Systematic Review

Systematic Review: The Gut Microbiome and Its Potential Clinical Application in Inflammatory Bowel Disease

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Abstract: Inflammatory bowel disease (IBD) is a chronic relapsing–remitting systemic disease of the gastrointestinal tract. It is well established that the gut microbiome has a profound impact on IBD pathogenesis. Our aim was to systematically review the literature on the IBD gut microbiome and its usefulness to provide microbiome-based biomarkers. A systematic search of the online bibliographic database PubMed from inception to August 2020 with screening in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines was conducted. One-hundred and forty-four papers were eligible for inclusion. There was a wide heterogeneity in microbiome analysis methods or experimental design. The IBD intestinal microbiome was generally characterized by reduced species richness and diversity, and lower temporal stability, while changes in the gut microbiome seemed to play a pivotal role in determining the onset of IBD. Multiple studies have identified certain microbial taxa that are enriched or depleted in IBD, including bacteria, fungi, viruses, and archaea. The two main features in this sense are the decrease in beneficial bacteria and the increase in pathogenic bacteria. Significant differences were also present between remission and relapse IBD status. Shifts in gut microbial community composition and abundance have proven to be valuable as diagnostic biomarkers. The gut microbiome plays a major role in IBD, yet studies need to go from casualty to causality. Longitudinal designs including newly diagnosed treatment-naïve patients are needed to provide insights into the role of microbes in the onset of intestinal inflammation. A better understanding of the human gut microbiome could provide innovative targets for diagnosis, prognosis, treatment and even cure of this relevant disease.

Keywords: gut microbiome; inflammatory bowel disease; Crohn’s disease; ulcerative colitis; biomarkers

1. Introduction

The gastrointestinal microbiota comprises a collection of microbial communities, including viruses, bacteria, archaea and fungi, inhabiting the gastrointestinal tract [1]. The constitution and diversity of the microbiota in different sections of the gastrointestinal tract are highly variable and its concentration increases steadily along it, with small numbers in the stomach and very high concentrations in the colon [2,3]. This community has been linked to many diseases, including inflammatory bowel disease (IBD) [4].

IBD encompasses a group of chronic inflammatory bowel pathologies of idiopathic origin that affect millions of people throughout the world; the two most important pathologies covered by this term are Crohn’s disease (CD) and ulcerative colitis (UC) [5]. IBD is not curable and shows a chronic evolution, with alternating periods of exacerbation and remission. This situation entails a high burden on health care systems, which try to provide treatment and to ensure quality of life for these complex patients who often require lifelong medical attention.
The microbiota of the gastrointestinal tract is frequently proposed as one of the key players in the etiopathogenesis of IBD. Studies in animal models and humans have shown that there is a persistent imbalance of the intestinal microbiome (which refers to the gut microbiota and their collective genetic material) related to IBD, with a substantial body of literature providing evidence for the relation of the human gut microbiome and IBD [4,6–10]. Despite all this evidence, it has been difficult to determine whether these changes in the microbiome are the cause of IBD or rather the result of inflammation after IBD onset. The consequence of this relationship between the human gut and microbes is that pharmacological therapies, diet and other interventions targeted to the host will also significantly impact the gut microbiota. Most of the existing studies attempting to determine whether dysbiosis is causative or a consequence of inflammation had certain limitations, such as disparities in methodologic approaches, including different techniques used to analyze the gut microbiome, different sampling sites (stool/mucosa) or site of inflammation, lack of prospective data, small cohort sizes, restricted focus on bacteria, different disease activities and influence of treatment interventions.

We conducted a systematic review to comprehensively collate the body of evidence surrounding the relationship between the gut microbiome and IBD. Our objective was to describe the associations between IBD and dysbiosis and the potential clinical translation of microbiome-based biomarkers.

2. Methodology

2.1. Search Strategy

An electronic search was conducted using the MEDLINE database via PubMed to identify published articles on the gut microbiome and IBD, from inception to August 2020. The search strings used were:

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[("ulcerative colitis" [MeSH Terms]) OR ("colitis" [All Fields] AND "ulcerative" [All Fields]) OR ("ulcerative colitis" [All Fields]) OR ("crohn disease" [MeSH Terms]) OR ("crohn" [All Fields] AND "disease" [All Fields]) OR ("crohn disease" [All Fields]) OR ("crohn’s disease" [All Fields]) OR ("inflammatory bowel diseases" [MeSH Terms]) OR ("inflammatory bowel diseases" [All Fields]) AND ("microbiome" [All Fields]) OR "microbiota" [All Fields]).
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Moreover, the reference lists of the included studies were revised to identify further relevant studies.

The work was conducted in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement in Appendix A [11].

2.2. Eligibility Criteria

The inclusion criteria were intestinal microbiome studies comparing IBD patients with controls; performed on fecal, intestinal lavage or intestinal tissue samples; focused on human adults; written in English.

Studies were excluded if they reported data on IBD complications or postsurgery (pouchitis, fistulae, among others); studied other conditions in addition to IBD (irritable bowel syndrome, Clostridium difficile infection, primary sclerosing cholangitis, among others); were abstracts from conference proceedings, letters to editor, reviews or reported only one patient.

3. Results

A total of 5267 records were identified from the PUBMED database. Of 190 papers remaining after screening, 23 did not include controls, 22 included other pathologies and 2 were in silico studies. A total of 143 papers were ultimately included.

3.1. Gut Microbiome Studies in IBD: Methodologic Aspects

The main methodologic characteristics of the studies included in this review are summarized in Table 1 (IBD gut microbiome studies using non-next-generation sequencing [NGS] approaches) and Table 2 (IBD gut microbiome studies using NGS approaches).
| Reference | Year | Treatment | No. of Participants | Disease State | Specimen | Histology | Design | Microbiome Analysis Method | Focus | Microbiota Findings |
|-----------|------|-----------|---------------------|---------------|----------|-----------|--------|----------------------------|-------|-----------------------|
| Macfarlane et al. [12] | 2004 | Not naïve | NA 9 NA 10 | Active | Biopsy | NA | Cross-sectional | Culture, FISH | Bacteria | UC | Only differences in bifidobacteria were statistically significant.  
* Peptostreptococci were only present in UC patients. |
| Lepage et al. [13] | 2005 | Not naïve | 20 11 NA 4 | Active/Inactive | Stool and biopsy | Non-inflamed | Cross-sectional | TTGE (16S rDNA V6–V8 region) | Bacteria | CD and UC | Dominant species differ between the mucosa-associated and fecal microbiota.  
* The microbiota is relatively stable along the distal digestive tract. |
| Manichanh et al. [14] | 2006 | Not naïve | 6 NA NA 6 | Inactive | Stool | NA | Cross-sectional | Cloning, Sequencing (16S rDNA) | Bacteria | CD | Reduced Firmicutes diversity. |
| Bibiloni et al. [15] | 2006 | Naïve | 20 15 NA 14 | Active | Biopsy | Inflamed/non-inflamed | Cross-sectional | DGGE (16S rDNA V3 region) and qPCR | Bacteria | CD and UC | Bacteria associated with inflamed and non-inflamed tissues did not differ.  
* UC had more bacteria associated with biopsies than CD.  
* Bacteroidetes were more prevalent in CD than in UC. |
| Sokol et al. [16] | 2006 | Not naïve | NA 9 NA 9 | Active | Stool | NA | Cross-sectional | TTGE (16S rDNA V6–V8 region) | Bacteria | UC | Reduced bacterial diversity. |
| Gophna et al. [17] | 2006 | Not naïve | 6 5 NA 5 | Active/Inactive | Biopsy | Inflamed/non-inflamed | Cross-sectional | PCR, cloning, sequencing (16S rDNA) | Bacteria | CD and UC | No significant difference between inflamed and non-inflamed tissues.  
* In CD, increased Proteobacteria and Bacteroidetes and reduced Clostridia.  
* No difference between UC and HC. |
| Scanlan et al. [18] | 2006 | Not naïve | 16 NA NA 6 | Active/Inactive | Stool | NA | Longitudinal | DGGE (16S rDNA) | Bacteria | CD | Lower temporal bacterial stability but higher stability for remission patients.  
* Lactic acid bacteria spp. varied significantly between the CD groups.  
* Decrease in Clostridium and Bacteroides spp. |
Table 1. Cont.

