



COMPUTATIONAL ANDSTRUCTURAL BIOTECHNOLOGY

JOURNAL



journal homepage: www.elsevier.com/locate/csbj

# Analysis methods for the gut microbiome in neuropsychiatric and neurodegenerative disorders



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## ARTICLE INFO

Article history: Received 6 November 2021 Received in revised form 24 February 2022 Accepted 24 February 2022 Available online 26 February 2022

Keywords: Animal model Brain Genomic analysis Gut microbiome Metabolomics

# ABSTRACT

For a long time, the central nervous system was believed to be the only regulator of cognitive functions. However, accumulating evidence suggests that the composition of the microbiome is strongly associated with brain functions and diseases. Indeed, the gut microbiome is involved in neuropsychiatric diseases (e.g., depression, autism spectrum disorder, and anxiety) and neurodegenerative diseases (e.g., Parkinson's disease and Alzheimer's disease). In this review, we provide an overview of the link between the gut microbiome and neuropsychiatric or neurodegenerative disorders. We also introduce analytical methods used to assess the connection between the gut microbiome and the brain. The limitations of the methods used at present are also discussed. The accurate translation of the microbiome information to brain disorder could promote better understanding of neuronal diseases and aid in finding alternative and novel therapies.

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#### https://doi.org/10.1016/j.csbj.2022.02.024

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# 1. Introduction

Over the past decades, a considerable amount of research has shown the gut microbiome to be associated with brain functions. Despite Hippocrates' insightful claims, "All diseases begin in the gut," the interaction between the gut and brain is not fully understood yet. For a long time, neuropsychiatric and neurodegenerative diseases have been investigated by solely focusing on the central nervous system. However, recent groundbreaking studies have indicated that microorganisms living in the gut could influence the development of brain disorders by changing metabolites and neurochemicals [1,2]. The chemicals produced by intestinal microorganisms can be transported across the intestinal epithelium [3]. The gut microbiome communicates with the brain via different pathways such as the neurological pathways [4,5], endocrine pathways [6-8], and immune pathways [9-12]. The vagus nerve is also a connection that transmits information about the inner organs from the gastrointestinal tract to the brain [13]. Under stress conditions, the hypothalamic-pituitaryadrenal (HPA) axis responds by inducing hormone secretion, such as cortisol, through the release of adrenocorticotropic hormone. Cortisol regulates immune signaling responses, which affect the intestinal barrier [14]. Short-chain fatty acids (SCFAs), one of the metabolites produced by bacteria, can be transported across the intestinal epithelium and reduce the gene expression related to tight junction formation in the brain microvascular endothelial cells, which makes the blood-brain barrier more permeable and could consequently regulate microglia homeostasis [12].

This interesting association between the gut microbiome and the brain has drawn researchers and encouraged them to unravel this association, particularly in terms of neurological diseases. Despite their limitations, most studies have relied on animal models to mimic the details of human diseases. Advanced analysis methods have been established to study the relationship between the gut and the brain and confirm the underlying biological mechanisms to develop treatments against brain disorders. In this review, the current state of the gut microbiome-brain research was discussed, and the evidence was summarized in favor of the involvement of the gut microbiota in neuropsychiatric and neurodegenerative disorders (Fig. 1).

# 2. Analysis methods and bioinformatics tools for analyzing the microbiome

Various techniques for analyzing the microbiome are available. Here we discuss the basic analysis methods, the primary tools available for each method (Table 1), and previous studies about the relationship between the gut microbiome and neurodegenerative or neuropsychiatric diseases.

# 2.1. Marker gene analysis

Marker gene analysis is the most popular bioanalytical method to survey microbial populations and phylogenies. The 16S ribosomal RNA (rRNA) is globally found in bacteria. It contains a highly conserved domain and a species-dependent hypervariable sequence region [28]. Owing to these qualities, 16S rRNA can be easily targeted for amplification using polymerase chain reaction (PCR) using primers that bind to the conserved domain. 18S rRNA or internal transcribed spacers are used in fungi and yeasts as marker genes [29]. The microbial phylogeny and relative abundance of certain microbes in samples are determined from the amplified marker gene proportion [30]. Preparing the sample for cloning, sequencing the marker gene, and analyzing the resulting amplicon are tasks that are relatively inexpensive and straightforward. Wellknown bioinformatics tools, such as QIIME [31], and large-scale public datasets, such as the Ribosomal Database Project [32], can be used to enhance the analysis. However, the results are subjected to PCR-related biases [33]. Indeed, biases may occur during the amplification process because the primer binding affinity for the conserved regions is not equal for the 16S rRNA component from various species. Some PCR parameters, including variable region selection, amplicon size, and the number of PCR cycles, may be further sources of biases [34]. These biases can be decreased by



Fig. 1. Overall analysis process of the microbiome and neuronal diseases.

#### Table 1

Tools for microbiome analysis.

Name	Description	Analysis	Link
Mothur [15]	Includes useful tools such as SONS, DOTOUR, TreeClimber, LIBSHUFF, $\int$ -LIBSHUFF, and UniFrac. Implemented over 25 calculators for quantifying parameters to estimate $\alpha$ and $\beta$ diversity.	Marker gene analysis	https://mothur.org/
QIIME 2 [16]	Plugin containing q2-cscs, q2-metabolomics, q2-shotgun, q2-metaphlan2, and q2- picrust methods. Provides various interactive visualization tools.	Marker gene analysis	http://qiime2.org/
DADA 2 [17]	Performs the full amplicon process: filtering, dereplication, sample inference, chimera-filtered, and merging paired reads.	Marker gene analysis	https://benjjneb.github.io/dada2/ index.html
Phyloseq [18]	Converts data output of OUT clustering pipelines into a suitable form amenable to modern analysis methods such as discriminant analysis, and canonical correspondence analysis.	Marker gene analysis	http://www.bioconductor. org/packages/release/bioc/ html/phyloseq.html
VEGAN [19]	Provides basic functions of diversity analysis and multivariate analysis.	Marker gene analysis	https://github.com/vegandevs/ vegan
DESeq 2 [20]	Focused on quantitative analysis rather than on differential expression. Uses the negative binomial distribution, the Wald, and the Likelihood Ratio Tests.	Marker gene analysis	https://bioconductor. org/packages/release/bioc/html/ DESeq2.html
IDBA-UD [21]	Based on the iterative de Bruijn graph assembler for standard metagenomics and single-cell analysis.	Shotgun Metagenomics	http://www.cs.hku.hk/~alse/idba_ ud
SPAdes [22]	Uses k-mers to construct the de Bruijn graph for mate-pair, pair-end reads, and unpaired reads. SPAdes pipeline with two separate modules: 1) BayesHammer and 2) SPAdes.	Shotgun Metagenomics	http://cab.spbu.ru/software/ spades/
MEGAHIT [23]	Based on the construction of succinct de Bruijn graphs with CPU-favored graph module.	Shotgun Metagenomics	https://www.metagenomics. wiki/tools/assembly/megahit
MetaPhlAn 3 [24]	Performs unambiguous taxonomic assignment by MetaPhlAn markers of clade- specific. Possibility to get additional information on identifying metagenomics data using StrainPhlAn 3, PanPhlAn 3, PhyloPhlAn 3, and HUMAnM 3.	Shotgun Metagenomics	https://huttenhower.sph.harvard. edu/metaphlan/
MG-RAST [25]	Web application server based on SEED framework for metagenomics data. Performs five pipelines: 1) data hygiene, 2) feature extraction, 3) feature annotation, 4) profile generation, 5) data loading.	Shotgun Metagenomics	https://www.mg-rast.org/
PICRUSt 2.0 [26]	Enables output of MetaCyc predictions that will be linked with shotgun metagenomics results.	Shotgun Metagenomics	https://github.com/picrust/ picrust2/wiki
SOAP denovo 2 [27]	Consists of six modules: 1) handle read error correction, 2) de Bruijn graph construction, 3) contig assembly, 4) paired-end reads mapping, 5) scaffold construction, and 6) gap closure.	Metatranscriptomics	https://sourceforge.net/ projects/soapdenovo2/

optimizing primers and PCR parameters [35]. After proper optimization, marker gene analysis can be applied to different research pipelines and sample types. However, the methodology only offers limited and low-resolution information about the microbial community, without microbial genome expression.

## 2.2. Whole genome analysis

Whole genome analysis is a sequencing method that detects whole genomic contents in a sample and detects bacteria, fungi, yeasts, and viral DNAs [36]. With dramatic advances in sequencing technologies, parallel and rapid total DNA sequencing can be performed with next-generation sequencing (NGS), also known as high-throughput sequencing (HTS). Various commercial NGS platforms based on different sequencing technologies are available [37]. Additionally, shotgun sequencing allows profiling functional representations or metabolic pathways in the microbiota community [38]. In silico alignment tools have been introduced to analyze sequencing data, which consist of billions of short sequence fragments called reads [39]. Read profiling techniques provide taxonomy information and annotate genes from the sample by comparing unassembled DNA reads to reference data. Many read-based profiling tools extract all short sequences with length k from the sequencing result (k-mer) and build a de Bruin graph to reconstruct long DNA sequences [40]. Burrows–Wheeler transform algorithm can also be used in read-based profiling [41]. Rapid and accurate mapping for entire reads are achieved by compressing large-scale genomic sequence databases with the algorithm. Another method for reading alignment consist in assembling reads into longer sequences also called contigs with the overlap-layout-consensus (ORC) algorithm. The algorithm function in three steps: 1) search overlapping sequences between

the read sequences, 2) layout and assemble the overlapping reads, and 3) search for the most reliable sequences from the assembled contigs [42]. Although processes including sample preparation, whole-genome sequencing, and analysis are relatively complex and expensive, whole metagenome analysis offers detailed information about the microbial community [43].

# 2.3. Meta-transcriptome analysis

Metagenomic analysis provides information about the microbial populations or the microbial genomic contents; however, it cannot identify transcriptionally active microbial genes or functions. In metatranscriptome analysis, a whole-genome sequencing technique is applied to conduct RNA sequencing. Microbial transcription, gene expression, and functional outputs are profiled from the sequencing data [44]. Metatranscriptome analysis allows to estimate the transcription activity and identify active metabolic pathways of species from the microbiome community [45]. However, because of the instability of RNA molecules, sample preparation, amplification, and analysis are the most expensive and complex among the microbial analyse techniques. Despite the cost, metatranscriptome analysis offers unique information to the microbiome research community.

# 2.4. Metabolome and meta-proteome analyses

Metabolome analysis is an emerging tool that can be used to investigate the status of the microbial community by focusing on functional dysregulation of the gut microbiome because small metabolites are the actual players of the gut-brain communication [46]. The extraction and quantification of metabolites enables the study of active metabolic pathways expressed by the gut microbiome and investigation of the interactions of these metabolites with host metabolism. Of the metabolites produced by gut microorganisms, SCFAs are the main bacterial metabolites associated with gut physiology. Of SCFAs, acetate (C1), propionate (C2), and butyrate (C3) are the most abundant ones [47]. Acetate is generally produced from acetyl-CoA, a metabolite produced *via* glycolysis. Propionate and butyrate are the products of metabolites produced via glycolysis or amino acid metabolism [47].

