPI use, IBD phenotype, and sequence read depth.

ns = FDR $\geq 0.1$; * = FDR < 0.1; ** = FDR < 0.05; *** = FDR < 0.025
significantly decreased compared with healthy controls in IBD-twins and unrelated patients with IBD, but not in healthy cotwins (Figures 3B and 4D).

Two pathways involved in the biosynthesis of family-specific surface enterobacterial common antigen, which is specific to the family Enterobacteriaceae, among which the genus *Escherichia*23 (ECASYN_PWY_enterobacterial_common_antigen_biosynthesis, 7315_dTDP_N_acetylthomosamine_biosynthesis) were increased in the healthy cotwins, IBD-twins, and unrelated patients with IBD, as compared with the healthy individuals (FDR <0.1) (Figures 3C and 5A and 5B). The relative abundance of pathways for the biosynthesis of siderophores, iron-chelating molecules that are known potential virulence factors,24 were increased in healthy cotwins, IBD-twins, and unrelated patients with IBD, namely enterobactin, (ENTBACSYN_PWY_enterobactin_biosynthesis, Figure 5C), and in healthy cotwins and IBD-twins, namely aerobactin (AEROBACTINSYN_PWY_aerobactin_biosynthesis, Figure 3C). Moreover, the relative abundance of genes encoding the degradation of amino acid L-arginine, which is known to promote gut integrity via tight junctions between enterocytes,25,26 (AST_PWY_L_arginine_degradation_II_AST_pathways) was increased in healthy cotwins, IBD-twins, and unrelated patients with IBD (Figure 5D) and the degradation of short-chain fatty acid (SCFA) propionate (PWY0_1277_3 phenylpropionate_and_3_3_hydroxyphenylpropanoate_degradation, HCAMHPDEG_PWY_3 phenylpropanoate_and_3_3_hydroxyphenyl_propanoate_degradation_to_2 oxopent_4 enoate) was increased in the gut microbiomes of healthy cotwins and IBD-twins, as compared with the healthy individuals (Figure 5E, FDR <0.1). In line with the decrease in

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**Figure 5.** Relative abundance of a selection of pathways. The relative abundance of pathways of (A) enterobacterial common antigen biosynthesis, (B) N-acetylthomosamine (an enterobacterial common antigen component) biosynthesis, (C) enterobactin (a siderophore and virulence factor) biosynthesis, (D) L-arginine degradation, and (E) propionate (an SCFA) degradation are increased in healthy cotwins and shared differentially abundant in the gut microbiomes of IBD-twins and unrelated patients with IBD compared with healthy controls. (F) Butyrate biosynthesis is decreased in unrelated patients with IBD and IBD-twins, but not in healthy cotwins compared with healthy controls. The FDR values are based on multivariable general linear model analyses adjusted for age, sex, BMI, antibiotics use, PPI use, IBD phenotype, and sequence read depth.
the relative abundance of *F. prausnitzii*, the relative abundance of 2 butyrate synthesis pathways (CENTFERM_PWY_pyr-uvate_fermentation_to_butanoate and 6590_superpathway_of_Clostridium_acetobutylicum_acidogenic_fermentation) were only statistically significantly decreased in IBD-twins and unrelated patients with IBD, and not in healthy cotwins (Figure 5F), although 1 butyrate biosynthesis pathway was decreased in healthy cotwins and IBD-twins compared with healthy controls (5676_acetyl_CoA_fermentation_to_butanoateII).

Taken all together, an increase in potentially pathogenic species and proinflammatory pathways was noted in the gut microbiome of healthy cotwins of IBD-discordant twin pairs compared with healthy controls. A substantial proportion of these differences are shared with IBD-twins and unrelated patients with IBD.

**Discussion**

In this unique cross-sectional study analyzing the gut microbiome in IBD twin pairs, using metagenomic shotgun sequencing, we observed no differences in α and β diversity and the relative abundance of individual species and pathways between healthy cotwins and their IBD-twins. Compared with age-, sex-, and BMI-matched healthy controls, healthy cotwins displayed a large overlap in differentially abundant species and pathways, not only with their IBD-twins, but also with unrelated patients with IBD. Among these overlapping species and pathways, species associated with IBD and potentially inflammation-related pathways were present. This implies that either the diagnosis of IBD might be preceded by microbial compositional changes or an increased risk of IBD is associated with an altered microbial composition or a combination of both.