| Reference        | Year | Treatment     | Disease State | Microbiome Analysis Method | Focus | Microbiota Findings |
|------------------|------|---------------|---------------|----------------------------|-------|---------------------|
| Zhang et al.     | 2007 | Not naïve     | NA 24 NA 24   | DGGE (16S rDNA V3 region) | Bacteria | Lactobacilli and the *Clostridium leptum* subgroup were significantly different between the inflamed and non-inflamed tissues. They were also affected by UC location. |
| Sepehri et al.   | 2007 | Not naïve     | 10 15 NA 16   | ARISA, T-RFLP             | Bacteria | Differences between inflamed and non-inflamed tissues were found. The non-inflamed tissues form an intermediate population between HC and inflamed tissue for both CD and UC. |
| Andoh et al.     | 2007 | Not naïve     | NA 44 NA 44   | T-RFLP (16S rDNA)         | Bacteria | Bacterial communities are different between HC and active UC patients and between active and inactive patients. *Eubacterium*, and *Fusobacterium* were predominantly detected in the active patients. *Lactobacillus* were more predominant in the inactive patients. |
| Frank et al.     | 2007 | Not naïve     | 68 61 NA 61   | PCR, cloning, sequencing (16S rDNA) | Bacteria | Significant differences between the microbiotas of CD and UC and those of non-IBD controls. Depletion of members of the phyla Firmicutes and Bacteroidetes. |
| Ott et al.       | 2008 | Not naïve     | 13 NA 5 5     | PCR, cloning and sequencing | Bacteria | Temporal instability and bacterial richness decreased in relapsing patients compared to remission. |
| Ott et al.       | 2008 | Not naïve     | 31 26 NA 47   | DGGE, clone libraries, sequencing, in situ hybridization (18S rDNA) | Fungi | Increased fungal richness and diversity in CD. |
| Reference            | Year | Treatment | No. of Participants | Disease State | Specimen | Histology | Design       | Microbiome Analysis Method | Focus                  | Microbiota Findings                                                                                       |
|----------------------|------|-----------|---------------------|---------------|----------|-----------|--------------|---------------------------|------------------------|----------------------------------------------------------------------------------------------------------------|
| Martinez et al. [25] | 2008 | Not naïve | NA 16 NA            | Inactive      | Stool    | NA        | Longitudinal | DGGE (16S rDNA V6–V8 region) | Bacteria               | UC Temporal instability and reduced diversity in remission patients. |
| Dicksved et al. [26] | 2008 | Not naïve | 14 NA NA            | Active/Inactive| Stool    | NA        | Cross-sectional | T-RFLP, cloning and sequencing (16S rDNA) | Bacteria               | CD Decreased bacterial diversity. Decreased *Bacteroides uniformis* and increased *B. ovatus* and *B. vulgatus*. Ileal CD bacterial communities were significantly different from HC and colonic CD. |
| Kuehbacher et al. [27] | 2008 | Not naïve | 42 31 NA            | Active        | Biopsy   | Inflamed  | Cross-sectional | T-RFLP (16S rDNA) | Bacteria               | CD and UC TM7 (subgroup of Gram-positive uncultivable bacteria) were more diverse in CD than in UC and non-IBD controls. |
| Andoh et al. [28]   | 2008 | Not naïve | 34 NA NA            | Active/Inactive| Stool    | NA        | Cross-sectional | T-RFLP (16S rDNA) | Bacteria               | CD Decrease in *Clostridium* cluster IV, *Clostridium* cluster XI and subcluster XV. Increase in *Bacteroides* and Enterobacteriales. |
| Nishikawa et al. [29] | 2009 | Not naïve | 9 NA NA             | Active/Inactive| Biopsy   | Inflamed/Non-inflamed | Longitudinal | T-RFLP (16S rDNA) | Bacteria               | UC Decreased diversity due to loss of commensals. Decreased diversity in inactive patients compared to active patients. |
| Willing et al. [30]  | 2009 | Not naïve | 14 NA NA            | Active/Inactive| Biopsy   | Inflamed/Non-inflamed | Cross-sectional | T-RFLP, cloning and sequencing, qPCR (16S rDNA) | Bacteria               | CD Ileal CD had a lower abundance of *Faecalibacterium prausnitzii* and an increased abundance of *Escherichia coli* compared to healthy co-twins and colonic CD. Dysbiosis was significantly correlated to the disease phenotype. |
| Reference         | Year | Treatment | No. of Participants | Disease State | Specimen | Histology | Design       | Microbiome Analysis Method                  | Focus                                      | Microbiota Findings                                                                 |
|-------------------|------|-----------|---------------------|---------------|----------|-----------|--------------|---------------------------------------------|--------------------------------------------|------------------------------------------------------------------------------------------|
| Andoh et al.      | 2009 | Not naïve | NA 2 NA 3Ur         | Inactive      | Stool    | NA        | Cross-sectional | T-RFLP (16S rDNA)                      | Bacteria                                  | Increase in *Clostridium* cluster IX and decreases in *Clostridium* cluster XIVa.     |
| Gillevet et al.   | 2010 | Not naïve | 4 2 NA 4            | NA            | Stool and biopsy | NA        | Cross-sectional | LH-PCR, cloning, sequencing, and multitagged pyrosequencing (16S rDNA) | Bacteria                                  | CD and UC. Mucosal microbiome is distinct from the luminal microbiome in HC. Mucosal microbiome appears to be dysbiotic in IBD. |
| Rehman et al.     | 2010 | Not naïve | 10 10 NA 10         | Active        | Biopsy   | Inflamed  | Cross-sectional | PCR, cloning, sequencing (16S rDNA)      | Bacteria                                  | CD and UC. Increase in *Escherichia* sp.                                                |
| Kang et al.       | 2010 | Not naïve | 6 NA NA 6           | Inactive      | Stool    | NA        | Cross-sectional | Microarray (16S rDNA)                  | Bacteria                                  | CD. Decreased *Eubacterium rectale*, *B. fragilis* group, *B. vulgatus*, *Ruminococcus albus*, *R. callidus*, *R. bromii*, and *F. prausnitzii*. Increased *Enterococcus* sp., *Clostridium difficile*, *E. coli*, *Shigella flexneri*, and *Listeria* sp. |
| Rowan et al.      | 2010 | Not naïve | NA 20 NA 19         | Active/Inactive | Biopsy   | NA        | Cross-sectional | PCR, qPCR (16S rDNA)                  | Bacteria                                  | UC. Increase in *Desulfovibrio*, more marked in acute phase.                            |
| Andoh et al.      | 2011 | Not naïve | 31 31 NA 30         | Active/Inactive | Stool    | NA        | Cross-sectional | T-RFLP (16S rDNA V4-V9)                | Bacteria                                  | CD and UC. Decrease in the *Clostridium* family in active UC and inactive/active CD. Increase in *Bacteroides*. Inactive UC tended to be closer to that of HC. |
| Mondot et al.     | 2011 | Not naïve | 16 NA NA 16         | Active        | Stool    | NA        | Cross-sectional | qPCR, RT qPCR (16S rDNA)               | Bacteria                                  | CD. Decrease in *F. prausnitzii* *Ruminococcus* bromii, *Oscillo bacter valericigenes*, *Bifidobacterium* bifidum, and *E. rectale*. Increase in *E. coli* and *Enterococcus faecium*. More marked increase in *E. coli* in ileal CD. |
| Reference          | Year  | Treatment       | No. of Participants | Disease State | Specimen | Histology | Design         | Microbiome Analysis Method | Focus | Microbiota Findings                                                                 |
|--------------------|-------|-----------------|---------------------|---------------|----------|-----------|----------------|---------------------------|-------|------------------------------------------------------------------------------------|
| Joossens et al.    | 2011  | Not naïve       | 68                  | NA            | Stool    | NA        | Cross-sectional | DGGE (16S rDNA V3), qPCR | CD    | Decrease in Dialister invisus species of Clostridium cluster XIVa, F. prausnitzii and Bifidobacterium adolescentis. Increase in R. gnavus. |
| Lepage et al.      | 2011  | Not naïve       | NA                  | 8             | Biopsy   | NA        | Cross-sectional | PCR, cloning, sequencing (16S rDNA) | UC    | Decreased bacterial diversity. Increase in Actinobacteria and Proteobacteria. Healthy siblings from discordant twins had more bacteria from the Lachnospiraceae and Ruminococcaceae families than twins who were both healthy. |
| Benjamin et al.    | 2012  | Not naïve       | 103                 | NA            | Stool    | NA        | Cross-sectional | FISH (16S rDNA) | CD    | Increase in Bacteroides-Prevotella in smokers (38.4%) compared with nonsmokers (28.1%). Increase in bifidobacterial and Bacteroides-Prevotella. Decrease in F. prausnitzii. |
| Hotte et al.       | 2012  | Not naïve       | 15                  | 14            | Biopsy   | Non-inflamed | Cross-sectional | T-RFLP (16S rDNA) | CD and UC | Increase in Proteobacteria compared with HC and UC. |
| Pistone et al.     | 2012  | Not naïve       | 35                  | 18            | Biopsy   | Inflamed/non-inflamed | Cross-sectional | PCR M. avium subspecies paratuberculosis | CD and UC | Increase in M. avium subspecies paratuberculosis compared to controls. |
| Andoh et al.       | 2012  | Not naïve       | 67                  | NA            | Stool    | NA        | Longitudinal    | T-RFLP (16S rDNA V1-V9) | CD    | Decrease in Clostridia in active disease and remission and in Bifidobacterium in active phase but increased during remission. Increase in Bacteroides genus in active. Decreased bacterial diversity. |
Table 1. Cont.

| Reference | Year  | Treatment       | No. of Participants | Disease State | Specimen | Histology              | Design         | Microbiome Analysis Method          | Focus                  | Microbiota Findings                                                                 |
|-----------|-------|-----------------|---------------------|---------------|----------|------------------------|----------------|-------------------------------------|------------------------|---------------------------------------------------------------------------------------|
| Li et al. [44] | 2012  | Not naïve       | 18 NA NA 9          | Active        | Stool and biopsy        | Inflamed/non-inflamed | Cross-sectional | DGGE (16S rDNA V3 region), sequencing | Bacteria               | Decreased bacterial diversity. Increase in γ-Proteobacteria (especially E. coli and S. flexneri). Decrease in reduced Bacteroidetes and Firmicutes. In ulcerated mucosa, E. coli was increased and F. prausnitzii, Lactobacterium coliforms, Bacteroides sp and Streptococcus gallolyticus were decreased compared with the non-ulcerated. |
| Nemoto et al. [45] | 2012  | Not naïve       | NA 48 NA 36         | Active/Inactive| Stool     | NA                     | Cross-sectional | Culture, T-RFLP, qPCR              | Bacteria               | Decreased bacterial diversity. Decrease in Bacteroides and Clostridium subcluster XI and Enterococcus. Increase in Enterococcus. |
| Vignaes et al. [46] | 2012  | Not naïve       | NA 12 NA 6          | Active/Inactive| Stool     | NA                     | Cross-sectional | DGGE (16S rDNA, 16S-23S rDNA intergenic spacer region), qPCR | Bacteria               | Different microbiota in active UC compared to HC but in inactive UC compared to HC. Decrease in Lactobacillus spp. and Akkermansia muciniphila in active disease. |
| de Souza et al. [47] | 2012  | Not naïve       | 11 7 NA 14          | NA            | Stool and biopsy        | Inflamed/non-inflamed | Cross-sectional | Culture                           | E. coli                | CD and UC Only the mucosa-associated population of E. coli was increased, not in stool. The increase was prominent in the ileal CD and rectum and sigmoid of both UC and CD. |
| Duboc et al. [48] | 2013  | Not naïve       | 12 30 NA 26         | Active/Inactive| Stool     | NA                     | Cross-sectional | PCR (rDNA), culture            | Bacteria               | CD and UC Decrease in the ratio between F. prausnitzii and E. coli |
| Sha et al. [49] | 2013  | Not naïve       | 10 26 NA 14         | Active/Inactive| Stool     | NA                     | Cross-sectional | DGGE (16S rDNA V6-V8 region), qPCR | Bacteria               | CD and UC Decrease in the numbers of Bacteroides–Porphyromonas–Prevotella, Bifidobacterium and B. fragilis in active phase. Decrease in Helicobacter and Clostridium phylogenetic clusters XI and XIVa in active and inactive phases. Increase in E. coli in active phases. |
### Table 1. Cont.

| Reference          | Year | Treatment | No. of Participants | Disease State | Specimen | Histology | Design           | Microbiome Analysis Method | Focus                        | Microbiota Findings                                      |
|--------------------|------|-----------|---------------------|---------------|-----------|-----------|------------------|-----------------------------|-----------------------------|----------------------------------------------------------|
| Kabeerdoss et al. [50] | 2013 | Not naïve | 20 22 NA 17        | Active/Inactive | Stool   | NA        | Cross-sectional | TTGE (16S rDNA V1–V9), qPCR | C. leptum group, F. prausnitzii | CD and UC                                                                 |
|                    |      |           |                     |               |          |           |                  |                             | Decrease in C. leptum group bacteria and F. prausnitzii. | Decreased bacterial diversity. |
| Varela et al. [51]  | 2013 | Not naïve | NA 116 NA 29 Ur + 31 | Inactive       | Stool   | NA        | Cross-sectional and longitudinal | PCR (16S rDNA), qPCR | F. prausnitzii | UC                                                                 |
|                    |      |           |                     |               |          |           |                  |                             | Decrease in F. prausnitzii in patients and relatives. | Recovery of the F. prausnitzii population after relapse was associated with remission. |
| Midtvedt et al. [52] | 2013 | Not naïve | 4 NA NA 5          | Active         | Stool and biopsy | Inflamed | Cross-sectional | Microarray          | Bacteria                   | CD                                                                 |
|                    |      |           |                     |               |          |           |                  |                             | Decrease in Bacteroides in both stool and biopsies. |
| Fujimoto et al. [53] | 2013 | Not naïve | 47 NA NA 20        | Active/Inactive | Stool   | NA        | Cross-sectional | qPCR, PCR (16S rDNA V4–V9), T-RFLP | F. prausnitzii and Bilophila wadsworthia | CD                                                                 |
|                    |      |           |                     |               |          |           |                  |                             | Decrease in Clostridia, including the genus Faecalibacterium. | Decreased bacterial diversity. |
| Fite et al. [54]   | 2013 | Not naïve | NA 33 NA 18        | Active         | Biopsy  | Inflamed | Longitudinal    | qPCR                         | Bacteria                   | UC                                                                 |
|                    |      |           |                     |               |          |           |                  |                             | High clinical activity indices were associated with enterobacteria, desulfovibrios, type E Clostridium perfringens, and Enterococcus fecalis. | Decreased bacterial diversity. |
| Rajilic-Stojanovic et al. [55] | 2013 | Not naïve | NA 15 NA 15        | Inactive       | Stool   | NA        | Longitudinal    | Microarray                  | Bacteria                   | UC                                                                 |
|                    |      |           |                     |               |          |           |                  |                             | Decrease in members of the Clostridium cluster IV R. bromii et rel. E. rectale et rel., Roseburia sp., and Akkermansia sp. | Decrease in C. coccoides and C. leptum clusters. |
| Kumari et al. [56] | 2013 | Not naïve | NA 26 NA 14        | Active/Inactive | Stool   | NA        | Cross-sectional | FISH, flow cytometry, qPCR (16S rDNA) | Bacteria                   | UC                                                                 |
|                    |      |           |                     |               |          |           |                  |                             | F. prausnitzii and Roseburia intestinales were differentially present in patients with different disease activity. |
Table 1. Cont.