SCFAs can be separated and quantified using different methods such as nuclear magnetic resonance, mass spectroscopy (MS), liquid chromatography (LC), and gas chromatography (GC) [48]. It is difficult to identify the source of the detected metabolites or to differentiate the microbial metabolites from host-derived products based solely on metabolome analysis. However, the gut microbial phylogeny or functional alterations can be established by measuring relative amounts of metabolites in the samples and comparing this profile to reference libraries [49].

As in metabolome analysis, functional changes in the microbiome can be characterized by focusing on proteins as markers. The metaproteomic analysis pipeline consists of extracting proteins from microbiome samples, digesting the proteins into peptides, separating these peptides using LC-MS, and identifying and quantifying proteins by searching and comparing data to metagenome databases [50]. After annotating proteins from the samples using various protein databases, such as UniProt [51] or BRENDA [52], functional analysis or taxonomic assignment can be conducted. Moreover, phylogenetic or metagenomic information about the microbial community in the samples can be linked to the proteomic data for further analysis. Proteomic analysis provides information about real activities of the microbiome or allows to identify potential drug targets for certain pathways or diseases [53]. However, the sample preparation is very complex and the computational analysis is also difficult due to the large and complex proteomics datasets [54].

# 3. Gut-brain axis (GBA)

The gut-brain axis (GBA) consists of the biochemical signaling between the gut microbiome and the brain resulting in changes in the central nervous system state [55]. In recent years, the research focus was on the gut microbiome functions influencing the GBA bidirectional communication [6,56–58], which is important to maintain homeostasis of the gastrointestinal and central nervous systems of animals [59–61]. It is likely that the GBA mediates various aspects of the pathogenesis of brain diseases, and intensive research is necessary to clarify the underlying mechanisms. Various studies have shown the potential relationships between the gut microbiome and neurodegenerative or neuropsychiatric diseases (Table 2).

# 3.1. Gut microbiome and neuropsychiatric disorders

Many studies in animal models and human patients have shown an association between the gut microbiome and the development of various neuropsychiatric disorders. Most studies reported changes in the bacterial community but lacked explaining how these changes affected brain physiology. Although many mediators, such as SCFAs, have been identified recently, the bacterial species that produce such mediators and how they influence brain functions needs further clarification.

# 3.1.1. Schizophrenia (SCZ)

SCZ is a psychiatric disorder characterized by positive symptoms such as delusions, hallucinations, and thought abnormalities. Patients with SCZ also exhibit negative symptoms such as anhedonia, abolition, social withdrawal, and poverty of thought [103]. SCZ is associated with dysregulation of the dopaminergic, glutamatergic,  $\gamma$ -aminobutyric acid (GABA)ergic, and cholinergic neurotransmitters [104].

The comparison of differences in fecal microbiota between SCZ patients and healthy controls was performed using 16S rRNA sequencing and diversity analyses [83]. In the study, a lower microbiome diversity in patients with SCZ patients. At the phylum level, Proteobacteria were considerably enriched in patients with SCZ. At the genus level, Blautia, Coprococcus, and Roseburia were found less often in patients with SCZ, whereas Succinivibrio, Megasphaera, Collinsella, Clostridium, Klebsiella, and Methanobrevibacter were present to a larger extent. In another study, 16S rRNA gene sequence analysis and diversity analysis were performed [84], and a decreased microbiome alpha-diversity, that was assessed using the species richness indices and species diversity indices. and disturbance of gut microbial composition in SCZ patients were observed. Specifically, the abundance of Lachnospiraceae and Ruminococcaceae, which are the most abundant in the order Clostridiales, was reduced in the gut microbiome of SCZ patients. In addition, significant deviations in Aerococcaceae, Bifidobacteriaceae, Brucellaceae, Pasteurellaceae, and Rikenellaceae between SCZ patients and healthy controls were observed. Based on the deviations, a stepwise regression analysis was conducted and showed the potential of SCZ diagnosis by gut microbiome analysis. This finding suggests that patients with SCZ have abnormal microbial status. Furthermore, the changes in the gut microbiota, resulting from the transfer of human SCZ fecal microbiome to mice, led to SCZ-relevant hyperactive behavior in mice who received SCZ microbiota, relative to healthy control microbiota mice [84,105]. The study indicates that glycerophospholipids, which are key regulators of synaptic function are reduced in the hippocampus and SCZ mice serum, but increased in fecal samples of SCZ mice. This suggests that lower levels of glycerophospholipids are consistent with symptoms of patients with SCZ having synaptic deficits and disconnectivity.

There have been attempts to identify specific bacterial metabolites that are associated with SCZ symptoms [106]. SCFA-producing bacteria, such as *Blautia*, *Coprococcus*, and *Roseburia*, were reduced in patients with SCZ [83]. Butyrate, an SCFA, can cross the BBB and inhibit histone deacetylase 1 (HDAC1) [107,108]. Because HDAC1 level is elevated in the PFC and hippocampus of patients with SCZ, the elevation might be due to the reduced butyrate level produced by the bacterial species [109,110]. A study hypothesized that the increased populations of *Clostridiales*, *Prevotella*, and *Lactobacillus ruminis* increased SCFA levels that stimulated TNF production, and the increased SCFA activated microglia and disrupted membrane metabolism in patients with SCZ [111]. However, specific SCFAs were not measured in the study.

Germ-free (GF) mice transplanted with the microbiome of SCZ patients have low glutamate and high GABA levels in the hippocampus and display SCZ-relevant behaviors [84]. Dickerson *et al.* showed that SCZ subjects treated with probiotics containing *L. rhamnosus* strain GG and *Bifidobacterium lactis* strain Bb12 had less severe bowel difficulties, although there was no difference in the psychiatric symptoms compared with those of the placebo group [112]. Using the same strain as Dickerson *et al.*, Tomasik *et al.* showed a decrease in the von Willebrand factor (vWF) amounts and an increase in monocyte chemotactic protein-1 (MCP-1) and brain-derived neurotrophic factor (BDNF) levels in probiotic-treated SCZ patients. It also modulated immune and intestinal epithelial cells through the IL-17 cytokine family. They hypothesized that this controlled gastrointestinal leakage [113].

Severance *et al.* conducted a longitudinal, double-blind, and placebo-controlled pilot study on 56 schizophrenia subjects to

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# Table 2

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Disease	Number of subjects	Sample	Analysis method	Alterations of microbiota
AD [62]	AD patients = 61, healthy controls = 30	Human	Gastrointestinal endoscopy	The elimination of pathogenic bacteria such as <i>Helicobacter pylori</i> by triple eradication antibiotic regimen (omeprazole,clarithromycin, and amoxicillin) led to improved cognitive and functional status parameters in AD patients. The eradication of <i>Helicobacter milori</i> influenced AD manifectations
AD [63]	AD patients = 60	Human blood	ELISA, cognitive test	Probiotic consumption (Lactobacillusacidophilus, Lactobacilluscasei, Bifidobacteriumbifidum, and Lactobacillusfermentum) induced significant cognitive improvements and affected metabolic pathways in AD
AD [64]	AD patients = 25, non-AD = 94	Human fecal samples	16S rRNA gene sequencing	Differences in bacterial abundance, including decreased <i>Firmicutes and Bifidobacterium levels</i> as well as increased <i>Bacteroidetes</i> amounts, in the microbiome of AD participants
AD [65]	AD patients = 43, healthy controls = 43	Human fecal samples	16S rRNA gene sequencing	Several bacteria taxa were different in AD patients than those in controls, such as <i>Bacteroides</i> , <i>Actinobacteria</i> , <i>Ruminococcus</i> , <i>Lachnospiraceae</i> , and <i>Selenomonadales</i> .
AD [66]	n = 32	Mice fecal samples	16S rRNA gene sequencing	The composition and diversity of gut microbiota changed with aging in a tauopathy mice model. In detail, at the phylum level, the relative abundance of <i>Bacteroidetes</i> was increased, whereas that of <i>Firmicutes</i> was decreased in 3-month-old P301L mice compared with that in aged-matched wild-type mice. <i>Actinobacteria</i> levels were decreased in 3 to 6-month- old P301L mice. Less <i>Tenericutes</i> was found in 10- month-old P301L mice. Several specific macrobiota were highly associated with the levels of AT8-tau or pT231-tau protein in the brain
AD [67]	n = 24	Mice fecal samples	16S rRNA gene sequencing	At the phylum level, <i>Proteobacteria</i> and <i>Verrucomicrobia</i> levels were increased in AD mice. At the genus level, <i>Ruminococcus</i> and <i>Butricicocus</i> were less abundant in AD mice.
AD [68]	n = 18	Mice fecal samples	16S rRNA gene sequencing (qPCR)	At the phylum level, <i>Firmicutes</i> levels were increased in AD mice.
PD [65]	PD patients = 75, healthy controls = 45	Human fecal samples	16S rRNA gene sequencing	Increase in the abundance of four bacterial families ( <i>Eubacteriaceae</i> , <i>Bifidobacteriaceae</i> , <i>Aerococcaceae</i> , and <i>Desulfovibrionaceae</i> ) and significant decrease in the abundance of 17 bacterial families ( <i>Lachnospiraceae</i> , etc.) in PD patients compared to the levels found in control
PD [69]	PD patients = 76, healthy controls = 78	Human fecal	16S/18S rRNA gene sequencing	Increased Akkermansia levels in PD patients.
PD [70]	PD patients = 64, healthy controls = 64	fecal samples	16S rRNA gene sequencing	Increased <i>Bifidobacterium</i> levels and decreased amounts of <i>Prevotellaceae</i> and <i>Roseburia</i> in PD patients.
PD [71]	PD patients = 52, healthy controls = 36	Human fecal samples	16S rRNA gene sequencing (with qRT- PCR)	The amounts of <i>Lactobacillus</i> were increased, whereas the <i>Clostridium coccoides</i> and <i>Bacteroides fragilis</i> groups were less abundant in PD patients.
PD [72]	PD patients = 31, healthy controls = 28	Human fecal samples	Shotgun metagenomic sequencing	Increased amounts of Verrucomicrobiaceae (Akkermansia muciniphila) and unclassified Firmicutes in PD patients. Prevotellaceae (Prevotella copri) and Erysipelotrichaceae (Eubacterium biforme) were markedly less abundant in PD patients than they were in controls
PD [73]	PD patients = 197, healthy controls = 130	Human fecal samples	16S rRNA gene sequencing	Increased Bifidobacteriaceae, Lactobacillaceae, Tissierellaceae, Christensenellaceae, and Verrucomicrobiaceae levels and decreased amounts of Lachnospiraceae Pasteurellaceae in PD patients
PD [74]	PD patients = 89, healthy controls = 66	Human fecal samples	16S rRNA gene sequencing	Increased abundance of Christensenella, Catabacter, Lactobacillus, Oscillospira, Bifdobacterium, Christensenella minuta, Catabacter hongkongensis, Lactobacillus mucosae, Ruminococcus bromii, and Papillibacter cinnamivorans, and decreased levels of Dorea, Bacteroides, Prevotella, Faecalibacterium, Bacteroides massiliensis, Stoquefichus massiliensis, Bacteroides coprocola, Blautia glucerasea, Dorea longicatena, Bacteroides dorei, Bacteroides plebeus, Prevotella copri, Coprococcus eutactus, and Ruminococcus callidus in PD patients.