In the present study, the relative abundances of taxa and pathways were not statistically significantly different between IBD-twins and their healthy cotwins. This is in contrast with previous, smaller-sized, 16S rRNA sequencing based microbiome IBD twin studies,\(^{11,13-15}\) in which gut microbiome differences between IBD-affected twins, especially individuals with ileal CD, and their healthy cotwins were reported.\(^{11,13}\) In these previous twin studies, furthermore, a microbial profile linked to ileal CD that differed from healthy cotwins and patients with colonic CD,\(^{11-12}\) and a less diverse microbial composition at the intestinal mucosa in UC-affected twins compared with their healthy cotwins was observed.\(^{14}\) This last phenomenon was found to be less pronounced in healthy cotwins, but was still decreased compared with healthy non-IBD related twins.\(^{14}\) A Belgian study in siblings and parents of CD-affected patients, using denaturing gradient gel electrophoresis, found a fecal microbiome dysbiosis in the unaffected relatives as compared with healthy controls characterized by mucin degrading microbiota.\(^{27}\) An English study among siblings of patients with CD also noted a dysbiosis in the fecal microbiome of the IBD-unaffected siblings.\(^{28}\) The results of these smaller family (ie, non-twin) studies based on techniques with lower resolution are in line with our findings with a changed composition of the fecal microbiome in healthy cotwins with IBD-affected twins.

The question arises whether shared genetics and (childhood) environment, rather than IBD phenotype or disease activity, might have resulted in a more similar composition of the gut microbiome between healthy cotwins and their IBD-twins. Previous work underscored the large impact of environmental factors during childhood on the adult gut microbiome composition.\(^{3}\) To further unravel the processes shaping the gut microbiome and explore the putative microbial drivers of IBD, we included unrelated patients with IBD, in addition to the IBD-twins. Unrelated patients with IBD showed only minor differences in individual taxa and pathways with IBD-twins. Interestingly, healthy cotwins did not only have overlap in differentially abundant species and pathways with their IBD-twins, but also with the unrelated patients with IBD compared with healthy controls. This renders shared environment and genetics as sole explanation for the overlap in microbiome signatures between IBD-twins and their healthy cotwins less probable, and suggests a mechanistic role for the gut microbiome in the development of IBD in the prediagnostic state whether or not related to subclinical inflammation, which we cannot rule out. Although prediagnostic microbiome data are lacking for IBD, in type 1 diabetes mellitus gut microbiome changes, albeit moderate, have been shown to occur before diagnosis,\(^{29}\) hinting toward the possibility of gut microbial changes occurring in the mechanistic chain of disease development.

One of the IBD signatures detected in healthy cotwins was an increase in the relative abundance of *Escherichia unclassified* (Figure 4A). *Escherichia unclassified* belongs to the genus of gram-negative, facultative anaerobic taxa that display *lipopolysaccharide*, a potent stimulator of innate immune responses, on their outer surface. *Escherichia unclassified* has previously been associated with UC and CD, and is considered a key pathogenic driver in IBD.\(^{30,31}\) Moreover, 2 pathways involved in the biosynthesis of family-specific surface enterobacterial common antigen, which is shared by all members of, and restricted to, the family Enterobacteriaceae,\(^{32}\) and a pathway for the biosynthesis of enterobactin, a siderophore that chelates Fe\(^{3+}\) and is known to be a virulence factor,\(^{33}\) were increased in the healthy cotwins, IBD-twins, and unrelated patients with IBD, as compared with the healthy individuals. The relative abundance of the genes encoding L-arginine degradation was increased in healthy cotwins, IBD-twins, and unrelated patients with IBD, as compared with the healthy controls, as well. L-Arginine is a precursor for the synthesis of polyamines. Polyamines contribute to the integrity of the gut and reduced expression of proinflammatory cytokines by monocytes and macrophages.\(^{32}\) L-arginine has been associated with a protective effect of colitis in mice models,\(^{33,34}\) and decreased levels of L-arginine have been observed in the intestinal epithelium in active UC possibly caused by decreased cellular uptake and increased consumption by nitric oxide synthase 2.\(^{32}\) A decreased L-arginine biosynthesis or increased L-arginine degradation has been reported previously in metagenomic shotgun sequencing
based microbiome studies in IBD.\textsuperscript{3,35} Likewise, an increase in the relative abundance of the gene encoding the degradation of propionate, an SCFA, was observed in healthy cotwins, IBD-twins and unrelated patients with IBD, as compared with healthy controls. SCFAs are an important energy source for enterocytes, and induce tolerogenic and anti-inflammatory enterocyte and T-cell phenotypes by multiple mechanisms.\textsuperscript{36,37} SCFAs constitute an increasingly convincing link between the gut microbiome and the IBD phenotype, and SCFA supplementation has been found to attenuate colonic inflammation.\textsuperscript{38} Interestingly, the relative abundance of \textit{F. prausnitzii} (often inversely associated with IBD\textsuperscript{3}), a well-known butyrate producer, and 2 butyrate biosynthesis pathways was found to be decreased in IBD-twins and unrelated patients with IBD, but not in healthy cotwins. We could thus not replicate the data from 1 Spanish\textsuperscript{39} and 1 English\textsuperscript{28} smaller-sized 16S rRNA quantitative polymerase chain reaction–based study, in which a decrease of \textit{F. prausnitzii} in relatives of patients with UC\textsuperscript{39} and relatives of patients with CD\textsuperscript{38} was found.