| Reference          | Year | Treatment     | No. of Participants | Disease State | Specimen | Histology | Design  | Microbiome Analysis Method | Focus | Microbiota Findings                                                                                     |
|--------------------|------|---------------|---------------------|---------------|----------|-----------|---------|------------------------------|-------|----------------------------------------------------------------------------------------------------------------|
| Hedin et al. [57]  | 2014 | Not naïve     | 22                  | NA            | NA       | 25 + 21Ur | Inactive | Stool                        | NA    | Cross-sectional qPCR (16S rDNA) Bacteria CD - Siblings shared dysbiosis pattern with patients (lower concentrations of F. prausnitzii, Clostridia cluster IV and Roseburia spp.). |
| Lennon et al. [58] | 2014 | Not naïve     | NA                  | NA            | 19       | 34        | Active  | Biopsy                       | NA    | Cross-sectional qPCR (16S rDNA) Desulfovibrio species UC - No significant differences in Desulfovibrio sp. were found between cohorts or at each sampling region between the cohorts. |
| Machiels et al. [59]| 2014 | Not naïve     | NA                  | 127           | NA       | 447       | Active/Inactive | Stool                        | NA   | Cross-sectional PCR (16S rDNA V3 region) DGGE, sequencing, qPCR Bacteria UC - Decrease in Roseburia hominis and F. prausnitzii showed an inverse correlation with disease activity. |
| Wang et al. [60]   | 2014 | Not naïve     | 21                  | 34            | NA       | 21        | Active/Inactive | Stool and biopsy                | Inflamed/non-inflamed | Cross-sectional qPCR (16S rDNA) Bacteria CD and UC - Bifidobacterium was increased in biopsies of active UC patients, and higher in the biopsies than in the fecal samples in active CD patients. - *Lactobacillus* group was increased in biopsies of active CD patients. - *F. prausnitzii* was decreased in both the fecal and biopsy specimens of the active patients. |
| Blais Lecours et al. [61] | 2014 | Not naïve    | 18                  | 11            | NA       | 29        | Active/Inactive | Stool                        | NA   | Cross-sectional qPCR Archaea and bacteria CD and UC - Increase in Methanosphaera stadtmanae. |
| Fukuda et al. [62] | 2014 | Not naïve     | NA                  | 69            | NA       | 80Ur      | Active/Inactive | Stool                        | NA   | Cross-sectional PCR (16S rDNA, V4-V9 region), T-RFLP Bacteria UC - Development of a Discriminant Score based on selected OTUs. - Five differential clusters were obtained indicating a strong association between the gut microbiota and UC. |
Table 1. Cont.

| Reference        | Year | Treatment                   | No. of Participants | Disease State | Specimen | Histology          | Design          | Microbiome Analysis Method | Focus          | Microbiota Findings                                                                 |
|------------------|------|-----------------------------|---------------------|---------------|----------|---------------------|-----------------|----------------------------|----------------|--------------------------------------------------------------------------------------|
| Li et al. [63]   | 2014 | Not naïve                   | 19                  | NA            | NA       | Active              | Stool and biopsy| Inflamed/non-inflamed     | Cross-sectional| DGGE (18S rDNA), cloning, sequencing        | Fungi          | Increase in fungal richness and diversity in the inflamed mucosa compared with the non-inflamed mucosa.  
|                  |      |                             |                     |               |          |                     |                 |               |                                             |                | Increase in Candida spp., Gibberella moniliformis, Alternaria brassicicola, and Cryptococcus neoformans.  
|                  |      |                             |                     |               |          |                     |                 |               |                                             |                | In stool, increase in fungal diversity and prevalence in Candida albicans, Aspergillus clavatus, and C. neoformans.  
| Andoh et al. [64]| 2014 | Not naïve                   | 160                 | NA            | NA       | Active/Inactive     | Stool           | NA            | Longitudinal              | T-RFLP (16S rDNA V1–V9) | Bacteria            | CD  
|                  |      |                             |                     |               |          |                     |                 |               |                                             |                | Decision tree based on selected OTUs, obtaining 9 groups.  
|                  |      |                             |                     |               |          |                     |                 |               |                                             |                | Microbiota profiles may differ according to disease activity.  
| Wisittipanit et al. [65] | 2015 | Not naïve                   | 101                 | 89            | NA       | Active/Inactive     | Biopsy and lumen aspiration | NA           | Cross-sectional | LH-PCR (16S rDNA V1–V2 region) | Bacteria | Development of a computational pipeline to characterise the gut microbial communities.  
|                  |      |                             |                     |               |          |                     |                 |               |                                             |                | Model could classify IBD from HC at specific locations and based on disease state *  
| Kabeerdoss et al. [66] | 2015 | Naïve and not naïve         | 28                  | 32            | NA       | Biopsy              | Inflamed/non-inflamed | Cross-sectional | RT-qPCR (16S rDNA) | Bacteria | CD and UC  
|                  |      |                             |                     |               |          |                     |                 |               |                                             |                | Increase in Bacteroides and Lactobacillus in UC patients compared with controls or CD.  
|                  |      |                             |                     |               |          |                     |                 |               |                                             |                | Increase in E. coli in UC compared with controls.  
|                  |      |                             |                     |               |          |                     |                 |               |                                             |                | Decrease in C. coccoides group and C. leptum group in CD compared with controls.  
|                  |      |                             |                     |               |          |                     |                 |               |                                             |                | Decrease in Firmicutes to Bacteroidetes ratio in UC and CD.  
|                  |      |                             |                     |               |          |                     |                 |               |                                             |                | No differences between inflamed and non-inflamed tissues were found, nor between treated and untreated patients.  

| Reference                  | Year | Treatment          | No. of Participants | Disease State | Specimen | Histology | Design          | Microbiome Analysis Method | Focus | Microbiota Findings                                                                 |
|----------------------------|------|---------------------|---------------------|---------------|----------|-----------|-----------------|----------------------------|-------|-------------------------------------------------------------------------------------|
| Takeshita et al. [67]      | 2016 | Not naïve           | NA 48 NA 34         | Active/Inactive | Stool    | NA        | Cross-sectional | RT-qPCR                   | Bacteria | Decrease bacterial diversity in active phase. Fusicatenibacter saccharivorans was decreased in active patients and increased in quiescent. |
| Zhang et al. [68]          | 2017 | Not naïve           | 132 NA NA 71        | Active/Inactive | Stool    | NA        | Cross-sectional | Culture                   | Bacteria | CD Increase in E. coli and Enterococcus spp. in active phase compared with inactive and controls. |
| Vrakas et al. [69]         | 2017 | Naïve and not naïve | 12 20 NA 20         | Active/Inactive | Biopsy   | Inflamed  | Cross-sectional | RT-qPCR (16S rDNA)         | Bacteria | CD and UC Increased total bacterial DNA concentration levels in active phase compared to the inactive. Increase in Bacteroides spp. in active and inactive phases. Decrease in C. leptum group (IV), and F. prausnitzii in active and inactive phases. |
| Zamani et al. [70]         | 2017 | Not naïve           | NA 35 NA 60         | Active        | Biopsy   | Inflamed  | Cross-sectional | Culture, qPCR               | Bacteria | UC No association between B. fragilis and UC. Enterotoxigenic B. fragilis was more prevalent in UC patients with diarrhoea. |
| Ghavami et al. [71]        | 2018 | Not naïve           | 9 45 NA 47          | Active/Inactive | Stool    | NA        | Cross-sectional | PCR, qPCR (16S rDNA)               | Bacteria and Methanobrevibacter smithii (Archaea) | CD and UC Decrease in Methanobrevibacter smithii. More marked increase in Mbb. smithii in remission than in active phase. |
| Le Baut et al. [72]        | 2018 | Not naïve           | 262 NA NA 76        | Resected tissue and biopsy | Inflamed/non-inflamed | Cross-sectional | PCR | Yersinia Species | CD | Increase in Yersinia species. |
| Al-Bayati et al. [73]      | 2018 | Not naïve           | NA 40 NA 40         | Biopsy        | Inflamed  | Cross-sectional | Culture, PCR (16S rDNA) | Bacteria | UC Decrease in F. prausnitzii, Prevotella, and Peptostreptococcus productus. |
Table 1. Cont.

| Reference                | Year | Treatment                  | No. of Participants | Disease State | Specimen | Histology | Design       | Microbiome Analysis Method | Focus       | Microbiota Findings                                                                 |
|--------------------------|------|----------------------------|---------------------|---------------|----------|-----------|--------------|----------------------------|-------------|------------------------------------------------------------------------------------|
| Heidarian et al. [74]    | 2019 | Not naïve                  | 7 22 NA 29          | Active/Inactive | Stool    | NA        | Cross-sectional | qPCR | Bacteria                                                                 |
|                          |      |                            |                     |               |          |           |              |                             | CD and UC   | • Decrease in Bacteroides, F. prausnitzii, Prevotella spp., and Methanobrevibacterium. |
|                          |      |                            |                     |               |          |           |              |                             |             | • Decrease in Bacteroides spp., F. prausnitzii, and Prevotella spp. in UC patients with disease activity score greater than 4. |
|                          |      |                            |                     |               |          |           |              |                             |             | • Increase in Streptococcus and Haemophilus in the patients who were at flare.       |
| Vatn et al. [75]         | 2020 | Naïve and not naïve        | 68 84 12 160        | Active/Inactive | Stool    | NA        | Cross-sectional | GA-map™ (16S rDNA V3-V9 region) | Bacteria    | CD and UC                                                                 |
|                          |      |                            |                     |               |          |           |              |                             |             | • Decrease in Firmicutes and Eubacterium hallii.                                    |
|                          |      |                            |                     |               |          |           |              |                             |             | • Increase in Bifidobacterium spp., E. hallii, Actinobacteria and Firmicutes in ulcerative proctitis, compared to extensive colitis. |
|                          |      |                            |                     |               |          |           |              |                             |             | • No association with disease location in CD.                                       |

Abbreviations: CD, Crohn’s disease; UC, ulcerative colitis; IBD, inflammatory bowel disease; IBDU, inflammatory bowel disease unclassified; HC, healthy control; C, control; NA, not applicable; FISH, fluorescence in situ hybridization; TTGE, temporal temperature gradient gel electrophoresis; DGGE, denaturing gradient gel electrophoresis; qPCR, quantitative real-time polymerase chain reaction; ARISA, automated ribosomal intergenic spacer analysis; T-RFLP, terminal restriction fragment length polymorphism; Ur, unaffected relatives; LH-PCR, length heterogeneity polymerase chain reaction; OTUs, operational taxonomic unit. * No microorganisms specified.

Table 2. Gut microbiome studies in inflammatory bowel disease using next-generation sequencing approaches.

| Reference        | Year | Treatment | No. of Participants | Disease State | Specimen | Histology | Design          | Microbiome Analysis Method | Focus       | Microbiota Findings                                                                 |
|------------------|------|-----------|---------------------|---------------|----------|-----------|-----------------|----------------------------|-------------|------------------------------------------------------------------------------------|
| Willing et al.   | 2010 | Not naïve | 29 16 NA 35         | Active/Inactive | Stool and biopsy | Non-inflamed | Cross-sectional | 16S rDNA sequencing          | Bacteria    | CD and UC                                                                 |
|                  |      |           |                     |               |          |           |                 |                            |             | • Ileal CD differed from colonic CD.                                                |
|                  |      |           |                     |               |          |           |                 |                            |             | • In ileal CD, decrease in Faecalibacterium and Roseburia, and increase in Enterobacteriaceae and Ruminococcus gravis. |
| Rausch et al.    | 2011 | Not naïve | 29 NA NA 18         | Inactive      | Biopsy   | Non-inflamed | Cross-sectional | 16S rDNA V1–V2 region sequencing | Bacteria    | CD                                                                 |
|                  |      |           |                     |               |          |           |                 |                            |             | • Decrease in bacterial diversity.                                                  |
|                  |      |           |                     |               |          |           |                 |                            |             | • Prevotella, Lactobacillus, Coprococcus, Clostridium, Faecalibacterium, and Stenotrophomonas were only present in HC. |
Table 2. Cont.

| Reference          | Year | Treatment         | No. of Participants | Disease State | Specimen | Histology         | Design   | Microbiome Analysis Method | Focus                                      |
|--------------------|------|-------------------|---------------------|---------------|----------|-------------------|----------|----------------------------|--------------------------------------------|
| Walker et al. [78] | 2011 | Not naïve         | 6, 6, NA            | Active        | Biopsy   | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V1–V8 region sequencing | Bacteria                                    |
| Erickson et al. [79]| 2012 | Not naïve         | 8, NA, NA           | Active/Inactive| Stool    | NA                | Cross-sectional | 16S rDNA V1–V2 region and WGS | Bacteria                                    |
| Morgan et al. [80] | 2012 | Not naïve         | 121, 75, 8         | Active/Inactive| Stool and biopsy | NA            | Cross-sectional | 16S rDNA V3–V5 region and WGS | Bacteria                                    |
| Ricanek et al. [81]| 2012 | Naïve             | 4, NA, NA           | Active        | Biopsy   | Inflamed           | Cross-sectional | 16S rDNA sequencing            | Bacteria                                    |
| Thorkildsen et al. [84]| 2013 | Naïve             | 30, 33, 3         | Active        | Stool    | NA                | Cross-sectional | 16S rDNA (all regions) sequencing | Bacteria                                    |

CD and UC
- Decrease in bacterial diversity.
- Decrease in Firmicutes and increase in Bacteroidetes, and in CD only, Enterobacteriaceae.
- Differences between inflamed and non-inflamed tissues were found.