(continued on next page)

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# Table 2 (continued)

Disease	Number of subjects	Sample	Analysis method	Alterations of microbiota
PD [75]	PD patients = 9, healthy controls = 13	Human fecal	16S rRNA gene sequencing	Increased Akkermansia and Bifidobacterium levels and decreased Prevotella amounts in PD patients.
PD [76]	PD patients = 10, healthy controls = 10	samples Human fecal samples	16S rRNA gene sequencing	Increased abundance of Ruminococcaceae, Verrucomicrobiaceae, Porphyromondaceae, Hydrogenoanaerobacterium, and Lachnospiraceae and decreased amounts of Bacteroides and Prevotellaceae in
PD [77]	PD-MCI (mild cognitive impairment) = 13, PD-NC (normal cognition) = 14, healthy controls = 13	Human fecal samples	16 s rRNA gene sequencing, gas chromatography-mass spectrometry	PD patients. The fecal microbial diversities of PD-MCI and PD-NC were higher than that of healthy controls. In PD-MCI, <i>Alistipes, Barnesiella, Butyricimonas</i> , and <i>Odoribacter</i> levels were higher than those in the other two groups. Compared with the PD-NC group, the genus <i>Blautia</i> and <i>Ruminococcus</i> were less abundant in the PD-MCI group.
PD [78]	PD patients = 38, healthy controls = 34	Human mucosal and fecal samples	16S rRNA gene sequencing	Increased abundance of <i>Faecalibacterium</i> and <i>Ralstonia</i> in mucosal samples. Increased levels of <i>Blautia, Coprococcus,</i> and <i>Roseburia</i> in fecal samples.
PD [79]	PD patients = 29, healthy controls = 29	Human fecal samples	16S rRNA gene sequencing (next- generation-sequencing)	Increased abundance of <i>Lactobacillaceae</i> , <i>Barnesiellaceae</i> , and <i>Enterococcaceae</i> in PD patients.
PD [80]	PD patients = 24, healthy controls = 14	Human fecal samples	16S rRNA gene sequencing	More Escherichia-Shigella, Streptococcus, Proteus, and Enterococcus were found in PD patients than in healthy controls. Blautia, Faecalibacterium, and Ruminococcus levels were decreased in PD patients.
PD [81]	PD patients = 45, healthy controls = 45	Human fecal samples	16S rRNA gene sequencing	Clostridium IV, Aquabacterium, Holdemania, Sphingomonas, Clostridium XVIII, Butyricicoccus, and Anaerotruncus levels were increased in PD patients.
PD [82]	PD patients = 34, healthy controls = 34	Human fecal samples	16 s rRNA gene sequencing (RT-qPCR)	Enterobacteriaceae Phylum was more abundant in PD patients, whereas the levels of Bacteroidetes and Prevotellaceae were decreased.
Schizophrenia [83]	SCZ patients = 64, healthy controls = 53	Human fecal samples	16S rRNA gene sequencing (metagenomes, PICRUSt analysis)	Proteobacteria levels were significantly increased in SCZ patients. At the genus level, <i>Succinivibrio, Megasphaera,</i> <i>Collinsella, Clostridium, Klebsiella,</i> and <i>Methanobrevibacter</i> were more abundant in SCZ patients than they were in healthy controls, whereas <i>Blautia, Coprococcus,</i> and <i>Roseburia</i> levels were decreased in SCZ patients.
Schizophrenia [84]	SCZ patients = 63, healthy controls = 69	Human fecal samples	16S rRNA gene sequence analysis	Unmedicated and medicated SCZ patients had a decreased microbiome alpha-diversity index and marked disturbances of gut microbial composition compared to those of healthy controls. Several unique bacterial taxa like <i>Veillonellaceae</i> and <i>Lachnospiraceae</i> were linked to SCZ severity. <i>Aerococcaceae, Bifdobacteriaceae, Brucellaceae</i> , <i>Pasteurellaceae</i> , and <i>Rikenellaceae</i> were different in SCZ patients than these found in healthy controls.
ALS [85]	ALS patients = 50, healthy controls = 50	Human fecal samples	16S rRNA gene sequencing with qPCR	<i>Escherichia coli and Enterobacteriaceae</i> levels were higher in ALS patients than those of healthy controls, whereas the amounts of <i>Clostridium</i> were lower in ALS patients.
ALS [86]	ALS patients = 6, healthy controls = 5	Mice fecal samples	16S rRNA gene sequencing, Shotgun metagenomic sequencing	Decreased levels of Akkermansia muciniphila, Parabaceroides distasonis, Rikenellaceae, Prevotella, Lactobacillus murinus, Alistipes unclassified, and Eggertella unclassified as well as increased Sutterella, Allobaculum, Desulfovibrionaceae, Coprococcus, Oscillospira, Bifidobacterium, Helicobacter hepaticus, Lactobacillus johnsonii, and Lactobacillus reuteri in ALS mice.
	ALS patients = 37, healthy controls = 29	Human fecal samples	Shotgun metagenomic sequencing	The amounts of Anaerostipes hadrus and Bacteroidales bacterium were increased marginally in ALS patients and Bifidobacterium pseudocatenulatum, Clostridium leptum, and Escherichia coli levels were decreased.
ALS [87]	ALS patients = 6, healthy controls = 5	Human fecal samples	16S rRNA gene sequencing	Dorea levels were increased and the amounts of Oscillibacter, Anaerostipes, and Lachnospiraceae were decreased in ALS patients compared with those of healthy controls.
ALS [88]	not mentioned (454 16rRNA sequencing data)	Mice fecal samples	16S rDNA qRT-PCR with pyrosequencing	A gut dysbiosis was evidenced in ALS mice, particularly in terms of reduced levels of butyrate-producing bacteria, including <i>Butyrivibriofibrisolvens</i> and <i>E. coli</i> .
ALS [89]	ALS patients = 8, healthy controls = 8	Human fecal samples	16S rRNA gene sequencing	Increased amounts of <i>Methanobrevibacter</i> and decreased levels of <i>Faecalibacterium</i> and <i>Bacteroides</i> in ALS patients.

 Table 2 (continued)

Disease	Number of subjects	Sample	Analysis method	Alterations of microbiota
Depression [90]	Depressed patients = 5, healthy controls = 5	Mice fecal samples	16S rRNA genes pyrosequencing	Stressed and depressed mice showed changes in microbial diversity with more <i>Desulfovibrionaceae</i> , <i>Rikenellaceae</i> , and <i>Lachnospiraceae</i> families and less
Depression [91]	Depressed patients = 13, healthy controls = 15	Rats fecal samples	16 s rRNA sequencing	Allobaculum and Mucispirillum. At the phylum level, the relative abundances of Actinobacteria and Candidate Division TM7 were decreased in rats that received the depression fecal microbiota transplantation (FMT rats). At the family level, the relative proportions of Bifidobacteriaceae, Coriobacteriaceae, Porphyromonadaceae, Candidate Division TM7 uncultured bacterium, Caldicoprobacteraceae, Alcaligenaceae were decreased in FMT rats. Propionibacteriaceae levels were increased in FMT rats. At the genus level, the relative abundances of Bifidobacterium uncultured, Coriobacteriaceae uncultured, Caldicoprobacter, Roseburia, Allobaculum, Burkholderiales were decreased and Freudenreichii, Staphyloccus, Peptococcus levels were increased in FMT rats.
	depressed patients = 34, healthy controls = 33	Human fecal samples	16 s rRNA sequencing	At the family level, the relative proportions of <i>Prevotellaceae</i> were decreased, whereas those of <i>Thermoanaerobacteriaceae</i> were increased in depressed patients. At the genus level, there was an increase of the relative proportions of <i>Eggerthella</i> , <i>Holdemania</i> , <i>Gelria</i> , <i>Turicibacter</i> , <i>Paraprevotella</i> and <i>Anaerofilum</i> , and decreased amounts of <i>Prevotella</i> and <i>Dialister</i> in depressed patients.
Depression [92]	depressed patients = 58, healthy controls = 63	Human fecal samples	16S rRNA gene sequencing	Actinobacteria levels were increased, whereas Bacteroidetes amounts were decreased in depressed patients compared with the levels in healthy controls
Depression [93]	Active-Major Depressive Disorder (A-MDD) patients = 29, Responded-Major Depressive Disorder (R-MDD) patients = 17, healthy controls = 30	Human fecal samples	16S rRNA gene pyrosequencing	The fecal bacterial α-diversity was increased in A-MDD patients compared with the reversion in carticy controls. The fecal bacterial α-diversity was increased in A-MDD patients compared with that in healthy controls, but there was no difference between R-MDD patients and the HC group. The levels of <i>Bacteroidetes, Proteobacteria,</i> and <i>Actinobacteria</i> were strongly increased, whereas those of <i>Firmicutes</i> were significantly reduced in the A-MDD and R-MDD patients compared with the levels in healthy controls. Depressed patients had increased <i>Enterobacteriaceae</i> and <i>Alistipes</i> levels and decreased amounts of <i>Faecalibacterium</i> .
Autism/ASD [94]	Autism patients = 30, healthy controls = 24	Human fecal samples	16S rRNA gene amplicon in NGS	ASD patients have a higher relative abundance of the families <i>Lactobacillaceae</i> , <i>Bifidobacteraceae</i> , and <i>Veillonellaceae</i> , whereas healthy controls have higher levels of the <i>Prevotellaceae</i> family.
Autism/ASD [95]	Autism patients = 20, healthy controls = 10	Human fecal samples	16S rDNA and 16S rRNA sequencing (by using Bacterial tag-encoded FLX- titanium amplicon pyrosequencing [bTEFAP])	Higher microbial diversity in autism patients. Autism patients have higher levels of the genera <i>Caloramator</i> , <i>Sarcina</i> and <i>Clostridium</i> , and higher amounts of the species of <i>Alistipes</i> and <i>Akkermansia</i> .
Autism/ASD [96]	Autism patients = 13, healthy controls = 8	Human fecal samples	16S rRNA gene sequencing	Autism patients have higher amounts of <i>clostridial</i> species. The abundance and type of <i>Clostridium</i> and <i>Ruminococcus</i> species in autism patients are differ from those of healthy controls. Nine species of <i>Clostridium</i> were found in autism patients but not in controls. In controls, there were three species not found in children with autism.
Autism/ASD [97]	Autism patients = 21, healthy controls = 19	Human fecal samples	16S rRNA gene pyrosequencing	The abundance of the genus <i>Burkholderia</i> was higher and that of the genus <i>Neisseria</i> was lower in autism patients. At the species level, the amounts of two <i>Bacteroides</i> species and <i>Escherichia coli</i> were decreased in autism patients.
Autism/ASD [98]	Autism patients = 40, healthy controls = 40	Human fecal samples	16S rRNA gene pyrosequencing	Significant increase of the Firmicutes/Bacteroidetes ratio in autistic patients due to a reduction of the Bacteroidetes relative abundance. At the genus level, decreased levels of Alistipes, Bilophila, Dialister, Parabacteroides, and Veillonella, and increased amounts of Collinsella, Corynebacterium, Dorea, and Lactobacillus in autistic patients.
Autism/ASD [99]	Autism patients = 58, healthy controls = 39	Human fecal samples	16S rDNA sequencing	Much lower levels of <i>Bifidobacterium</i> , slightly lower levels of <i>Enterococcus</i> , and the much higher levels of <i>Lactobacillus</i> in autism patients. Autism patients were likely to have <i>Bacillus</i> spp. and less likely to have <i>Klebsiella oxytoca</i> .