To date, insights in the preclinical phase of IBD are scarce.\textsuperscript{40} Previous epidemiological studies have shown an increased risk of developing IBD in healthy cotwins from IBD-discordant twin pairs.\textsuperscript{41} It is therefore tempting to speculate that changes in the gut microbiome of healthy cotwins precede IBD development and are involved in the pathogenesis. An alternative explanation might be that these microbiome alterations reflect the impact of shared genetic makeup or environmental factors in these individuals, but do not necessarily lead to IBD development. Longitudinal studies, with sampling at multiple timepoints in high-risk individuals before the onset of IBD (for example the GEM-project\textsuperscript{41}), are therefore needed to get a true insight in the preclinical phase of IBD.

Our study is, to the best of our knowledge, the largest twin study in the field of IBD and the microbiome up to date with high-resolution assessment of the gut microbiome. An important strength of our study included the comparison of carefully phenotyped IBD-discordant and -concordant twin pairs, and a large cohort of age-, sex-, and BMI-matched healthy controls and unrelated patients with IBD, allowing us to study several aspects of the overlap of the microbiome between healthy cotwins and patients with IBD. Although the participants came from 2 cohorts, the DNA isolation, library preparation, and sequencing were for all samples performed in the same way, at the same location and time by the same technician, minimizing the risk for batch effects. Furthermore, we adjusted our analyses for potential confounding factors, thereby reducing the chance of identifying spurious associations between the microbiome composition and IBD-status.

Our study does, however, have its limitations. Further increasing the sample size could have increased the power to detect more subtle differences in microbiome composition. Furthermore, multicollinearity prevented us from correcting for the use of IBD medication (which has been shown, however, to be only mildly associated with the gut microbiome composition\textsuperscript{42}), and stool consistency. Patients with CD and patients with UC were grouped together in our analyses. However, by including IBD phenotype as a covariate in our regression models we corrected for its potential confounding effects. Although we sought, as described, to minimize batch effects as much as possible, this could not completely be avoided, given the fact that the fecal samples were collected from individuals included in 2 different cohorts. Last, the unrelated patients with IBD were self-reported and, therefore, detailed information on Montreal classification, IBD-medication use, and disease activity was not available. Nonetheless, the gut microbiome composition of IBD-twins and unrelated patients with IBD considerably overlapped, as would have been expected when comparing 2 cohorts of patients with IBD.

In conclusion, we found that the gut microbiome of healthy cotwins from IBD-discordant twin pairs displays IBD-like signatures, both at a taxonomic and functional level. The gut microbiome of these individuals at increased risk of developing IBD displays similarities to the gut microbiome of their IBD-affected twins and unrelated patients with IBD, and is different from healthy controls. These IBD-like microbiome signatures could be a reflection of a shared genetic background and environment and might precede IBD development. However, longitudinal studies are needed to infer a causal relationship.

Supplementary Material

Note: To access the supplementary material accompanying this article, visit the online, version of \textit{Gastroenterology} at www.gastrojournal.org, and at https://doi.org/10.1053/j.gastro.2021.01.030.

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