CD and UC
- Decrease in bacterial diversity.
- Decrease in Firmicutes in ileal CD.

CD and UC
- Disease status influenced Firmicutes and Enterobacteriaceae abundances.

CD and UC
- Microbiota of Norwegian CD patients was found to be similar to that of CD patients in other countries.

CD and UC
- Decrease in C. coccoides-E. rectales group in ileal CD compared to control non-IBD.
- Decrease in F. prausnitzii in CD.

CD and UC
- Decrease in Firmicutes and increase in Proteobacteria and Actinobacteria.
- Decrease in microbial diversity.

CD and UC
- Increase in Escherichia/Shigella in CD.
- Decrease in Faecalibacterium in CD compared to both UC and controls.
| Reference          | Year   | Treatment | No. of Participants | Disease State | Specimen | Histology                    | Design          | Microbiome Analysis Method      | Focus                                                                 | Microbiota Findings                                                                                                                                                                                                 |
|--------------------|--------|-----------|---------------------|---------------|----------|----------------------------|----------------|-------------------------------|---------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Prideaux et al. [85] | 2013   | Not naïve | 22, 30, NA          | CD, UC        | Biopsy   | Inflamed/non-inflamed       | Cross-sectional | Microarray, 16S rDNA V1–V3 region sequencing | Bacteria                                                                 | CD and UC<br>• Decrease in microbial diversity and in Faecalibacterium, Coprococcus, Dorea, Roseburia, and 2 unclassified genera (from Lachnospiraceae and Clostridiales) in CD.<br>• In UC, diversity was reduced in Chinese subjects.<br>• Actinobacteria was significantly different between the UC groups.<br>• Decrease in Coprococcus and Dorea genera in UC. |
| Chiodini et al. [86] | 2013   | Not naïve | 14, NA              | NA            | Resected tissue | NA                          | Cross-sectional | 16S rDNA V3–V6 region sequencing | Bacteria                                                                 | CD<br>• Separation of the submucosal and mucosal microbiome and existence of a submucosal bacterial population within diseased tissues. |
| Pérez-Brocal et al. [87] | 2013   | Naïve and not naïve | 11, NA, NA          | CD           | Stool    | NA                          | Cross-sectional | Viral DNA and 16S rDNA V1–V3 region sequencing | Bacteria and viruses                                                                 | CD<br>• Decreased bacterial and viral diversity.<br>• Synchroococcus phage S CBS1 and Retroviridae family viruses were more represented in CD.<br>• Increase in Proteobacteria and decrease in Tenericutes, the order Bacteroidales and Collinsella aerofaciens. |
| Davenport et al. [88] | 2014   | Not naïve | 13, 14, NA          | CD, UC        | Biopsy   | Inflamed                   | Cross-sectional | 16S rDNA V4 region sequencing | Bacteria                                                                 | CD and UC<br>• Decreased bacterial diversity.<br>• No phylum-level significant differences in Firmicutes or Proteobacteria<br>• Bacteroidetes were only increased in CD. |
Table 2. Cont.

| Reference       | Year | Treatment | No. of Participants | Disease State | Specimen         | Histology                          | Design        | Microbiome Analysis Method | Focus                      |
|-----------------|------|-----------|---------------------|---------------|------------------|------------------------------------|---------------|---------------------------|----------------------------|
| Chen et al.     | 2014 | Not naïve | 26 41 NA 21         | Active/Inactive | Stool and biopsy | Inflamed/non-inflamed              | Cross-sectional | 16S rDNA V1–V3 region sequencing | Bacteria CD and UC          |
|                 |      |           |                     |               |                  |                                    |               |                           | - Decrease in Roseburia, Coprococcus, and Ruminococcus. |
|                 |      |           |                     |               |                  |                                    |               |                           | - Increase in Escherichia-Shigella and Enterococcus. |
|                 |      |           |                     |               |                  |                                    |               |                           | - Fecal- and mucosa-associated microbiota were similar between CD and UC and differed from HC. |
| Wang et al.     | 2015 | Not naïve | 6 4 NA 5            | NA            | Biopsy           | NA                                 | Cross-sectional | RNA sequencing             | Bacteria and viruses CD and UC |
|                 |      |           |                     |               |                  |                                    |               |                           | - Increase in bradyrhizobiaceae, enterobacteriaceae, comamonadaceae, and moraxellaceae families. |
|                 |      |           |                     |               |                  |                                    |               |                           | - Human adenovirus and Herpesviridae sequences were predominant in IBD. |
| Lavelle et al.  | 2015 | Not naïve | NA 9 NA 4           | NA            | Luminal brush, mucosal biopsy, mucus gel layer | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V4 region sequencing | Bacteria UC |
|                 |      |           |                     |               |                  |                                    |               |                           | - Spatial variation between the luminal and mucosal communities in both cohorts. |
|                 |      |           |                     |               |                  |                                    |               |                           | - Decrease in Bacteroidaceae and Akkermanniaceae. |
|                 |      |           |                     |               |                  |                                    |               |                           | - Increase in Clostridiaceae, Peptostreptococcaceae, Enterobacteriaceae, Ruminococcaceae, Bifidobacteriaceae, Actinomyctaeae and FJ440089, an uncultured member of the Prevotellaceae family. |
| Chiordini et al. | 2015 | Not naïve | 20 NA NA 15         | NA            | Biopsy           | Inflamed                           | Cross-sectional | 16S rDNA V4 region sequencing | Bacteria CD |
|                 |      |           |                     |               |                  |                                    |               |                           | - Distinct sub-mucosal microbiome compared to mucosa and/or fecal material. |
|                 |      |           |                     |               |                  |                                    |               |                           | - Desulfovibrionales were present within the submucosal tissues. |
|                 |      |           |                     |               |                  |                                    |               |                           | - Increase in Firmicutes in the subjacent submucosa as compared to the parallel mucosal tissue. |
|                 |      |           |                     |               |                  |                                    |               |                           | - Increase in Propionibacterium spp., Cloacibacterium spp., Parasutterella spp. and Methylobacterium spp. |
| Reference          | Year | Treatment               | No. of Participants | Disease State | Specimen | Histology          | Design       | Microbiome Analysis Method | Focus                  | Microbiota Findings                                                                 |
|--------------------|------|-------------------------|---------------------|---------------|----------|--------------------|--------------|-----------------------------|------------------------|-------------------------------------------------------------------------------------|
| Pérez-Brocal et al. [93] | 2015 | Naïve and not naïve     | 20 NA NA 20        | Active        | Stool and biopsy | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V1–V3 region and viral DNA/RNA sequencing | Bacteria and viruses | Decrease bacterial diversity in all CD groups. Increased richness and diversity were observed in feces compared with biopsies. Increase in Actinobacteria, Gammaproteobacteria, and Fusobacteria. |
| Vidal et al. [94]   | 2015 | Not naïve               | 13 NA NA 7         | Active/Inactive| Biopsy   | Non-inflamed       | Cross-sectional | 16S rDNA V1–V5 region sequencing | Bacteria              | Decrease in Clostridia and increase in Bacteroidetes and Proteobacteria. No detection of F. prausnitzii. |
| Norman et al. [95]   | 2015 | Not naïve               | 18 42 NA 12        | Active/Inactive| Stool    | NA                 | Cross-sectional | VLP DNA sequencing | Viruses                | CD and UC Increase in Caudovirales bacteriophages. It did not appear that expansion and diversification of the enteric virome was secondary to changes in the microbiota. |
| Eun et al. [96]     | 2016 | Not naïve               | 35 NA NA 15        | Inactive      | Stool and biopsy | NA                | Cross-sectional | 16S rDNA V1–V3 region sequencing | Bacteria              | Decrease in bacterial diversity. Increase in Proteobacteria was increased in both fecal and mucosal tissues, and Fusobacteria only in tissue samples. Increase in Gammaproteobacteria and Fusobacteria in both fecal and mucosal tissue samples in active phase. |
| Chiodini et al. [97] | 2016 | Not naïve               | 20 NA NA 15        | NA            | Biopsy   | Inflamed           | Cross-sectional | 16S rDNA V1–V3 region sequencing | Bacteria              | CD Increase in Sphingomonadaceae, Alicyclobacillaceae, Methylbacteriaceae, Pseudomonadaceae and Prevotellaceae in the submucosa at the advancing disease margin when compared to the superjacent mucosa (translocation). |
| Reference          | Year | Treatment       | No. of Participants | Disease State | Specimen | Histology | Design       | Microbiome Analysis Method        | Focus                                                                 |
|--------------------|------|-----------------|---------------------|---------------|----------|-----------|--------------|------------------------------------|----------------------------------------------------------------------|
| Rehman et al. [98] | 2016 | Not naïve       | 28                  | Inactive      | Biopsy   | NA        | Cross-sectional | 16S rDNA V1–V2 region sequencing | Bacteria                                                             |
|                    |      |                 | 30                  |               |          |           |              |                                    | CD and UC                                                            |
|                    |      |                 | NA                  |               |          |           |              |                                    | • Proteobacteria decrease in UC compared with CD and HC.               |
|                    |      |                 | 30                  |               |          |           |              |                                    | • Different microbial pattern based on geographical origin.          |
| Takahashi et al. [99] | 2016 | Not naïve       | 68                  | Active/Inactive| Stool    | NA        | Cross-sectional | qPCR and 16S rDNA V3–V4 region sequencing | Bacteria                                                             |
|                    |      |                 | NA                  |               |          |           |              |                                    | CD and UC                                                            |
|                    |      |                 | NA                  |               |          |           |              |                                    | • Increase in Actinomyces and Bifidobacterium.                        |
| Forbes et al. [100] | 2016 | Not naïve       | 15                  | NA            | Biopsy   | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V6 region sequencing | Bacteria                                                             |
|                    |      |                 | 21                  |               |          |           |              |                                    | CD and UC                                                            |
|                    |      |                 | NA                  |               |          |           |              |                                    | • No difference between inflamed and non-inflamed tissues were found.|
|                    |      |                 | 7                   |               |          |           |              |                                    | • There were only differences between the inflamed and non-inflamed  |
|                    |      |                 |                     |               |          |           |              |                                    | • mucosa between CD and UC.                                          |
|                    |      |                 |                     |               |          |           |              |                                    | • Increase in Bacteroides and Fusobacteria in inflamed CD mucosa      |
|                    |      |                 |                     |               |          |           |              |                                    | • than in inflamed UC mucosa.                                       |
|                    |      |                 |                     |               |          |           |              |                                    | • Proteobacteria and Firmicutes were more frequently in the inflamed  |
|                    |      |                 |                     |               |          |           |              |                                    | • UC mucosa.                                                         |
| Liguori et al. [101] | 2016 | Not naïve       | 23                  | Active/Inactive| Biopsy   | Inflamed/non-inflamed | Cross-sectional | qPCR (16S or 18S rDNA) 16S rDNA V3–V4 region and ITS2 sequencing | Bacteria and fungi                                                   |
|                    |      |                 | NA                  |               |          |           |              |                                    | CD and UC                                                            |
|                    |      |                 | NA                  |               |          |           |              |                                    | • Decrease in bacterial diversity.                                 |
|                    |      |                 | 10                  |               |          |           |              |                                    | • Increase in Proteobacteria and Fusobacteria.                       |
|                    |      |                 |                     |               |          |           |              |                                    | • Increase in fungal load in active phase.                           |
|                    |      |                 |                     |               |          |           |              |                                    | • Cystofilobasidiaceae family and Candida glabrata species were       |
|                    |      |                 |                     |               |          |           |              |                                    | • overrepresented.                                                   |
| Mar et al. [102]   | 2016 | Not naïve       | NA                  | NA            | Stool    | NA        | Cross-sectional | 16S rDNA V3–V4 region and ITS2 sequencing | Bacteria and fungi                                                   |
|                    |      |                 | 30                  |               |          |           |              |                                    | UC                                                                  |
|                    |      |                 | NA                  |               |          |           |              |                                    | • Decrease in bacterial diversity.                                 |
|                    |      |                 | 13                  |               |          |           |              |                                    | • Decrease in Bacteroides and Prevotella species and Alternaria       |
|                    |      |                 |                     |               |          |           |              |                                    | • alternata, Aspergillus flavus,                                     |
|                    |      |                 |                     |               |          |           |              |                                    | • Aspergillus cibarius, and Candida soja.                            |
|                    |      |                 |                     |               |          |           |              |                                    | • Increase in Streptococcus, Bifidobacterium, and Enterococcus        |
|                    |      |                 |                     |               |          |           |              |                                    | • and Candida albicans and Debaryomyces.                            |
Table 2. Cont.