(continued on next page)

Table 2 (continued)

Disease	Number of subjects	Sample	Analysis method	Alterations of microbiota
Autism/ASD [100]	Autism patients = 33, healthy controls = 7	Human fecal samples	DNA pyrosequencing	At the phylum level, autistic patients have higher Bacteroidetes and lower Firmicutes levels compared with those of healthy controls. Amounts of Desulfovibrio species and Bacteroides vulgatus were increased in autistic patients compared with those of healthy controls.
Autism/ASD [101]	Autism patients = 58, healthy controls = 22	Human fecal samples	FISH analysis using a collection of 59 Cy3-labeled 16S rRNA oligonucleotide probes	ASD patients have the higher amounts of the Clostridium histolyticum group (Clostridium clusters I and II) compared with the levels in healthy controls.
Huntington's disease (HD) [102]	HD patients = 18, healthy controls = 17	Mice fecal samples	16S rRNA gene sequencing	Sex differences: in males, <i>Bacteroidales</i> levels were increased and <i>Clostridiales</i> amounts were decreased. In females, <i>Coriobacteriales, Erysipelotrichales,</i> <i>Bacteroidales,</i> and <i>Burkholderiale</i> levels were increased and <i>Clostridiales</i> amounts were decreased.

explore the effects of probiotic treatment on antibodies against *Candida albicans*. Probiotic administration significantly reduced *Candida albicans* antibody levels, *Candida albicans*-associated gut discomfort, and positive psychiatric symptoms in male individuals [114]. Okubo *et al.* investigated the effects of *Bifidobacterium breve* A-1 administration in 29 SCZ patients with anxiety and depression symptoms. Their HADS total score and PANSS anxiety/depression score were improved and their interleukin 22 (IL-22) and tumor necrosis factor related activation-induced cytokine (TRANCE) levels were increased after treatment with the probiotics [115].

## 3.1.2. Autism spectrum disorder (ASD)

ASD is a neurodevelopmental disorder characterized by repetitive behaviors and impairments in social communication and cognitive functions [116,117]. A combination of genetic and nongenetic factors contributes to ASD. Candidate genes involved in ASD etiology include genes coding for postsynaptic scaffolding proteins, contactins, neurexin family, and chromatin remodeling proteins [118]. Nongenetic factors include parental age, preterm birth, and other environmental factors [118]. Gastrointestinal symptoms and high levels of intestinal immune inflammation linked with gut dysbiosis are common in children with ASD [114,119]. The gut microbiome of ASD patients differs from that of healthy individuals. The increased levels of Lactobacillus and Clostridium have been consistently observed in ASD patients in a few reports [95,96,98-101]. However, controversial results were also observed for Bifidobacterium, Alistipes, Bacteroidetes, and Bacteroidetes in ASD patients [98–100].

NGS of 16S rRNA gene amplicons was used to compare fecal samples from autism patients and healthy controls. The relative abundance of the *Lactobacillaceae*, *Bifidobacteraceae*, and *Veillonellaceae* families was higher in ASD patients than that in healthy controls [94]. For comparison, fecal samples from two groups (Indian and American patients with ASD) having different diets were also analyzed. Despite differences in the diets of Indian and American patients with ASD, *Lactobacillus* species were the most dominant in both groups. *Lactobacillus* is one of the common lactic acid producers present predominantly in the gut of infants and its level reduces with age because of different diets. However, the association between the increased populations of *Lactobacillus* with autism pathology is unclear.

Bacterial tag-encoded FLX-titanium amplicon pyrosequencing (bTEFAP) of the 16S rDNA and 16S rRNA analyses of fecal samples from autism patients and healthy controls were conducted [95]. The microbial alpha-diversity was higher in autism patients. Additionally, autism patients displayed higher levels of the genera *Caloramator*, *Sarcina*, and *Clostridium* and of the *Alistipes* and *Akkermansia* species.

The microbiota composition of fecal samples from autism patients and healthy controls was also analyzed by 16S rRNA gene sequencing [97]. There was no difference in the microbiome diversity, namely species richness and evenness, between autism patients and healthy controls. However, a higher abundance of the genus *Burkholderia* and lower amounts of the genus *Neisseria* were found in autism patients. Moreover, two *Bacteroides* species and *Escherichia coli* were decreased in autism patients.

Compared with control mice, GF mice transplanted with ASD donor feces had increased repetitive behavior, decreased sociability, and decreased mobility [120]. The offspring of GF mice implanted with an ASD fecal sample also displayed ASD-like behaviors. Mice lacking ephrin type-B receptor 6 (EphB6) exhibited ASD-like behaviors as well as microbial dysbiosis and their fecal microbiota contained less *Deferribacteres* [121]. Specific pathogen-free (SPF) C57BL/6J mice transplanted by the fecal microbiota from EphB6-defective mice displayed ASD-like behaviors measure with the open field test, elevated plus maze test, and social behavior tests. On the other hand, ASD-like behaviors were improved in EphB6-defective mice transplanted with fecal microbiota from wild-type mice.

It has been reported that the administration of bacterial metabolites, such as taurine and 5-aminovaleric acid, into an autism mouse model, enhanced ASD-like behavior [120]. Arentsen *et al.* investigated the effect of bacterial peptidoglycan cognitive protein (PGN) in brain development [122]. PGN2-deficient mice showed alteration in the autism risk gene c-Met and social behavior change. Lipopolysaccharides produced by Gram-negative bacteria and lipoproteins and peptidoglycan produced by Gram-positive bacteria are recognized by Toll-like receptors and then stimulate the production of cytokines. Additionally, they are transported to the brain across the blood–brain barrier and are associated with the expression of the ASD risk gene c-Met.

## 3.1.3. Attention deficit hyperactivity disorder (ADHD)

ADHD is a neurodevelopmental disorder characterized by impulsivity, hyperactivity, and inattention. ADHD is usually present from birth and its onset is always before the age of seven. Risk factors of developing ADHD are genetic and environmental [123,124]. The causes of ADHD are unknown. However, the dopamine (DA) transporter gene (DAT1) and the dopamine D4 receptor gene were associated with ADHD [124]. ADHD patients showed a density of DA receptors lower than normal. Recently, the mutations A559V and R615C of DAT-1 were identified in ADHD patients, suggesting a hyperactive DA response [125].

Arts *et al.* examined the gut microbiome of adolescents and adults diagnosed with ADHD. They observed increased levels of bacteria belonging to the *Bifidobacterium* and *Eggerthella* genera in patients with ADHD [126]. They postulated that this increase was associated with enhanced dopamine precursor synthesis, which was associated with changes in reward expectation responses, a hallmark of ADHD [126]. In another study, *Faecalibac*-

*terium* was reduced in children with ADHD [127]. In patients with ADHD, the cytokine level is increased, and there is a positive correlation between the symptoms of ADHD and cytokines [128]. *Faecalibacterium* has an anti-inflammatory effect on the host [129,130], which suggests that a low level of *Faecalibacterium* increases inflammation [131].

GF mice transplanted with fecal microbiota of ADHD participants showed anxiety in the open field test and a decreased Lachnoclostridium content, which was also found in ADHD patients [132]. Pärtty et al. conducted a double-blind randomized placebo-controlled trial in which 75 infants received Lactobacillus rhamnosus GG (ATCC 53103) or placebo during the first 6 months of life. The diagnostic of ADHD or Asperger syndrome (AS) was posed when the children were 13 years old based on the ICD-10 diagnostic criteria. Interestingly, 17.1% of the children in the placebo group were diagnosed with ADHD or AS, whereas none of the children in the probiotics group were diagnosed [133]. Kumperscak et al. investigated the effects of supplementation with the probiotic strain Lactobacillus rhamnosus GG (ATCC53103). The scores obtained in the PedsQLTM Child-Self Report were significantly improved in children and adolescents with ADHD who received probiotics supplementation (n = 18) for 3 months compared with those of placebo controls (n = 14). In addition, the proinflammatory factors IL-12 p70, IL-10, and tumor necrosis factor- $\alpha$  (TNF- $\alpha$ ) were significantly decreased in the probiotics group. The proinflammatory IL-6 was significantly reduced in both groups compared with the levels 3 months before. These controversial results were suggested to be the result of the small sample size and the short observation period [134].

#### 3.1.4. Depression

Depression is a common psychiatric disorder characterized by depressed mood, loss of interest, and loss of pleasure for 2 weeks or longer. It also comes with symptoms such as sleep disorder, loss or gain of weight, and diminished ability to think or concentrate [135]. Depression was associated to abnormalities in neurotransmitters like a decrease in serotonin (5-HT) transporter binding and 5-HT1A receptor binding [136].

The association between depression and gut microbiota has been studied in depression patients. Obvious differences in gut microbiota composition between depressed patients and healthy controls were evidenced using 16S rRNA gene sequencing and shotgun metagenomic analysis. Briefly, the lower abundance of *Allobaculum* has been observed in patients with depression compared with healthy subjects [90,91]. However, a controversial result was also observed for *Bacteroidetes* and *Actinobacteria* [91– 93].

Kelly *et al.* analyzed the composition of human fecal samples from depressed patients and healthy controls by using 16S rRNA sequencing [91]. In depressed patients, the relative proportion of *Prevotellaceae* was decreased, whereas that of *Thermoanaerobacteriaceae* was increased. At the genus level, the relative proportions of *Eggerthella, Holdemania, Gelria, Turicibacter, Paraprevotella,* and *Anaerofilum* were increased in the depressed patients, whereas those of *Prevotella* and *Dialister* were decreased. Moreover, microbiota-depleted rats transplanted with the fecal microbiota from depressed patients developed depression-related behaviors, which suggested that depression was related to the decreased gut microbiota richness and diversity.

An increased abundance of *Actinobacteria* and decreased *Bacteroidetes* amounts were observed in depressed patients compared with the levels in healthy controls [92]. The mice that received fecal samples from depressed patients displayed depression-like behaviors. These results suggest that dysbiosis of the gut microbiome is associated with the development of depression. Naseribafrouei *et al.* showed an association between depression and gut

microbiota: specifically, Oscillibacter and Alistipes were abundant in depressed patients [40]. Jiang *et al.* found that *Enterobacteriaceae* and Alistipes levels were increased in patients with major depressive disorder (MDD), whereas Faecalibacterium was less abundant [93]. Moreover, the alpha-diversity was higher in the active-MDD patients. They suggested that Alistipes species was indole-positive and that it affects tryptophan availability, a precursor of serotonin closely related to depression [93,137,138]. In another study, the analysis of the fecal microbiota of patients with depression revealed the level of *Bifidobacterium* and *Lactobacillus* [139] to have decreased. Bravo et al. indicated that L. rhamnosus had a beneficial effect in treating depression and anxiety in stress-induced mice [140]. Long-term treatment with *L. rhamnosus* decreased corticosterone level and depressive-like behaviors in stress-induced mice. GABA is one of the inhibitory neurotransmitters of the CNS and is associated with the pathogenesis of anxiety and depression. The administration of L. rhamnosus decreased GABA<sub>B1b</sub> mRNA levels in the hippocampus, which was consistent with the antidepressant-like effect of GABA<sub>B</sub> receptor antagonists [140].