| Reference          | Year | Treatment | No. of Participants | Disease State | Specimen | Histology | Design        | Microbiome Analysis Method | Focus            | Microbiota Findings                                                                 |
|--------------------|------|-----------|---------------------|---------------|----------|-----------|---------------|----------------------------|------------------|----------------------------------------------------------------------------------|
| Hoarau et al. [103]| 2016 | Not naive | 20                  | NA            | NA       | NA        | Cross-sectional | 16S rDNA V4 region and ITS1 sequencing | Bacteria and fungi | CD Increase in Serratia marcescens and E. coli, and Candida tropicalis. |
| Hedin et al. [104] | 2016 | Not naive | 21                  | NA            | NA       | NA        | Cross-sectional | 16S rDNA V1–V3 region sequencing | Bacteria | CD Decrease in bacterial diversity. Decrease in *F. prausnitzii*. |
| Naftali et al. [105]| 2016 | Not naive | 31                  | NA            | NA       | NA        | Cross-sectional | 16S rDNA V1–V3 region sequencing | Bacteria | CD Difference between ileal CD compared with colonic CD. This separation was unaffected by the biopsy’s location, its inflammatory state or disease state. *Faecalibacterium* was strongly reduced in ileal CD compared with colonic CD, whereas *Enterobacteriaceae* were more abundant in the former. |
| Pedamallu et al. [106]| 2016 | Not naive | 12                  | NA            | NA       | NA        | Cross-sectional | WGS | Bacteria | CD Decrease in Bacteroidetes and Clostridiales. Enrichment of enterotoxigenic *Staphylococcus aureus* and an environmental *Mycobacterium* species within deeper layers of the ileum. |
| Sokol et al. [107] | 2016 | Not naive | 149                 | 86            | NA       | 38 Active/Inactive | Stool | Cross-sectional | 16S rDNA V3–V5 region and ITS2 sequencing | Bacteria and fungi | CD and UC Increase in Basidiomycota/Ascomycota ratio and *C. albicans*. Decrease in *Saccharomyces cerevisiae*. Correlations between bacterial and fungal components. |
| Santorl et al. [108]| 2017 | Not naive | 50                  | 82            | NA       | 51 Active/Inactive | Stool | Cross-sectional | 16S rDNA V3–V4 region sequencing, qPCR | Bacteria | CD and UC Increase in Firmicutes, Proteobacteria, Verrucomicrobia, and Fusobacteria. Decrease in Bacteroidetes and Cyanobacteria. |
Table 2. Cont.

| Reference         | Year | Treatment | No. of Participants | Disease State | Specimen | Histology | Design | Microbiome Analysis Method | Focus                     | Microbiota Findings                                                                 |
|-------------------|------|-----------|---------------------|---------------|----------|-----------|--------|----------------------------|---------------------------|-------------------------------------------------------------------------------------|
| Pascal et al. [109] | 2017 | Not naïve | 34 33 NA 40 + 71Ur | Active/Inactive | Stool    | NA        | Longitudinal | 16S rDNA V4 region sequencing | Bacteria                  | CD and UC: Dysbiosis was greater in CD than UC, as shown by a more reduced diversity, a less stable microbial community. |
| Chen et al. [110]  | 2017 | Not naïve | NA 8 NA 8 NA       | NA            | Stool    | NA        | Cross-sectional | 16S rDNA V3–V4 region sequencing | Bacteria                  | UC: Decrease in Firmicutes, (Blautia, Clostridium, Coprococcus and Roseburia). Decreased bacterial diversity. |
| Hall et al. [111]  | 2017 | Not naïve | 9 10 1 12           | Active/Inactive | Stool    | NA        | Longitudinal | WGS           | Bacteria                  | CD and UC: Increase in facultative anaerobes. Increase in R. genus, often co-occurring with increased disease activity. |
| Qiu et al. [112]   | 2017 | Not naïve | NA 14 NA 15          | Active        | Biopsy   | Inflamed  | Cross-sectional | 18S rDNA sequencing | Fungi                     | UC: Increase in Wickerhamomyces, unidentified genus of Saccharomycotales, Aspergillus, Sterigmatomyces, and Candida. Decrease in Exophiala, Alternaria, Emersiella, Epicoccum, Acremonium, Trametes, and Penicillium. |
| Kennedy et al. [113] | 2018 | Not naïve | 37 NA NA 54          | Inactive      | Stool    | NA        | Cross-sectional | 16S rDNA V1–V2 region sequencing | Bacteria                  | CD: Decrease in bacterial diversity. Decrease in Ruminococcaceae, Rikenellaceae, and Christensenellaceae. Increase in Enterobacteriaceae. |
| Ji et al. [114]    | 2018 | Not naïve | 51 66 NA 66          | Active/Inactive | Stool    | NA        | Cross-sectional | 16S rDNA V4 region sequencing | Bacteria                  | CD and UC: Results were different between HC and IBD patients and between active and inactive patients. |
| Imbann et al. [115] | 2018 | Not naïve | 188 107 18 582       | Active/Inactive | Stool    | NA        | Cross-sectional | 16S rDNA V4 region sequencing | Bacteria                  | CD and UC: Colonic CD was different from that of patients with ileal CD, with a decrease in alpha diversity associated with ileal CD. Decrease in the genus Roseburia was associated with higher IBD risk score. |
Table 2. Cont.

| Reference        | Year  | Treatment                      | No. of Participants | Disease State | Specimen | Histology   | Design   | Microbiome Analysis Method | Focus | Microbiota Findings                                                                 |
|------------------|-------|--------------------------------|---------------------|---------------|----------|-------------|----------|----------------------------|-------|-----------------------------------------------------------------------------------|
| Nishino et al.   | 2018  | Not naïve                      | 26, 43 NA 14        | Active/Inactive | Macosal brush | Non-inflamed | Cross-sectional | 16S rDNA V3–V4 region sequencing | Bacteria | CD and UC:  
|                  |       |                                |                     |               |          |             |          |                            |       | - No significant difference among anatomical sites within individuals.  
|                  |       |                                |                     |               |          |             |          |                            |       | - Increase in Proteobacteria and decrease in Firmicutes in CD and Bacteroidetes in CD.  
|                  |       |                                |                     |               |          |             |          |                            |       | - Greater abundance of Escherichia, Ruminococcus (R. gnavus), Clostridium, Cetobacterium, Peptostreptococcus in CD, and the Faecalibacterium, Blautia, Bifidobacterium, Roseburia and Citrobacter in UC.  |
| Rojas-Feria et al. | 2018  | Naïve                          | 13 NA NA 16         | Onset         | Stool    | NA          | Cross-sectional | 16S rDNA V1–V3 region sequencing | Bacteria | CD:  
|                  |       |                                |                     |               |          |             |          |                            |       | - Decrease in bacterial diversity.  
|                  |       |                                |                     |               |          |             |          |                            |       | - Decrease in Firmicutes and an increase in Bacteroidetes.  |
| Schirmer et al.  | 2018  | Naïve and not naïve            | 30, 21 NA 11        | Active/Inactive | Stool    | NA          | Longitudinal| WGS           | Bacteria | CD and UC:  
|                  |       |                                |                     |               |          |             |          |                            |       | - Decrease in bacterial diversity.  
|                  |       |                                |                     |               |          |             |          |                            |       | - Decrease in Firmicutes and increase in Enterobacteriaceae.  
|                  |       |                                |                     |               |          |             |          |                            |       | - Longitudinal profiles showed taxonomic shifts in community composition over time that coincided with changes in disease severity.  |
| Chiodini et al.  | 2018  | Not naïve                      | 20 NA NA 15         | Resected tissue | Inflamed  | Cross-sectional | 16S rDNA V1–V3 region sequencing | Bacteria | CD:  
|                  |       |                                |                     |               |          |             |          |                            |       | - Increase in bacterial richness.  
|                  |       |                                |                     |               |          |             |          |                            |       | - Bacterial translocation, with two bacterial families (Comamonadaceae and Xanthomonadaceae), having penetrated the mucosal surfaces.  |
| Hirano et al.    | 2018  | Not naïve                      | NA 14 NA 14         | Active         | Biopsy   | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V4 region sequencing | Bacteria | UC:  
|                  |       |                                |                     |               |          |             |          |                            |       | - Decrease in bacterial diversity.  
|                  |       |                                |                     |               |          |             |          |                            |       | - Increase in Clostridium and the Tissierellaceae and decrease in Neisseria in inflamed site when compared to the non-inflamed site.  |
| Reference                  | Year | Treatment         | No. of Participants | Disease State | Specimen | Histology | Design       | Microbiome Analysis Method | Focus | Microbiota Findings                                                                 |
|----------------------------|------|-------------------|---------------------|---------------|----------|-----------|--------------|-----------------------------|-------|-------------------------------------------------------------------------------------|
| Ma et al. [121]            | 2018 | Not naïve         | 15                  | Active/Inactive | Stool    | NA        | Cross-sectional | 16S rDNA V4 region sequencing | Bacteria | Increase in Proteobacteria. Decrease in Bacteroidetes in the active CD compared to inactive CD. Bacteroidetes showed a negative correlation with the CD activity index scores. |
| Walujkar et al. [122]      | 2018 | Not naïve         | NA                  | Active         | Biopsy   | Inflamed  | Longitudinal  | 16S rDNA V4 region sequencing | Bacteria | UC Increase in bacterial count in active UC. Increase in *Stenotrophomonas, Parabacteroides, Elizabethkingia, Pseudomonas, Micrococcus, Ochrobactrum and Achromobacter* in active UC. |
| Moen et al. [123]          | 2018 | Naïve             | NA                  | Onset          | Biopsy   | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V4 region sequencing | Bacteria | UC No difference in bacterial diversity. Proteobacteria were higher in the inflamed tissue compared with the non-inflamed. |
| Laserna-Mendieta et al. [124] | 2018 | Not naïve         | 71                  | Active/Inactive | Stool    | NA        | Cross-sectional | 16S rDNA V3–V4 region sequencing | Bacteria | CD and UC Decreased bacterial diversity. Decreased in *Clostridium* cluster IV, *Roseburia*, and *F. prausnitzii* only in CD. |
| Libertucci et al. [127]    | 2018 | Not naïve         | 43                  | Active/Inactive | Biopsy   | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V3 region and ITS2 sequencing | Bacteria and fungi | CD Increase in *Escherichia* and a decrease in *Firmicutes* in inflamed tissue. Bacterial diversity did not correlate with inflammation. |
| Moustafa et al. [126]      | 2018 | Not naïve         | 45                  | Active/Inactive | Stool    | NA        | Cross-sectional | WGS                          | Bacteria | CD and UC Decreased bacterial diversity. Increase in *Proteobacteria* and decrease in Bacteroidetes and Firmicutes. |
| O’Brien et al. [127]       | 2018 | Not naïve         | 24                  | NA            | Biopsy   | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V1–V3 region sequencing | Bacteria | CD No bacterial imbalance or reduced diversity in CD aphthous ulcers and adjacent mucosa, relative to control biopsies. |
| Reference          | Year | Treatment | No. of Participants | Disease State | Specimen | Histology          | Design   | Microbiome Analysis Method | Focus                                                                 | Microbiota Findings                                      |
|--------------------|------|-----------|---------------------|---------------|----------|--------------------|----------|----------------------------|-----------------------------------------------------------------------|----------------------------------------------------------|
| Zakrzewski et al. [128] | 2019 | Not naïve | 15                  | NA            | NA       | 58                 | Active   | Inflamed/non-inflamed      | Cross-sectional | 16S rDNA V3–V4 region sequencing | Bacteria                                                                 | Decrease in bacterial diversity and richness.            |
|                    |      |           |                     |               |          |                    |          |                            | VLP and 16S rDNA sequencing |                                                                                                        | Decrease in F. prausnitzii.                                |
| Zuo et al. [129]   | 2019 | Not naïve | NA                  | 91            | NA       | 76                 | Active/Inactive | Biopsy        | Inflamed/non-inflamed      | Cross-sectional |                                                                                                        | Viruses                                                                                 |
|                    |      |           |                     |               |          |                    |          |                            | VLP and 16S rDNA sequencing |                                                                                                        | Increase in Caudovirales bacteriophages, but decrease in mucosa Caudovirales diversity, richness and evenness. |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | Virome correlated with intestinal inflammation.                                  |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | Increase in Escherichia phage and Entrobacteria phage.                      |
| Altomare et al. [130] | 2019 | Not naïve | 10                  | NA            | NA       | 11                 | Active/Inactive | Stool and biopsy | Inflamed/non-inflamed      | Cross-sectional | 16S rDNA V1–V3 region sequencing | Bacteria                                                                 | Fecal microbiota was more similar to controls than mucosal microbiota. |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | In the colon district some specific bacterial biomarkers were identified: Entrobacteriaceae for IBD stools, Bacteroides for IBD biopsies. |
| Franzosa et al. [131] | 2019 | Not naïve | 68                  | 53            | NA       | 34                 | Active/Inactive | Stool | NA                        | Cross-sectional | WGS                        | Bacteria                                                                 | CD and UC                                                                 |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | Decreased bacterial diversity.                                              |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | Decrease in Firmicutes and increase in Proteobacteria.                      |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | Disease localization did not have a significant effect among CD subjects.   |
| Lloyd-Price et al. [132] | 2019 | Not naïve | 67                  | 38            | NA       | 27                 | Active/Inactive | Stool and biopsy | Inflamed/non-inflamed      | Longitudinal | 16S rDNA sequencing and WGS | Bacteria and viruses |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | CD and UC                                                                 |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | Increase in facultative anaerobes at the expense of obligate anaerobes.     |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | Periods of disease activity were marked by increases in temporal taxonomic variability. |
| Imai et al. [133]  | 2019 | Not naïve | 20                  | 18            | NA       | 20                 | Inactive | Stool | NA                        | Cross-sectional | 16S rDNA V3–V4 region and ITS sequencing | Bacteria and fungi |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | CD and UC                                                                 |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | Decrease in bacterial diversity in CD compared to HC and UC.               |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | No difference in fungal diversity.                                        |
|                    |      |           |                     |               |          |                    |          |                            |                                                                                                        | Increase in Candida in CD compared to HC and UC.                            |
### Table 2. Cont.