Akkasheh et al. conducted a randomized, double-blind, and placebo-controlled clinical trial that included 40 patients with a diagnosis of MDD based on the criteria from the diagnostic and statistical manual of mental disorders, fourth edition (DSM-IV). The patients were given probiotics (Lactobacillus acidophilus, Lactobacillus casei, and Bifidobacterium bifidum) or placebo for 8 weeks. Probiotic supplements had beneficial effects on Beck Depression Inventory total scores, serum insulin levels, homeostasis model assessment of insulin resistance, and serum high sensitivity Creactive protein (hs-CRP) concentrations [141]. Ghorbani et al also investigated the effects of symbiotic supplement on depression [142]. They provided synbiotic supplements to 40 patients (moderate depression and placebo-controlled) who met the DSM-V criteria for moderate depression. All patients received fluoxetine (20 mg/day) for 4 weeks. Either a synbiotic (Lactobacillus casaei, Lactobacillus acidofilus, Lactobacillus bulgarigus, Lactobacillus rhamnosus, Bifidobacterium breve, Bifidobacterium longum, Streptococus thermophilus, fructooligosaccharide as prebiotic) or the placebo was added to the administration for 6 weeks. The synbiotic group obtained a significantly reduced HAM-D score compared with that of the placebo group [142].

However, another study showed no beneficial effect of probiotics supplement on depression. Romijn *et al.* followed seventynine participants who received probiotics (*Lactobacillus helveticus* and *Bifidobacterium longum*) or placebo for 8 weeks [143]. They found no significant changes in depression severity and inflammatory markers in probiotics-supplemented depression patients.

## 3.2. Gut microbiome and neurodegenerative diseases

Researchers have identified a role of the gut microbiome in the development of neurodegenerative diseases such as Alzheimer's, Parkinson's, and amyotrophic lateral sclerosis. The gut microbiome also alters vulnerability to and progress of neurodegenerative diseases. In this section, we discuss the involvement of the microbiome in the pathogenesis of the neurodegenerative diseases.

## 3.2.1. Alzheimer's disease (AD)

AD, one of the major neurodegenerative disorders, is characterized by amyloid plaques and tau tangles. Patients with AD usually exhibit memory impairments and, less frequently, non-amnestic cognitive impairments [144]. Genetic factors, such as rare mutations, are critical risk factors in the development of AD [145].

The gut microbiome of patients with AD differs from that of other individuals. In reports, the decreased abundance of *Bifidobacterium* has been observed in AD patients, and supplementing *Bifidobacterium* alleviated cognitive symptoms of AD patients [63,64]. However, controversial results have been reported for the abundances of *Firmicutes* and *Bacteroidetes* in AD patients [64,66,68].

A study showed differences in bacterial abundance, including reduced Firmicutes and Bifidobacterium levels and increased abundance of Bacteroidetes, in fecal samples obtained from patients with AD using 16S rRNA gene sequencing [64]. Phylum Bacteroidetes includes different Gram-negative commensal bacteria in the gut such as Bacteroides. Gram-negative bacteria have lipopolysaccharide (LPS) that induces inflammation. It also revealed that patients with AD tended to have the lower microbial alpha-, beta-diversity compared with healthy controls. The intestinal absorption of significant LPS and amyloids from bacteria promotes the production of proinflammatory cytokines and is implicated in the pathogenesis of AD [146]. LPS and Gram-negative E. coli fragments were colocalized with an amyloid plaque in a postmortem brain tissue of patients with AD. The abundance of Gram-negative bacteria in the gut may increase LPS translocation, induce inflammation, and then aggravate pathology [64]. In another study, the analysis of fecal sample microbiota using 16S rRNA sequencing exhibited a decreased abundance of Bacteroidaceae and Lachnospiraceae. It increased the abundance of Actinobacteria and Ruminococcaceae in patients with AD compared with controls [65]. Although bacterial metabolites may be associated with AD, additional in-depth microbiome studies are required because the results of microbiome analysis are often controversial [64,65].

The fecal microbiomes and fecal SCFA composition of healthy and AD mice were also compared. The microbiota composition and diversity were perturbed and the levels of SCFAs were decreased in AD mice. At the phylum level, *Proteobacteria* and *Verrucomicrobia* levels were increased in AD mice as revealed by 16S rRNA gene sequencing of fecal samples [67]. At the genus level, *Ruminococcus* and *Butricicocus* were less abundant in AD mice. The findings showed that AD might exacerbate cognitive deficits through the reduction of SCFAs by altering the gut microbiota.

Mice transplanted with feces from AD patients had lower levels of nervous system-related metabolites such as GABA, taurine, and valine compared with those in mice transplanted with healthy human feces. They also displayed significant cognitive impairment [147]. Chen *et al.* demonstrated that mice with AD-linked 5 mutations (5xFAD), a rodent model for AD, experienced gut dysbiosis in an age-dependent manner, and that antibiotic treatment improved cognitive performances and reduced amyloidogenic processes [148]. Moreover, probiotic R13 treatment reduced amyloid aggregates in the gut of 5xFAD mice and boosted the beneficial effects of bacterium *L. salivarius*.

# 3.2.2. Parkinson's disease (PD)

PD is a neurodegenerative disorder characterized by movement dysfunctions. It also has a non-motor dimension that involves cognitive impairments and depression [149]. Misfolding and aggregation of  $\alpha$ -synuclein, mitochondrial dysfunction, malfunctioning protein clearance systems, including the ubiquitin–proteasome and autophagy-lysosome systems, and neuroinflammation, play vital roles in the progression and onset of PD [100]. The dark pigmented areas in the pars compacta of substantia nigra and locus coeruleus are lost in the brains of patients with PD due to the death of dopaminergic neuromelanin-containing neurons and noradrenergic neurons [100].

Several studies have analyzed the link between gut microbiota and PD from fecal samples of PD patients. In a few reports, the increased abundance of *Bifidobacteriaceae*, *Lactobacillaceae* and *Verrucomicrobiaceae* families, and the decreased abundance of *Prevotellaceae* family have been consistently observed in PD patients [72,73,76,79,82]. The higher level of *Lactobacillus* and lower level *Ruminococcus* have been also observed in PD patients compared with that of healthy controls [71,74,77,80]. However, there were controversial results about the abundance of *Faecalibac*-*terium* in PD patients [74,78,80].

Vidal-Martinez et al. observed the increased amounts of several microorganisms (Verrucomicrobiaceae [Akkermansia muciniphila] and unclassified Firmicutes) and decreased levels of other microorganisms (Prevotellaceae [Prevotella copri] and Erysipelotrichaceae [Eubacterium biforme]) in PD patients [72]. Akkermansia muciniphila is a mucin degrader that improves the gut barrier function by restoring the intestinal mucus layer and the underlying epithelium and is considered beneficial for human health [150]. However, Ring, C. et al. showed that A. muciniphila had the opposite outcome: degradation of mucin and increase in intestinal inflammation and intestinal permeability [151]. A. muciniphila is also abundant in constipated individuals, which is one of the major non-motor symptoms in PD [152]. In another study, 16S rRNA gene sequencing of fecal samples of 197 PD patients and 130 healthy controls was conducted. It revealed that the abundances of Bifidobacteriaceae, Lactobacillaceae, Tissierellaceae, Christensenellaceae, and Verrucomicrobiaceae were increased, and the abundances of Lachnospiraceae and Pasteurellaceae were decreased in PD patients [83]. Collectively, Verrucomicrobiaceae, Bifidobacteriaceae, Christensenellaceae, as well as Lactobacillaceae increased and Bacteroidaceae and Prevotellaceae decreased in patients with PD. In contrast, changes in Ruminococcaceae and Lachnospiraceae families were not consistent [65,69-76,78-80,82].

Higher levels of *Proteus* sp., *Bilophila* sp., and *Roseburia* sp. as well as loss of members of the families *Lachnospiraceae*, *Rikenellaceae*, and *Peptostreptococcaceae* were found in GF mice transplanted with PD patient fecal microbiota compared with the levels in GF mice transplanted with healthy control microbiota. GF mice transplanted with PD microbiota showed impairment in behavioral tests for motor dysfunction, such as the beam traversal, pole descent, and nasal adhesive removal tests [153]. The levels of beneficial bacteria such as *Bifidobacterium* increased in M83 transgenic mice treated with adoptive cellular therapy. It was also shown that immunotherapy outcomes had an impact on the gut microbiome of PD patients [154].

## 3.2.3. Amyotrophic lateral sclerosis (ALS)

ALS is a neurodegenerative disease inducing the loss of muscle control. Several studies have been conducted to investigate the association between gut microbiota and ALS. The found a strong link between several bacterial species and ALS.

The analysis of fecal samples from ALS patients and healthy controls with 16S rRNA gene sequencing revealed that *Escherichia coli* and *Enterobacteriaceae* abundance were higher in ALS patients than that in healthy controls. Additionally, *Clostridium* amounts were lower in ALS patients [85]. Blacher *et al.* employed shotgun metagenomic sequencing to analyze gut microbiota and found that *Anaerostipes hadrus* and *Bacteroidales bacterium* were marginally more abundant in ALS patients, whereas the levels of *Bifidobacterium pseudocatenulatum*, *Clostridium leptum*, and *Escherichia coli* were decreased [86]. Zhai *et al.* analyzed the fecal samples of ALS patients and healthy controls by 16S rRNA gene sequencing and found that *Methanobrevibacter* levels tended to be higher in ALS patients compared with those of healthy controls, whereas *Faecalibacterium* and *Bacteroides*, which are known beneficial microorganisms, were less abundant in ALS patients [89].

ALS presents with swallowing disturbance, muscle atrophy, alterations in metabolic activity, and weight loss due to eating problems [155]. Compared with healthy controls, the levels of SCFAs, NO<sub>2</sub>-N/NO<sub>3</sub>-N, and GABA, which are metabolites and substrates of the gut microbiome, were higher in patients with ALS [89]. Wu S *et al.* obtained fecal samples from ALS transgenic mice (G93A) and analyzed bacterial profiles using 16r RNA sequencing.

They showed that G93A mice exhibited increased gut permeability and higher levels of the inflammatory cytokine IL-17. G93A mice also showed a decrease level of in butyrate-producing bacteria (*Butyrivibrio fibrisolvens*) in young mice before the onset of ALS [88]. The changes in *E. coli* abundance and bacterial metabolites are still unclear in ALS. The abundance of identified microorganisms was often inconsistent. Thus, further in-depth research is required to determine the clear association between microbiota and ALS.

## 4. Summary

The advancement in microbiome analysis methods allowed us to understand which microorganism populations and bacterial metabolites were altered in a disease state compared with the normal state. In this review, the advantages and limitations of microbiome analysis methods and online tools for GBA discovery were summarized, such as metabolomic and metagenomic analyses. In addition, the potential relationship between alteration of the gut microbiome and brain disorders reported to date was summarized. These findings have provided researchers with a new insight into the pathophysiology of brain diseases and a new path to explore successful disease treatments based on probiotics or fecal transplantation [156].

## **CRediT authorship contribution statement**

Jae Gwang Song: Writing – original draft. Myeong-Sang Yu: Writing – original draft. Bomi Lee: Writing – original draft. Jingyu Lee: Writing – original draft. Su-Hee Hwang: Writing – original draft. Dokyun Na: Writing – review & editing. Hyung Wook Kim: Writing – review & editing.

### **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

## Acknowledgments

This work was supported by the Chung-Ang University Research Grants in 2021 and the National Research Foundation (NRF) of Korea (NRF- 2018R1A5A1025077 to D Na and NRF-2021R1F1A1059574 To HW Kim).