| Reference          | Year | Treatment | No. of Participants | Disease State | Specimen | Histology            | Design       | Microbiome Analysis Method                      | Focus       | Microbiota Findings                                                                 |
|--------------------|------|-----------|---------------------|---------------|----------|----------------------|-------------|-----------------------------------------------|-------------|-------------------------------------------------------------------------------------|
| Li et al. [134]    | 2019 | Not naïve | 106                 | NA            | 88       | 89                   | Resected tissue | Inflamed/non-inflamed | Longitudinal | 16S rDNA V3–V5 region sequencing, qPCR | Bacteria    | CD and UC: Proteobacteria was positively associated with ileal CD and more marked in non-inflamed tissue. |
| Vester-Andersen et al. [135] | 2019 | Not naïve | 58                  | 82            | NA       | 30                   | Active/Inactive | Stool             | Cross-sectional | 16S rDNA V3–V4 region sequencing | Bacteria    | CD and UC: Decrease in richness, diversity and Firmicutes in active and in aggressive CD. Increase in Proteobacteria in CD. |
| Clooney et al. [136] | 2019 | Not naïve | 27                  | 82            | NA       | 61                   | Active/Inactive | Stool             | NA           | Whole-virome analysis and 16S rDNA V3–V4 region sequencing | Bacteria and viruses | CD and UC: No changes in viral richness. Increase in Caudovirales. Changes in virome reflected alterations bacteriome. |
| Braun et al. [137] | 2019 | Not naïve | 45                  | NA            | NA       | 22                   | Inactive       | Stool             | NA           | 16S rDNA V4 region sequencing | Bacteria    | CD: Decrease in bacterial diversity. Inactive patients preceding flare showed a decrease in Christensenellaceae and S24.7, and increase in Gemellaceae compared with those in remission. |
| Galazzo et al. [138] | 2019 | Not naïve | 57                  | NA            | NA       | 15                   | Active/Inactive | Stool             | NA           | 16S rDNA V4 region sequencing | Bacteria    | CD: Decrease in bacterial diversity and richness. Microbial community structure was less stable over time. |
| Sun et al. [139]   | 2019 | Not naïve | NA                  | 58            | NA       | 30                   | Active/Inactive | Stool             | Cross-sectional | 16S rDNA V3–V4 region sequencing | Bacteria    | UC: Decreased bacterial diversity. Firmicutes and Bacteroidetes, were the most abundant active UC and inactive UC, respectively. Increase in Proteobacteria and Fusobacteria and decrease in Firmicutes and Bacteroidetes in active UC. |
| Yilmaz et al. [140] | 2019 | Not naïve | 270                 | 232           | NA       | 573                  | Active/Inactive | Biopsy            | Inflamed/non-inflamed | Longitudinal | 16S rDNA V5–V6 region sequencing | Bacteria    | CD and UC: Decrease in diversity in CD compared with UC and HC. Firmicutes were higher than Bacteroidetes in UC compared with CD. |
Table 2. Cont.

| Reference          | Year | Treatment | No. of Participants | Disease State | Specimen | Histology | Design          | Microbiome Analysis Method | Focus | Microbiota Findings                                                                 |
|--------------------|------|-----------|---------------------|---------------|----------|-----------|-----------------|------------------------------|-------|----------------------------------------------------------------------------------|
| Magro et al. [141] | 2019 | Not naïve | 18 NA NA 18         | Inactive      | Stool    | NA        | Cross-sectional | 16S rDNA V3–V4 region sequencing | Bacteria | Decrease in bacterial diversity. Increase in Proteobacteria and decrease in Deltaproteobacteria, Akkermansia, Oscillospira and Saccharomyces cerevisiae. |
| Zhang et al. [142] | 2019 | Not naïve | NA 63 NA 30         | Active/Inactive | Stool    | NA        | Cross-sectional | 16S rDNA V4 region sequencing | Bacteria | Decrease in Porphyromonadaceae, Rikeneliaceae, and Lachnospiraceae and increase in Enterococcus and Streptococcus. |
| Alam et al. [143]  | 2020 | Not naïve | 9 11 NA 10          | NA            | Stool    | NA        | Cross-sectional | 16S rDNA V1–V3 region sequencing | Bacteria | Decrease in bacterial diversity. Increase in Porphyromonadaceae, Rikeneliaceae, and Lachnospiraceae and increase in Enterococcus and Streptococcus. |
| Ryan et al. [144]  | 2020 | Not naïve | 80 50 NA 31         | Active/Inactive | Biopsy   | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V3–V4 region sequencing | Bacteria | Difference in inflamed and non-inflamed colonic segments in both CD and UC. Inflammatory status did not appear to affect diversity. |
| Butera et al. [145]| 2020 | No naïve  | NA 88 NA 24         | Active        | Biopsy   | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V3–V4 region sequencing | Bacteria | High IL-13mRNA patients are younger at diagnosis and show higher prevalence of extensive colitis than low IL-13mRNA patients. Increase in Prevotellaceae in patients with high IL-13mRNA tissue content and Sutterella and Acidaminococcus in patients with low IL-13mRNA tissue content. |
| Boland et al. [146]| 2020 | No naïve  | 101 99 15 48        | Active/Inactive | Biopsy   | NA        | Cross-sectional | 16S rDNA V4 region sequencing | Bacteria | CD mucosal biopsy who achieved mucosal healing had lower diversity than biopsies from patients with UC or HC. Diversity was differently related to mucosal healing in CD and UC. |
| Reference | Year | Treatment | No. of Participants | Disease State | Specimen | Histology | Design | Microbiome Analysis Method | Focus | Microbiota Findings |
|-----------|------|-----------|---------------------|---------------|----------|-----------|--------|-----------------------------|-------|------------------------|
| Olaisen et al. [147] | 2020 | No naïve | 51 | NA | NA | 40 | Active/Inactive | Biopsy | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V3–V4 region sequencing | Bacteria | CD |
| | | | | | | | | | | | Decreased bacterial diversity. |
| | | | | | | | | | | Overrepresentation of Tyzzerella 4. |
| | | | | | | | | | | No difference in diversity in inflamed and non-inflamed ileal mucosa. |
| Shahir et al. [148] | 2020 | No naïve | 125 | NA | NA | 23 | NA | Biopsy | Inflamed/non-inflamed | Cross-sectional | 16S rDNA V1–V2 region sequencing | Bacteria | CD |
| | | | | | | | | | | Decreased bacterial diversity. |
| | | | | | | | | | | Distinct profile in colon and ileum. |
| | | | | | | | | | | Increase in obligate anaerobes in the ileum, B. fragilis was dramatically increased. |
| Park et al. [149] | 2020 | No naïve | 370 | NA | NA | 740 | Active/Inactive | Stool | NA | Longitudinal | 16S rDNA V3–V4 region sequencing | Bacteria | CD |
| | | | | | | | | | | Diversity was more decreased in patients with worse prognosis. |
| | | | | | | | | | | E. coli might be causally involved in CD progression. |
| Clooney et al. [150] | 2020 | No naïve | 303 | 228 | NA | 161 | Active/Inactive | Stool | NA | Longitudinal | 16S rDNA V3–V4 region sequencing | Bacteria | CD and UC |
| | | | | | | | | | | Decrease in bacterial diversity but increase in variability. |
| | | | | | | | | | | Reduced temporal microbiota stability, particularly in patients with changes in disease activity. |
| Park et al. [151] | 2020 | No naïve | 10 | 6 | NA | 9Ur | Inactive | Stool | NA | Cross-sectional | 16S rDNA V3–V4 region sequencing | Bacteria | CD and UC |
| | | | | | | | | | | Decrease in bacterial diversity. |
| | | | | | | | | | | Different diversity and identification of differentially abundant taxa in affected IBD relatives. |
| Lo Sasso et al. [152] | 2020 | No naïve | 41 | 43 | NA | 42 | Active | Stool | NA | Cross-sectional | 16S rDNA V4 region and WGS | Bacteria | CD and UC |
| | | | | | | | | | | Increase in Proteobacteria, Actinobacteria, and Fusobacteria. |
| | | | | | | | | | | Decrease in Firmicutes, Bacteroidetes, and Verrucomicrobia. |
| Reference          | Year | Treatment | No. of Participants | Disease State | Specimen | Histology | Design   | Microbiome Analysis Method | Focus                                                                                       | Microbiota Findings                                                                 |
|-------------------|------|-----------|---------------------|---------------|----------|-----------|----------|----------------------------|---------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------|
| Borren et al.     | 2020 | No naïve  | 108                 | NA            | NA       | Inactive  | Stool    | NA            | Longitudinal WGS           | CD and UC                                                                                 | Increase in Proteobacteria and Fusobacteria and, at the species level, *Lachnospiraceae bacterium_2_1_56FAA* in relapse. Potential microbial biomarker to identify proinflammatory state in quiescent IBD that predisposes to clinical relapse. |
| Rubbens et al.    | 2020 | No naïve  | 29                  | NA            | NA       | Inactive  | Stool    | NA            | Flow cytometry and 16S rDNA sequencing     | CD                                                                                | Decrease in bacterial diversity. Potential of flow cytometry to perform rapid diagnostics of microbiome profile. |

Abbreviations: CD, Crohn’s disease; UC, ulcerative colitis; IBD, inflammatory bowel disease; IBDU, inflammatory bowel disease unclassified; HC, healthy control; C, control; NA, not applicable; Ur, unaffected relatives; WGS, whole-genome shotgun sequencing; qPCR, quantitative real-time polymerase chain reaction; VLP, virus-like particle; ITS, internal transcribed spacer.
3.1.1. Study Design

Across the included studies, populations ranged from 2 to 531 patients, many of them with a small sample size that reduced the precision of the estimations. Thus, since many results are limited by sample size, further studies with larger cohorts are desirable to confirm these results and to clarify the significance of the microbiome in the pathogenesis of IBD.