#### References

- Appleton J. The gut-brain axis: influence of microbiota on mood and mental health. Integr Med (Encinitas, Calif) 2018;17(4):28–32.
- [2] Rhee SH, Pothoulakis C, Mayer EA. Principles and clinical implications of the brain-gut-enteric microbiota axis. Nat Rev Gastroenterol Hepatol 2009;6 (5):306–14.
- [3] Ghosh S, Whitley CS, Haribabu B, Jala VR. Regulation of intestinal barrier function by microbial metabolites. Cell Mol Gastroenterol Hepatol 2021;11 (5):1463–82.
- [4] Lyte M, Varcoe JJ, Bailey MT. Anxiogenic effect of subclinical bacterial infection in mice in the absence of overt immune activation. Physiol Behav 1998;65(1):63–8.
- [5] Goehler LE, Gaykema RP, Opitz N, Reddaway R, Badr N, Lyte M. Activation in vagal afferents and central autonomic pathways: early responses to intestinal infection with Campylobacter jejuni. Brain Behav Immun 2005;19(4):334–44.
- [6] Sudo N, Chida Y, Aiba Y, Sonoda J, Oyama N, Yu XN, et al. Postnatal microbial colonization programs the hypothalamic-pituitary-adrenal system for stress response in mice. J Physiol 2004;558(Pt 1):263–75.
- [7] Azzam I, Gilad S, Limor R, Stern N, Greenman Y. Ghrelin stimulation by hypothalamic-pituitary-adrenal axis activation depends on increasing cortisol levels. Endocr Connect 2017;6(8):847–55.
- [8] Picciotto MR. Galanin and addiction. Cell Mol Life Sci 2008;65(12):1872-9.

- [9] Cheng HY, Ning MX, Chen DK, Ma WT. Interactions between the gut microbiota and the host innate immune response against pathogens. Front Immunol 2019;10:607.
- [10] Dantzer R. Neuroimmune interactions: from the brain to the immune system and vice versa. Physiol Rev 2018;98(1):477–504.
- [11] Braniste V, Al-Asmakh M, Kowal C, Anuar F, Abbaspour A, Toth M, et al. The gut microbiota influences blood-brain barrier permeability in mice. Sci Transl Med 2014;6(263):263ra158.
- [12] Mayer EA, Tillisch K, Gupta A. Gut/brain axis and the microbiota. J Clin Invest 2015;125(3):926–38.
- [13] Breit S, Kupferberg A, Rogler G, Hasler G. Vagus nerve as modulator of the brain-gut axis in psychiatric and inflammatory disorders. Front Psychiatry 2018;9(44):44.
- [14] Morais LH, Schreiber HLt, Mazmanian SK. The gut microbiota-brain axis in behaviour and brain disorders. Nat Rev Microbiol 2021;19(4):241-55.
- [15] Schloss PD, Westcott SL, Ryabin T, Hall JR, Hartmann M, Hollister EB, et al. Introducing mothur: open-source, platform-independent, communitysupported software for describing and comparing microbial communities. Appl Environ Microbiol 2009;75(23):7537–41.
- [16] Bolyen E, Rideout JR, Dillon MR, Bokulich NA, Abnet CC, Al-Ghalith GA, et al. Reproducible, interactive, scalable and extensible microbiome data science using QIIME 2. Nat Biotechnol 2019;37(8):852–7.
- [17] Callahan BJ, McMurdie PJ, Rosen MJ, Han AW, Johnson AJA, Holmes SP. DADA2: high-resolution sample inference from illumina amplicon data. Nat Methods 2016;13(7):581–3.
- [18] McMurdie PJ, Holmes S. Phyloseq: an R package for reproducible interactive analysis and graphics of microbiome census data. PLoS ONE 2013;8(4):e61217.
- [19] Oksanen J, Blanchet FG, Kindt R, Legendre P, Minchin PR, O'hara R, et al. Package 'vegan'. Community ecology package, version 2013;2(9):1-295.
- [20] Love M, Anders S, Huber W, Love MM, BiocGenerics I, Biobase B, et al. Package 'DESeq2'. 2013.
- [21] Peng Y, Leung HCM, Yiu SM, Chin FYL. IDBA-UD: a de novo assembler for single-cell and metagenomic sequencing data with highly uneven depth. Bioinformatics 2012;28(11):1420–8.
- [22] Bankevich A, Nurk S, Antipov D, Gurevich AA, Dvorkin M, Kulikov AS, et al. SPAdes: a new genome assembly algorithm and its applications to single-cell sequencing. J Comput Biol 2012;19(5):455–77.
- [23] Li DH, Luo RB, Liu CM, Leung CM, Ting HF, Sadakane K, et al. MEGAHIT v1.0: a fast and scalable metagenome assembler driven by advanced methodologies and community practices. Methods 2016;102:3–11.
- [24] Beghini F, McIver LJ, Blanco-Miguez A, Dubois L, Asnicar F, Maharjan S, et al. Integrating taxonomic, functional, and strain-level profiling of diverse microbial communities with bioBakery 3. Elife 2021;10:e65088.
- [25] Meyer F, Paarmann D, D'Souza M, Olson R, Glass EM, Kubal M, et al. The metagenomics RAST server-a public resource for the automatic phylogenetic and functional analysis of metagenomes. BMC Bioinform 2008;9:386.
- [26] Douglas GM, Maffei VJ, Zaneveld JR, Yurgel SN, Brown JR, Taylor CM, et al. PICRUSt2 for prediction of metagenome functions. Nat Biotechnol 2020;38 (6):685–8.
- [27] Luo RB, Liu BH, Xie YL, Li ZY, Huang WH, Yuan JY, et al. SOAPdenovo2: an empirically improved memory-efficient short-read de novo assembler. GigaScience 2012;1(1):18.
- [28] Schluenzen F, Tocilj A, Zarivach R, Harms J, Gluehmann M, Janell D, et al. Structure of functionally activated small ribosomal subunit at 3.3 angstrom resolution. Cell 2000;102(5):615–23.
- [29] Wu GD, Lewis JD. Analysis of the human gut microbiome and association with disease. Clin Gastroenterol Hepatol 2013;11(7):774–7.
- [30] Dave M, Higgins PD, Middha S, Rioux KP. The human gut microbiome: current knowledge, challenges, and future directions. Transl Res 2012;160 (4):246-57.
- [31] Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, et al. QIIME allows analysis of high-throughput community sequencing data. Nat Methods 2010;7(5):335–6.
- [32] Cole JR, Wang Q, Fish JA, Chai BL, McGarrell DM, Sun YN, et al. Ribosomal database project: data and tools for high throughput rRNA analysis. Nucleic Acids Res 2014;42(D1):D633–42.
- [33] Suzuki MT, Giovannoni SJ. Bias caused by template annealing in the amplification of mixtures of 16S rRNA genes by PCR. Appl Environ Microbiol 1996;62(2):625–30.
- [34] Walker AW, Martin JC, Scott P, Parkhill J, Flint HJ, Scott KP. 16S rRNA genebased profiling of the human infant gut microbiota is strongly influenced by sample processing and PCR primer choice. Microbiome 2015;3:26.
- [35] Schloss PD, Gevers D, Westcott SL. Reducing the effects of PCR amplification and sequencing artifacts on 16S rRNA-based studies. PLoS ONE 2011;6(12): e27310.
- [36] Knight R, Vrbanac A, Taylor BC, Aksenov A, Callewaert C, Debelius J, et al. Best practices for analysing microbiomes. Nat Rev Microbiol 2018;16(7):410–22.
- [37] Reuter JA, Spacek DV, Snyder MP. High-throughput sequencing technologies. Mol Cell 2015;58(4):586–97.
- [38] Nocker A, Burr M, Camper AK. Genotypic microbial community profiling: a critical technical review. Microb Ecol 2007;54(2):276–89.
- [**39**] Li H, Homer N. A survey of sequence alignment algorithms for nextgeneration sequencing. Brief Bioinform 2010;11(5):473–83.
- [40] Naseribafrouei A, Hestad K, Avershina E, Sekelja M, Linlokken A, Wilson R, et al. Correlation between the human fecal microbiota and depression. Neurogastroenterol Motil 2014;26(8):1155–62.

- [41] Li H, Durbin R. Fast and accurate long-read alignment with Burrows-Wheeler transform. Bioinformatics 2010;26(5):589–95.
- [42] Li ZY, Chen YX, Mu DS, Yuan JY, Shi YJ, Zhang H, et al. Comparison of the two major classes of assembly algorithms: overlap-layout-consensus and debruijn-graph. Brief Funct Genomics 2012;11(1):25–37.
- [43] Ranjan R, Rani A, Metwally A, McGee HS, Perkins DL. Analysis of the microbiome: advantages of whole genome shotgun versus 16S amplicon sequencing. Biochem Biophys Res Commun 2016;469(4):967–77.
- [44] Bashiardes S, Zilberman-Schapira G, Elinav E. Use of metatranscriptomics in microbiome research. Bioinform Biol Insights 2016;10:19–25.
- [45] Shakya M, Lo CC, Chain PSG. Advances and challenges in metatranscriptomic analysis. Front Genet 2019;10:904.
- [46] Chen MX, Wang SY, Kuo CH, Tsai IL. Metabolome analysis for investigating host-gut microbiota interactions. J Formos Med Assoc 2019;118:S10–22.
- [47] Parada Venegas D, De la Fuente MK, Landskron G, González MJ, Quera R, Dijkstra G, et al. Short chain fatty acids (SCFAs)-mediated gut epithelial and immune regulation and its relevance for inflammatory bowel diseases. Front Immunol 2019;10:277.
- [48] Vernocchi P, Del Chierico F, Putignani L. Gut microbiota profiling: metabolomics based approach to unravel compounds affecting human health. Front Microbiol 2016;7:1144.
- [49] Galloway-Pena J, Hanson B. Tools for analysis of the microbiome. Dig Dis Sci 2020;65(3):674–85.
- [50] Zhang X, Figeys D. Perspective and guidelines for metaproteomics in microbiome studies. J Proteome Res 2019;18(6):2370–80.
- [51] The UniProt Consortium. UniProt: the universal protein knowledgebase in 2021. Nucleic Acids Res 2021;49(D1):D480-9.
- [52] Chang A, Jeske L, Ulbrich S, Hofmann J, Koblitz J, Schomburg I, et al. BRENDA, the ELIXIR core data resource in 2021: new developments and updates. Nucleic Acids Res 2021;49(D1):D498–508.
- [53] Kolmeder CA, de Vos WM. Metaproteomics of our microbiome-developing insight in function and activity in man and model systems. J Proteomics 2014;97:3–16.
- [54] Lin H, He QY, Shi L, Sleeman M, Baker MS, Nice EC. Proteomics and the microbiome: pitfalls and potential. Expert Rev Proteomics 2019;16 (6):501–11.
- [55] Quigley EMM. Microbiota-brain-gut axis and neurodegenerative diseases. Curr Neurol Neurosci Rep 2017;17(12):94.
- [56] Neufeld KM, Kang N, Bienenstock J, Foster JA. Reduced anxiety-like behavior and central neurochemical change in germ-free mice. Neurogastroenterol Motil 2011;23(3):255–e119.
- [57] Diaz Heijtz R, Wang S, Anuar F, Qian Y, Bjorkholm B, Samuelsson A, et al. Normal gut microbiota modulates brain development and behavior. P NATL A SCI INDIA A 2011;108(7):3047–52.
- [58] Foster JA, McVey Neufeld KA. Gut-brain axis: how the microbiome influences anxiety and depression. Trends Neurosci 2013;36(5):305–12.
- [59] Ma Q, Xing C, Long W, Wang HY, Liu Q, Wang R. Impact of microbiota on central nervous system and neurological diseases: the gut-brain axis. J Neuroinflamm 2019;16(1):53.
- [60] Rutsch A, Kantsjo JB, Ronchi F. The Gut-brain axis: how microbiota and host inflammasome influence brain physiology and pathology. Front Immunol 2020;11:604179.
- [61] de Vadder F, Mithieux G. Gut-brain signaling in energy homeostasis: the unexpected role of microbiota-derived succinate. J Endocrinol 2018;236(2): R105-8.
- [62] Kountouras J, Boziki M, Gavalas E, Zavos C, Grigoriadis N, Deretzi G, et al. Eradication of Helicobacter pylori may be beneficial in the management of Alzheimer's disease. J Neurol 2009;256(5):758–67.
- [63] Akbari E, Asemi Z, Daneshvar Kakhaki R, Bahmani F, Kouchaki E, Tamtaji OR, et al. Effect of probiotic supplementation on cognitive function and metabolic status in Alzheimer's disease: a randomized, double-blind and controlled trial. Front Aging Neurosci 2016;8:256.
- [64] Vogt NM, Kerby RL, Dill-McFarland KA, Harding SJ, Merluzzi AP, Johnson SC, et al. Gut microbiome alterations in Alzheimer's disease. Sci Rep 2017;7 (1):1–11.
- [65] Zhuang Z-Q, Shen L-L, Li W-W, Fu X, Zeng F, Gui L, et al. Gut microbiota is altered in patients with Alzheimer's disease. J Alzheimer's Dis 2018;63 (4):1337-46.
- [66] Sun BL, Li WW, Wang J, Xu YL, Sun HL, Tian DY, et al. Gut microbiota alteration and its time course in a tauopathy mouse model. J Alzheimers Dis 2019;70(2):399–412.
- [67] Zhang L, Wang Y, Xiayu X, Shi C, Chen W, Song N, et al. Altered gut microbiota in a mouse model of Alzheimer's disease. J Alzheimers Dis 2017;60 (4):1241–57.
- [68] Brandscheid C, Schuck F, Reinhardt S, Schäfer K-H, Pietrzik CU, Grimm M, et al. Altered gut microbiome composition and tryptic activity of the 5xFAD Alzheimer's mouse model. J Alzheimers Dis 2017;56(2):775–88.
- [69] Heintz-Buschart A, Pandey U, Wicke T, Sixel-Doring F, Janzen A, Sittig-Wiegand E, et al. The nasal and gut microbiome in Parkinson's disease and idiopathic rapid eye movement sleep behavior disorder. Mov Disord 2018;33 (1):88–98.
- [70] Aho VTE, Pereira PAB, Voutilainen S, Paulin L, Pekkonen E, Auvinen P, et al. Gut microbiota in Parkinson's disease: temporal stability and relations to disease progression. EBioMedicine 2019;44:691–707.