In addition, to date, most published studies in IBD are cross-sectional (121 out of the 143 reviewed studies). However, longitudinal designs are required to capture changes that precede or coincide with disease and symptom onset, and to mechanistically relate microbiome shifts with disease pathogenesis. Overall, longitudinal studies in IBD (only 15% of the included studies) indicate that there is decreased stability in the microbiota composition in UC and CD patients [18,23,25,118,131,132,138,149]. These dynamic changes emphasize the importance of longitudinal sampling for a better understanding of taxa stability in individuals.

The IBD microbiome varies not only over time but also with treatment [80,155,156]. Newly diagnosed patients with no treatment provide an ideal scenario to study the potential etiopathogenesis related to intestinal dysbiosis that occurs in IBD. Mouse and human studies have proven that the gut microbiome is required for disease onset, as germ-free mice rarely develop the disease [157,158], antibiotics can prevent disease onset in mice [159] and ameliorate (but not cure) the disease in humans [160].

However, prior IBD microbiome studies have mostly included subjects with an established treatment; of the 143 microbiome studies included herein, only 11 included treatment-naïve patients [15,66,69,75,81,84,87,93,117,118,123], sometimes only on a small subset of the cohort, and only one was conducted prospectively.

Results on newly diagnosed treatment-naïve patients showed that gut dysbiosis is already established at the beginning of the disease. The dysbiotic profile in the gut of newly diagnosed treatment-naïve IBD patients presents reduced microbial abundance, less biodiversity in the structure of microbial communities, and differential bacterial abundances compared to the profile of established and treated IBD patients or control groups. Conversely, one study showed none or minor microbial differences between these patients and a control group [84].

Current knowledge, despite some controversy, provides valuable insights supporting the idea that microbial alterations may precede IBD onset. Given the limited number of studies in this type of patients, no consistent conclusion can be inferred, and further work is needed to investigate in depth the gut dysbiosis of newly diagnosed treatment-naïve IBD patients.

3.1.2. Microbiome Analysis Methods

Culture-independent and -dependent methods for microbial community analysis have both been used to describe microorganisms from different environments, including the human gut. However, due to the inability to culture the majority of the resident bacteria from the gastrointestinal tract, culture-independent methods have proven much more reliable and faster in profiling complex microbial communities.

Culture-independent techniques are based on sequence divergences of the small subunit ribosomal RNA (16S rRNA) or other target gene regions. Some of these techniques are quantitative real-time PCR (qPCR), denaturing gradient gel electrophoresis (DGGE), terminal restriction fragment length polymorphism (T-RFLP), fluorescence in situ hybridization (FISH), DNA microarrays, and NGS. All these techniques, except for NGS, are referred herein as non-NGS techniques.

Currently, there are many differences in study design and methodology among studies, making translation of basic science results into clinical practice a challenging task. Among the studies included in this review, very few used culture-dependent techniques (7 out of 143); and over the years, NGS became the most employed technique—79 studies used NGS, while 64 studies used non-NGS approaches.
Lately, the most widely used approaches are amplicon gene sequencing, predominantly the 16S rRNA gene (16S rDNA), and whole-genome shotgun sequencing, both NGS techniques.

Another recent technique much less used in this field is flow cytometry. A recent study demonstrated that cytometry fingerprints can be used as a diagnostic tool to classify samples according to CD state [154]. These results highlight the potential of flow cytometry as a tool to conduct rapid and cheaper diagnostics of microbiome-associated diseases.

3.1.3. Sample Type and Site

Currently, bacterial diversity in the human gut is determined through analysis of the luminal content (stool) and mucosal biopsies; however, the stool microbiome differs from the mucosa-associated microbiome [161]. Most of the bacteria are tightly adhered to the mucus and this mucosa-adhered microbiota may be associated with the pathogenesis of the disease [9,76]. Changes observed in stool samples likely represent an indirect measure of what is happening at the mucosal surface, where microorganisms interact more intimately with the host and induce disease.

The studies reviewed herein used fecal data, biopsy data or both, and most of them showed differences between fecal and biopsy samples [13,32,47,80,89,93,96,130], although a few studies found similarities [52,76]. The reported differences in microbial composition related to whether the sample origin was fecal or mucosal indicate that each biological sample represents a different environment thus emphasizing the importance of experimental design. Biopsies are primarily recommended for the dissection of the complex pathogenesis of IBD, whereas feces could effectively detect key biomarkers, enabling non-invasive continuous disease monitoring.

In biopsy samples, sampling site can also be a confounding factor. Many studies have compared the microbiome of inflamed and non-inflamed tissue from the same IBD patient. Regarding the effect of gut inflammation on the microbiota, there are some discrepancies among studies. Some researchers did not find significant differences in the mucosa-associated bacteria between apparently normal and inflamed mucosa in IBD patients [15,66,100,127,128,147]. Conversely, other studies found gut microbiome differences between inflamed and non-inflamed regions in mucosal biopsies [19,44,72,78,120,125,134,144]. This difference was also observed in fungal communities of inflamed mucosa, which are distinguishable from those of the non-inflamed area [63].

In spite of the controversial results, there is evidence supporting that inflamed and non-inflamed tissue samples in both CD and UC may present some differential microbiota composition suggesting that a comparison of mucosal samples obtained from identical sites in IBD patients and non-IBD controls is needed to avoid the confounding effect of inflammation in the assessment of the microbial profile.

3.1.4. Structural and Functional Analysis

IBD microbiome studies have typically focused on characterizing the composition of a community and less attention has been paid to functional profiles of the microbes within a community. Functional information can be inferred from the taxa through bioinformatic approaches or directly assessed via whole-genome shotgun sequencing.

Function is more informative than taxonomy [162] as it provides information on possible mechanisms acting on microbes and on microbe–host interactions, which are important for understanding microbial communities, specially microbiome-related diseases. The loss of a particular function could be more biologically meaningful than the loss of a single or a group of species.

The vast majority of the studies published on the IBD microbiome to date have focused on taxonomy and the reported associations in the IBD gut microbiome are largely limited to identifying high-level taxonomic classification (ranging from phyla to genera) given, for example, the limitations of amplicon gene sequencing for reliable species identification.
Some IBD gut microbiome studies have assessed the change in microbial function compared to healthy subjects. Outcomes of such studies showed a quite distinct change in microbial functions, such as fecal tryptic activity, oxidative response or lipid and glycan metabolism pathways [52,80,83,88,132]. Based on these results, it is necessary to redirect the study of dysbiosis from a purely compositional definition to a definition that includes functional changes of the microbiota.

3.2. Dysbiosis in IBD

The microbiome is different among healthy individuals around the globe [163], and the great differences found between the microbiomes of apparently healthy people complicate the definition of a healthy microbiome. Despite this divergence, the vast and diverse microbial gut community lives in relative balance in healthy individuals. Dysbiosis refers to an imbalance in microbial species, which is commonly associated with impaired gut barrier function and inflammatory activity [164]. It encompasses major traits such as loss of beneficial microbes, expansion of pathobionts, and loss of diversity [3] (Figure 1). The following sections will describe the key alterations found in the gut of IBD patients.

![Gut microbiota disturbance in inflammatory bowel disease compared to healthy individuals. Upward arrow indicates increase and downward arrow decrease.](image)

3.2.1. Defining the Gut Microbiome in IBD

Although the gastrointestinal tract contains trillions of resident microorganisms that include bacteria, archaea, fungi and viruses, the studies revised herein highlighted that current research on microbiome is mainly focused on bacteria.

Bacterial Dysbiosis

It has consistently been shown that there is a disease-dependent restriction of biodiversity and an imbalanced bacterial composition associated with IBD. The abundance of beneficial microorganisms such as Clostridium groups IV and XIVa, Bacteroides, Suterella, Roseburia, Bifidobacterium species and Faecalibacterium prausnitzii is reduced, whereas some pathogens such as Proteobacteria members (including invasive and adherent Escherichia coli), Veil-
*lonellaceae, Pasteurellaceae, Fusobacterium* species, and *Ruminococcus gnavus* are increased [4]. Most of the studies have revealed that in IBD patients, commensal bacteria are depleted and the microbial community is less diverse [14,22,37,48,80,94,106,108,126,143,150,152,153].

The increase in the phylum Proteobacteria, which includes multiple genera considered potentially pathogenic such as *Escherichia, Salmonella, Yersinia, Desulfovibrio, Helicobacter* or *Vibrio*, has been extensively reported in IBD patients [17,22,34,35,58,76,113,116,126,135,165].

In the *Firmicutes* phylum, *F. prausnitzii*, an anti-inflammatory commensal bacterium, is frequently decreased in CD, while less evidence has been reported in UC, where it is sometimes increased and in other studies decreased [14,22,51,59,73,89,109,140,142,166,167]. Specific decrease in *Roseburia* spp. in patients with IBD has also been consistently noted [56,59,76,99,115,116,130]. Both bacteria are known to be involved in the production of butyrate, an important energy source for intestinal epithelial cells, which strengthens gut barrier function and exerts important immunomodulatory functions [168]. In this same phylum, the mucin degrader *R. gnavus* is frequently increased in IBD patients’ gut, which may impair barrier stability and contribute to inflammation [38,76,103,111,116,132,140,144].

**Fungal Dysbiosis**

Despite the large body of literature on the IBD gut bacterial microbiome, little has been published on the gut mycobiome; specifically, only nine studies reviewed herein included fungal analysis.

Fungi are ubiquitous and their presence in the gastrointestinal tract has been demonstrated [169]. It was already evidenced many years ago that antibodies directed against mannanproteins of *Saccharomyces cerevisiae* (ASCA) were associated with CD, suggesting an inappropriate immune response to fungi in these patients [170].

Although fungi only constitute approximately 0.1% of the total microbial community in the gut [171], changes in gut mycobiota have been reported in IBD patients. However, results on fungal diversity are controversial; compared to controls, some studies have shown that fungal diversity is decreased in UC patients [107,112], and in CD, diversity and richness have been reported to be either increased [24,63], reduced [103,133], or unchanged [101]. Findings across fungal studies have consistently shown an increase in fungal load, especially in *Candida albicans* [24,63,101,102,107,133].

Nowadays, the exact mechanisms of intestinal fungi in IBD remain unclear and microbiome studies need to include fungi to properly address the complex challenges of this promising field.

**Viral Dysbiosis**

The human gut virome includes a diverse collection of viruses, mostly bacteriophages, directly impacting on human health [172]. In this systematic review, only seven studies included viral analysis [87,90,93,95,129,132,136]. Alterations in IBD gut virome showed an expansion of *Caudovirales* and an inverse correlation between the virome and bacterial microbiome, suggesting an hypothesis where changes in the gut virome may affect bacterial dysbiosis [90,95,129,136]. The use of data on both bacteriome and virome composition would contribute to improve classification between health and disease.

These findings suggest that the loss of virus-bacterium relationships can cause microbiota dysbiosis and intestinal inflammation. However, whether viruses have a direct role in IBD pathogenesis, or merely reflect underlying dysbiosis remains to be determined.

**Archaeal Dysbiosis**

The human gut microbiota also contains prokaryotes of the domain Archaea. Methane-producing archaea (methanogens) have been associated with disorders of the gastrointestinal tract and dysbiosis. Methanogens play an important role in digestion, improving polysaccharide fermentation by preventing accumulation of acids, reaction end-products and hydrogen gas [173].
The two reviewed studies including archaeal analysis have shown that the variable prevalence of methanogens in different individuals may play an important role on IBD pathogenesis [61,71]. Lecours et al. showed that the abundance of *Methanosphaera stadtmanae* in fecal samples was significantly higher in IBD patients than in healthy subjects. Interestingly, only IBD patients developed a significant anti-*Msp. stadtmanae* immunoglobulin G response, indicating that the composition of archaeal microbiome appears to be an important determinant of the presence or absence of autoimmunity [61].

The other study demonstrated an inverse association between *Methanobrevibacter smithii* load and susceptibility to IBD, which could be extended to IBD patients in remission as they found that *Mbb. smithii* load was markedly higher in healthy subjects compared to IBD patients [71]. Although archaeal diversity in the gastrointestinal tract is far lower than that of bacteria, these microorganisms can also exert inflammatory effects and their consideration in microbiome studies may be crucial for developing optimal diagnostics and prognostics tools.

### Disease Activity and Severity

Different disease activity and severity have been described among IBD patients with a given clinically defined condition, suggesting that, in the context of microbiome dysfunction, each condition may present different microbial profiles. The reviewed studies showed a clear difference in the gut microbiota associated with different disease activity and severity in IBD patients.

Dysbiosis was evidenced by Tong et al. [83] at remission, where highly preserved microbial groups accurately classified IBD status during disease quiescence, suggesting that microbial dysbiosis in IBD may be an underlying disorder not only associated with active disease. In general, compared to inactive disease, bacterial diversity and richness are reduced in active disease. Studies of intestinal microbiota in active/inactive IBD patients have consistently shown an increase in *F. prausnitzii* and *Clostridiales* in inactive IBD compared to active IBD, and the increase in *Proteobacteria* in active IBD compared to inactive IBD. Besides, *F. prausnitzii* and *R. hominis* display an inverse correlation with disease activity [51,54,56,59,60,68,114,135,137,139,149].

Some studies showed that the genus *Bifidobacterium* is significantly decreased in stool samples during the active phase of CD and UC compared to the remission phase [43,49,68]. On the contrary, biopsies showed a higher abundance of *Bifidobacterium* during active UC, and the proportion of *Bifidobacterium* was significantly higher in biopsies than in the fecal samples in active CD patients [60]. Some controversial results were also found as other researchers did not find a correlation between microbiota and disease activity [35,45,50,101,105,138].

Regarding IBD severity, different microbial abundance was detected in both biopsies and fecal samples from patients with more aggressive disease, and gut dysbiosis was not only related to current activity but also to the course of the disease. In biopsies, *Firmicutes* showed a significant decrease and *Proteobacteria* a significant increase in more aggressive CD [135], and *Bifidobacterium* was inversely correlated with IBD severity [54,135,149]. The risk of flare was associated with reduced microbial richness, increased dysbiosis index and higher individualized microbial instability [74,122,132,137,146,153].

This area is still in its infancy and some results are inconsistent between studies. Several studies have evidenced microbiota signatures of disease activity and severity and the likelihood of a flare-up. However, more research is necessary to identify specific microbial taxa.

### 3.3. Gut Microbiome-Based Biomarkers in IBD

Ideal biomarkers should be easy to obtain, easy to determine, non-invasive, cheap, and capable of providing rapid and reproducible results. Non-invasive tests for IBD are already available, including serum antibodies [174,175], imaging-based screenings [176], and fecal biomarkers [177]. However, endoscopy remains the gold standard for IBD diagnosis, as
the aforementioned non-invasive tests are limited to active disease and their outcome can be interfered by diseases other than IBD limiting their clinical utility.

As a non-invasive, cost-effective technique, microbiome-based biomarkers might have great potential for early-stage disease detection and disease course prognosis as well as for treatment based on patient stratification. To this end, several attempts have been made to develop indices of dysbiosis based on relative abundances of selected microbial taxa in IBD patients compared to those of a healthy population. In stool samples, a machine learning algorithm using a combination of 50 operational taxonomic units was able to differentiate remission from active CD [178], and the genera *Collinsella* and *Methanobrevibacter* could be used to differentiate between UC and CD [109]. In biopsies, *Faecalibacteria* and *Papillibacter* were indicators of IBD status [98], *F. prausnitzii* and *E. coli* were used for differential diagnosis of CD (ileal/colonic) [30], supervised learning classification models were able to classify IBD at specific intestinal locations [65], and microbiome shifts predicted patient outcome [62,64,132,137,145,154]. In biopsies, stool and blood a dysbiosis score accurately stratified IBD patients [132].

In the previous sections, differential results on the gut microbiome between CD and UC, IBD and healthy subjects or between different disease activities have been described. Such research on the IBD microbiome has evidenced that (1) alterations in the abundance of certain microbial taxa or (2) in the structure of the microbial community, (3) the decreased bacterial richness and/or diversity and (4) the decreased microbial community stability could be used as potential biomarkers in the field.

Nevertheless, due to the high microbiome diversity between individuals, and within the same individual over time, the predictive value of these potential indicators is currently far below the level required for utility in diagnosis, prognosis, or response to treatment. Nonetheless, the increasing number of microbiome studies along with the use of longitudinal approaches pave the way to the refinement of microbiome-based biomarkers as useful disease indicators.

4. Concluding Remarks and Future Perspectives

The study of the human microbiome and its involvement in human health is nowadays one of the most active research topics in biomedicine. A simple search for “Microbiota” and “disease” within PubMed Database reveals almost 28,000 hits to date (august 2020). Given the potential clinical application of the microbiome, the number of studies in this field is rapidly increasing. However, some limitations can be found across these studies, including different methodologic approaches, small cohort sizes, different microbiome analysis methods and sample types and sites, main focus on bacteria, and influence of disease activity and treatment interventions. Therefore, these limitations result in variable findings, difficulty to establish comparison between studies and lack of reproducibility of microbiome signatures across studies.

Recent studies based on novel DNA sequencing methods have revealed major differences in microbial taxonomic and functional composition between IBD patients and healthy individuals. The current knowledge guides us to move our focus from community composition to the understanding of the interactions between microbial functions and the IBD gut microbiome.

The microbiota is very specific to an individual and variable in time, and therefore studies need to go from searching for correlation to searching for causation through longitudinal approaches. One important factor that we must keep in mind when studying the microbiome is that it is a “living entity” subject to variability. This variability is even more evident in the IBD microbiome. To better understand the IBD–microbiome connection, we require prospective longitudinal studies, along with following populations with early-onset IBD. The question of whether dysbiosis precedes the development of IBD and sets the inflammatory process, or merely reflects the altered immune and metabolic environment of the inflamed mucosa, remains to be answered. For this reason, it is of paramount importance to study newly diagnosed treatment-naïve patients, where the
Microbiome can be studied at the beginning of the disease and without the influence of any IBD treatment. Developing unified approaches to the accurate quantitative assessment of the gut microbiome would contribute to comparisons among studies and to its further clinical application.

The main feature in IBD gut dysbiosis is the decrease in beneficial bacteria and the increase in pathogens. Gut microbiome studies are mainly focused on bacteria, yet beyond bacteria, the gut microbiome is composed of other microorganisms such as viruses, fungi or archaea, which play a role in IBD etiology and/or in bacterial population control. In addition, it is currently known that disease activity and severity influence the gut microbiome, thereby affecting the results. IBD can be considered as a “multimicrobial” disease with no single causative microorganism, in which more severe disease is linked to reduced gut microbial diversity, and proliferation or reduction in specific taxa. Therefore, future studies should include the whole community for a deeper understanding of this disease.

The usefulness of the gut microbiome as a tool towards targeted non-invasive biomarkers for IBD has been evaluated by compelling studies. An acceptable biomarker may help in early diagnosis and classification of IBD as well as in the prediction of disease outcome. Overall, IBD clinical management would benefit from the identification of microbiome-based biomarkers, which could provide less invasive assessment tools, enable personalized treatments, and reduce the health care economic burden associated with IBD. Collectively, these microbiome data represent a valuable data source that can be continually mined to identify associations between the microbiome and IBD for a deeper pathophysiological understanding which may promote the development of clinical strategies, including disease prevention, treatment, stratification and assessment of high-risk population.

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Conflicts of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Appendix A

Table A1. PRISMA Checklist.

| Section/Topic                  | # | Checklist Item                                                                 | Reported on Page # |
|-------------------------------|---|-------------------------------------------------------------------------------|--------------------|
| TITLE                         |   | Title 1 Identify the report as a systematic review, meta-analysis, or both.    | 1                  |
| ABSTRACT                      |   | Structured summary 2 Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number. | 1                  |
| INTRODUCTION                  |   | Rationale 3 Describe the rationale for the review in the context of what is already known. | 1–2                |
|                               |   | Objectives 4 Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS). | 2                  |
| METHODS                       |   | Protocol and registration 5 Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number. | 2                  |
Table A2. PRISMA Checklist.

| Section/Topic                  | #   | Checklist Item                                                                 | Reported on Page # |
|-------------------------------|-----|-------------------------------------------------------------------------------|--------------------|
| **TITLE**                     |     |                                                                               |                    |
| Eligibility criteria          | 6   | Specify study characteristics (e.g., PICOS, length of follow-up) and report    | 2                  |
|                               |     | characteristics (e.g., years considered, language, publication status) used as |                    |
|                               |     | criteria for eligibility, giving rationale.                                  |                    |
| Information sources           | 7   | Describe all information sources (e.g., databases with dates of coverage,     | 2                  |
|                               |     | contact with study authors to identify additional studies) in the search and  |                    |
|                               |     | date last searched.                                                          |                    |
| Search                        | 8   | Present full electronic search strategy for at least one database, including   | 2                  |
|                               |     | any limits used, such that it could be repeated.                             |                    |
| Study selection               | 9   | State the process for selecting studies (i.e., screening, eligibility,        | 2                  |
|                               |     | included in systematic review, and, if applicable, included in the            |                    |
|                               |     | meta-analysis).                                                              |                    |
| Data collection process       | 10  | Describe method of data extraction from reports (e.g., piloted forms,        | 2                  |
|                               |     | independently, in duplicate) and any processes for obtaining and confirming   |                    |
|                               |     | data from investigators.                                                     |                    |
| **RESULTS**                   |     |                                                                               |                    |
| Study selection               | 17  | Give numbers of studies screened, assessed for eligibility, and included in   | 2                  |
|                               |     | the review, with reasons for exclusions at each stage, ideally with a flow    |                    |
|                               |     | diagram.                                                                     |                    |
| Study characteristics         | 18  | For each study, present characteristics for which data were extracted (e.g.,  | 3–8                |
|                               |     | study size, PICOS, follow-up period) and provide the citations.               |                    |
| Risk of bias within studies   | 19  | Present data on risk of bias of each study and, if available, any outcome     | N/A                |
|                               |     | level assessment (see item 12).                                               |                    |
| Results of individual studies | 20  | For all outcomes considered (benefits or harms), present, for each study: (a) | N/A                |
|                               |     | simple summary data for each intervention group (b) effect estimates and       |                    |
|                               |     | confidence intervals, ideally with a forest plot.                            |                    |
| Synthesis of results          | 21  | Present results of each meta-analysis done, including confidence intervals and | N/A                |
|                               |     | measures of consistency.                                                      |                    |
| Risk of bias across studies   | 22  | Present results of any assessment of risk of bias across studies (see Item 15).| N/A                |
| Additional analysis           | 23  | Give results of additional analyses, if done (e.g., sensitivity or subgroup   | N/A                |
|                               |     | analyses, meta-regression [see Item 16]).                                    |                    |
| **DISCUSSION**                |     |                                                                               |                    |
| Summary of evidence           | 24  | Summarize the main findings including the strength of evidence for each main   | N/A                |
|                               |     | outcome; consider their relevance to key groups (e.g., healthcare providers,   |                    |
|                               |     | users, and policy makers).                                                    |                    |
| Limitations                   | 25  | Discuss limitations at study and outcome level (e.g., risk of bias), and at   | N/A                |
|                               |     | review-level (e.g., incomplete retrieval of identified research, reporting bias).|                    |
| Conclusions                   | 26  | Provide a general interpretation of the results in the context of other        | 8–9                |
|                               |     | evidence, and implications for future research.                              |                    |
| **FUNDING**                   |     |                                                                               |                    |
| Funding                       | 27  | Describe sources of funding for the systematic review and other support (e.g., | 9                   |
|                               |     | supply of data); role of funders for the systematic review.                  |                    |
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