- [71] Hasegawa S, Goto S, Tsuji H, Okuno T, Asahara T, Nomoto K, et al. Intestinal dysbiosis and lowered serum lipopolysaccharide-binding protein in Parkinson's disease. PLoS ONE 2015;10(11):e0142164.
- [72] Bedarf JR, Hildebrand F, Coelho LP, Sunagawa S, Bahram M, Goeser F, et al. Functional implications of microbial and viral gut metagenome changes in early stage L-DOPA-naïve Parkinson's disease patients. Genome Med 2017;9 (1):1–13.
- [73] Hill-Burns EM, Debelius JW, Morton JT, Wissemann WT, Lewis MR, Wallen ZD, et al. Parkinson's disease and Parkinson's disease medications have distinct signatures of the gut microbiome. Mov Disord 2017;32(5):739–49.
- [74] Petrov V, Šaltykova I, Zhukova I, Alifirova V, Zhukova N, Dorofeeva YB, et al. Analysis of gut microbiota in patients with Parkinson's disease. Bull Exp Biol Med 2017;162(6):734–7.
- [75] Vidal-Martinez G, Chin B, Camarillo C, Herrera GV, Yang B, Sarosiek I, et al. A pilot microbiota study in Parkinson's disease patients versus control subjects, and effects of FTY720 and FTY720-mitoxy therapies in parkinsonian and multiple system atrophy mouse models. J Parkinsons Dis 2020;10(1):185–92.
- [76] Li F, Wang P, Chen Z, Sui X, Xie X, Zhang J. Alteration of the fecal microbiota in North-Eastern Han Chinese population with sporadic Parkinson's disease. Neurosci Lett 2019;707:134297.
- [77] Ren T, Gao Y, Qiu Y, Jiang S, Zhang Q, Zhang J, et al. Gut microbiota altered in mild cognitive impairment compared with normal cognition in sporadic parkinson's disease. Front Neurol 2020;11:137.
- [78] Keshavarzian A, Green SJ, Engen PA, Voigt RM, Naqib A, Forsyth CB, et al. Colonic bacterial composition in Parkinson's disease. Mov Disord 2015;30 (10):1351–60.
- [79] Hopfner F, Kunstner A, Muller SH, Kunzel S, Zeuner KE, Margraf NG, et al. Gut microbiota in Parkinson disease in a northern German cohort. Brain Res 2017;1667:41–5.
- [80] Li W, Wu X, Hu X, Wang T, Liang S, Duan Y, et al. Structural changes of gut microbiota in Parkinson's disease and its correlation with clinical features. Sci China Life Sci 2017;60(11):1223–33.
- [81] Qian Y, Yang X, Xu S, Wu C, Song Y, Qin N, et al. Alteration of the fecal microbiota in Chinese patients with Parkinson's disease. Brain Behav Immun 2018;70:194–202.
- [82] Unger MM, Spiegel J, Dillmann KU, Grundmann D, Philippeit H, Burmann J, et al. Short chain fatty acids and gut microbiota differ between patients with Parkinson's disease and age-matched controls. Parkinsonism Relat D 201;32:66-72.
- [83] Shen Y, Xu J, Li Z, Huang Y, Yuan Y, Wang J, et al. Analysis of gut microbiota diversity and auxiliary diagnosis as a biomarker in patients with schizophrenia: a cross-sectional study. Schizophr Res 2018;197:470–7.
- [84] Zheng P, Zeng B, Liu M, Chen J, Pan J, Han Y, et al. The gut microbiome from patients with schizophrenia modulates the glutamate-glutamine-GABA cycle and schizophrenia-relevant behaviors in mice. Sci Adv 2019;5(2):eaau8317.
- [85] Mazzini L, Mogna L, De Marchi F, Amoruso A, Pane M, Aloisio I, et al. Potential role of gut microbiota in ALS pathogenesis and possible novel therapeutic strategies. J Clin Gastroenterol 2018;52:S68–70.
- [86] Blacher E, Bashiardes S, Shapiro H, Rothschild D, Mor U, Dori-Bachash M, et al. Potential roles of gut microbiome and metabolites in modulating ALS in mice. Nature 2019;572(7770):474–80.
- [87] Fang X, Wang X, Yang SG, Meng FJ, Wang XL, Wei H, et al. Evaluation of the microbial diversity in amyotrophic lateral sclerosis using high-throughput sequencing. Front. Microbiol 2016;7:1479.
- [88] Wu S, Yi J, Zhang YG, Zhou J, Sun J. Leaky intestine and impaired microbiome in an amyotrophic lateral sclerosis mouse model. Physiol Rep 2015;3(4): e12356.
- [89] Zhai CD, Zheng JJ, An BC, Huang HF, Tan ZC. Intestinal microbiota composition in patients with amyotrophic lateral sclerosis: establishment of bacterial and archaeal communities analyses. Chin Med J 2019;132(15):1815–22.
- [90] Aoki-Yoshida A, Aoki R, Moriya N, Goto T, Kubota Y, Toyoda A, et al. Omics studies of the murine intestinal ecosystem exposed to subchronic and mild social defeat stress. J Proteome Res 2016;15(9):3126–38.
- [91] Kelly JR, Borre Y, C OB, Patterson E, El Aidy S, Deane J, et al. Transferring the blues: depression-associated gut microbiota induces neurobehavioural changes in the rat. J Psychiatr Res 2016;82:109-18.
- [92] Zheng P, Zeng B, Zhou C, Liu M, Fang Z, Xu X, et al. Gut microbiome remodeling induces depressive-like behaviors through a pathway mediated by the host's metabolism. Mol Psychiatry 2016;21(6):786–96.
- [93] Jiang H, Ling Z, Zhang Y, Mao H, Ma Z, Yin Y, et al. Altered fecal microbiota composition in patients with major depressive disorder. Brain Behav Immun 2015;48:186–94.
- [94] Pulikkan J, Maji A, Dhakan DB, Saxena R, Mohan B, Anto MM, et al. Gut microbial dysbiosis in indian children with autism spectrum disorders. Microb Ecol 2018;76(4):1102–14.
- [95] De Angelis M, Piccolo M, Vannini L, Siragusa S, De Giacomo A, Serrazzanetti DI, et al. Fecal microbiota and metabolome of children with Autism and pervasive developmental disorder not otherwise specified. PLoS ONE 2013;8 (10):e76993.
- [96] Finegold SM, Molitoris D, Song Y, Liu C, Vaisanen ML, Bolte E, et al. Gastrointestinal microflora studies in late-onset Autism. Clin Infect Dis 2002;35(Suppl 1):S6–S16.
- [97] Kushak RI, Winter HS, Buie TM, Cox SB, Phillips CD, Ward NL. Analysis of the duodenal microbiome in autistic individuals: association with carbohydrate digestion. J Pediatr Gastroenterol Nutr 2017;64(5):e110–6.

- [98] Strati F, Cavalieri D, Albanese D, De Felice C, Donati C, Hayek J, et al. New evidences on the altered gut microbiota in autism spectrum disorders. Microbiome 2017;5(1):24.
- [99] Adams JB, Johansen LJ, Powell LD, Quig D, Rubin RA. Gastrointestinal flora and gastrointestinal status in children with Autism-comparisons to typical children and correlation with Autism severity. BMC Gastroenterol 2011;11 (1):1–13.
- [100] Finegold SM, Dowd SE, Gontcharova V, Liu C, Henley KE, Wolcott RD, et al. Pyrosequencing study of fecal microflora of autistic and control children. Anaerobe 2010;16(4):444–53.
- [101] Parracho HMRT, Bingham MO, Gibson GR, McCartney AL. Differences between the gut microflora of children with autistic spectrum disorders and that of healthy children. J Med Microbiol 2005;54(10):987–91.
- [102] Kong G, Cao KL, Judd LM, Li S, Renoir T, Hannan AJ. Microbiome profiling reveals gut dysbiosis in a transgenic mouse model of Huntington's disease. Neurobiol Dis 2020;135:104268.
- [103] Cowen P, Harrison P, Burns T. Shorter Oxford textbook of psychiatry. (6th ed.). Oxford University Press; 2012.
- [104] Deng C, Dean B. Mapping the pathophysiology of schizophrenia: interactions between multiple cellular pathways. Front Cell Neurosci 2013;7(238):238.
- [105] Romme IAC, de Reus MA, Ophoff RA, Kahn RS, van den Heuvel MP. Connectome disconnectivity and cortical gene expression in patients with Schizophrenia. Biol Psychiat 2017;81(6):495–502.
- [106] Kelly JR, Minuto C, Cryan JF, Clarke G, Dinan TG. The role of the gut microbiome in the development of Schizophrenia. Schizophr Res 2021;234:4–23.
- [107] Steele RD. Blood-brain barrier transport of the alpha-keto acid analogs of amino acids. Fed Proc 1986;45(7):2060-4.
- [108] Bourassa MW, Alim I, Bultman SJ, Ratan RR. Butyrate, neuroepigenetics and the gut microbiome: can a high fiber diet improve brain health? Neurosci Lett 2016;625:56–63.
- [109] Sharma RP, Grayson DR, Gavin DP. Histone deactylase 1 expression is increased in the prefrontal cortex of Schizophrenia subjects: analysis of the National Brain Databank microarray collection. Schizophr Res 2008;98(1-3):111-7.
- [110] Benes FM, Lim B, Matzilevich D, Walsh JP, Subburaju S, Minns M. Regulation of the GABA cell phenotype in hippocampus of schizophrenics and bipolars. Proc Natl Acad Sci USA 2007;104(24):10164–9.
- [111] He Y, Kosciolek T, Tang J, Zhou Y, Li Z, Ma X, et al. Gut microbiome and magnetic resonance spectroscopy study of subjects at ultra-high risk for psychosis may support the membrane hypothesis. Eur Psychiat 2018;53:37–45.
- [112] Dickerson FB, Stallings C, Origoni A, Katsafanas E, Savage CL, Schweinfurth LA, et al. Effect of probiotic supplementation on Schizophrenia symptoms and association with gastrointestinal functioning: a randomized, placebocontrolled trial. Prim Care Companion CNS Disord 2014;16(1):26294.
- [113] Tomasik J, Yolken RH, Bahn S, Dickerson FB. Immunomodulatory effects of probiotic supplementation in Schizophrenia patients: a randomized, placebocontrolled trial. Biomark Insights 2015;10:47–54.
- [114] Severance EG, Gressitt KL, Stallings CR, Katsafanas E, Schweinfurth LA, Savage CLG, et al. Probiotic normalization of Candida albicans in Schizophrenia: a randomized, placebo-controlled, longitudinal pilot study. Brain Behav Immun 2017;62:41–5.
- [115] Okubo R, Koga M, Katsumata N, Odamaki T, Matsuyama S, Oka M, et al. Effect of Bifidobacterium breve A-1 on anxiety and depressive symptoms in Schizophrenia: a proof-of-concept study. J Affect Disorders 2019;245:377–85.
- [116] Lord C, Brugha TS, Charman T, Cusack J, Dumas G, Frazier T, et al. Autism spectrum disorder. Nat Rev Dis Primers 2020;6(1):5.
- [117] American Physhiatric Association. Diagnostic and statistical manual of mental disorders: DSM-5 (5th ed.). American Psychiatric Pub; 2013.
- [118] Lyall K, Croen L, Daniels J, Fallin MD, Ladd-Acosta C, Lee BK, et al. The changing epidemiology of autism spectrum disorders. Annu Rev Publ Health 2017;38(1):81–102.
- [119] Hughes HK, Rose D, Ashwood P. The gut microbiota and dysbiosis in autism spectrum disorders. Curr Neurol Neurosci Rep 2018;18:81.
  [120] Sharon G, Cruz NJ, Kang DW, Gandal MJ, Wang B, Kim YM, et al. Human gut
- [120] Sharon G, Cruz NJ, Kang DW, Gandal MJ, Wang B, Kim YM, et al. Human gut microbiota from autism spectrum disorder promote behavioral symptoms in mice. Cell 2019;177(6):1600–18.
- [121] Li Y, Luo ZY, Hu YY, Bi YW, Yang JM, Zou WJ, et al. The gut microbiota regulates autism-like behavior by mediating vitamin B6 homeostasis in EphB6-deficient mice. Microbiome 2020;8(1):120.
- [122] Arentsen T, Qian Y, Gkotzis S, Femenia T, Wang T, Udekwu K, et al. The bacterial peptidoglycan-sensing molecule Pglyrp2 modulates brain development and behavior. Mol Psychiatry 2017;22(2):257–66.
- [123] Faraone SV, Asherson P, Banaschewski T, Biederman J, Buitelaar JK, Ramos-Quiroga JA, et al. Attention-deficit/hyperactivity disorder. Nat Rev Dis Primers 2015;1(1):15020.
- [124] Ougrin D, Chatterton S, Banarsee R. Attention deficit hyperactivity disorder (ADHD): review for primary care clinicians. London J Prim Care (Abingdon) 2010;3(1):45–51.
- [125] Sharma A, Couture J. A review of the pathophysiology, etiology, and treatment of attention-deficit hyperactivity disorder (ADHD). Ann Pharmacother 2014;48(2):209–25.

- [126] Aarts E, Ederveen THA, Naaijen J, Zwiers MP, Boekhorst J, Timmerman HM, et al. Gut microbiome in ADHD and its relation to neural reward anticipation. PLoS ONE 2017;12(9):e0183509.
- [127] Jiang H, Zhou Y, Zhou G, Li Y, Yuan J, Li X, et al. Gut microbiota profiles in treatment-naive children with attention deficit hyperactivity disorder. Behav Brain Res 2018;347:408–13.
- [128] Mitchell RHB, Goldstein BI. Inflammation in children and adolescents with neuropsychiatric disorders: a systematic review. J Am Acad Child Psy 2014;53(3):274–96.
- [129] Quevrain E, Maubert MA, Michon C, Chain F, Marquant R, Tailhades J, et al. Identification of an anti-inflammatory protein from Faecalibacterium prausnitzii, a commensal bacterium deficient in Crohn's disease. Gut 2016;65(3):415–25.
- [130] Qiu X, Zhang M, Yang X, Hong N, Yu C. Faecalibacterium prausnitzii upregulates regulatory T cells and anti-inflammatory cytokines in treating TNBS-induced colitis. J Chrons Colitis 2013;7(11):e558–68.
- [131] Bull-Larsen S, Mohajeri MH. The potential influence of the bacterial microbiome on the development and progression of ADHD. Nutrients 2019;11(11):2805.
- [132] Tengeler AC, Dam SA, Wiesmann M, Naaijen J, van Bodegom M, Belzer C, et al. Gut microbiota from persons with attention-deficit/hyperactivity disorder affects the brain in mice. Microbiome 2020;8(1):44.
- [133] Partty A, Kalliomaki M, Wacklin P, Salminen S, Isolauri E. A possible link between early probiotic intervention and the risk of neuropsychiatric disorders later in childhood: a randomized trial. Pediatr Res 2015;77 (6):823–8.
- [134] Kumperscak HG, Gricar A, Ulen I, Micetic-Turk D. A pilot randomized control trial with the probiotic strain Lactobacillus rhamnosus GG (LGG) in ADHD: children and adolescents report better health-related quality of life. Front Psychiatry 2020;11:181.
- [135] Vlainic JV, Suran J, Vlainic T, Vukorep AL. Probiotics as an adjuvant therapy in major depressive disorder. Curr Neuropharmacol 2016;14(8):952–8.
- [136] Kaltenboeck A, Harmer C. The neuroscience of depressive disorders: a brief review of the past and some considerations about the future. Brain Neurosci Adv 2018;2: 2398212818799269
- [137] Saulnier DM, Riehle K, Mistretta TA, Diaz MA, Mandal D, Raza S, et al. Gastrointestinal microbiome signatures of pediatric patients with irritable bowel syndrome. Gastroenterology 2011;141(5):1782–91.
- [138] Song Y, Kononen E, Rautio M, Liu C, Bryk A, Eerola E, et al. Alistipes onderdonkii sp. nov. and Alistipes shahii sp. nov., of human origin. Int J Syst Evol Micr 2006;56(8):1985–90.
- [139] Aizawa E, Tsuji H, Asahara T, Takahashi T, Teraishi T, Yoshida S, et al. Possible association of Bifidobacterium and Lactobacillus in the gut microbiota of patients with major depressive disorder. J Affect Disorders 2016;202:254–7.
- [140] Bravo JA, Forsythe P, Chew MV, Escaravage E, Savignac HM, Dinan TG, et al. Ingestion of Lactobacillus strain regulates emotional behavior and central GABA receptor expression in a mouse via the vagus nerve. Proc Natl Acad Sci USA 2011;108(38):16050–5.
- [141] Akkasheh G, Kashani-Poor Z, Tajabadi-Ebrahimi M, Jafari P, Akbari H, Taghizadeh M, et al. Clinical and metabolic response to probiotic administration in patients with major depressive disorder: a randomized, double-blind, placebo-controlled trial. Nutrition 2016;32(3):315–20.
- [142] Ghorbani Z, Nazari S, Etesam F, Nourimajd S, Ahmadpanah M, Jahromi SR. The effect of synbiotic as an adjuvant therapy to fluoxetine in moderate Depression: a randomized multicenter trial. Arch Neurosci 2018;5(2): e60507.
- [143] Romijn AR, Rucklidge JJ, Kuijer RG, Frampton C. A double-blind, randomized, placebo-controlled trial of Lactobacillus helveticus and Bifidobacterium longum for the symptoms of depression. Aust NZ J Psychiat 2017;51 (8):810–21.
- [144] Knopman DS, Amieva H, Petersen RC, Chetelat G, Holtzman DM, Hyman BT, et al. Alzheimer disease. Nat Rev Dis Primers 2021;7(1):33.
- [145] Haque RU, Levey AI. Alzheimer's disease: a clinical perspective and future nonhuman primate research opportunities. Proc Natl Acad Sci USA 2019;116 (52):26224–9.
- [146] Pasinetti GM, Ho L. Role of grape seed polyphenols in Alzheimer's disease neuropathology. Nutr Diet Suppl 2010;2010(2):97-103.
- [147] Fujii Y, Nguyen TTT, Fujimura Y, Kameya N, Nakamura S, Arakawa K, et al. Fecal metabolite of a gnotobiotic mouse transplanted with gut microbiota from a patient with Alzheimer's disease. Biosci Biotechnol Biochem 2019;83 (11):2144–52.
- [148] Chen C, Ahn EH, Kang SS, Liu X, Alam A, Ye K. Gut dysbiosis contributes to amyloid pathology, associated with C/EBPbeta/AEP signaling activation in Alzheimer's disease mouse model. Sci Adv 2020;6(31):eaba0466.
- [149] Poewe W, Seppi K, Tanner CM, Halliday GM, Brundin P, Volkmann J, et al. Parkinson disease. Nat Rev Dis Primers 2017;3(1):17013.
  [150] Reunanen J, Kainulainen V, Huuskonen L, Ottman N, Belzer C, Huhtinen H,
- [150] Reunanen J, Kainulainen V, Huuskonen L, Ottman N, Belzer C, Huhtinen H, et al. Akkermansia muciniphila adheres to enterocytes and strengthens the integrity of the epithelial cell layer. Appl Environ Microbiol 2015;81 (11):3655–62.
- [151] Ring C, Klopfleisch R, Dahlke K, Basic M, Bleich A, Blaut M. Akkermansia muciniphila strain ATCC BAA-835 does not promote short-term intestinal inflammation in gnotobiotic interleukin-10-deficient mice. Gut Microbes 2019;10(2):188–203.

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- [152] Romano S, Savva GM, Bedarf JR, Charles IG, Hildebrand F, Narbad A. Meta-analysis of the Parkinson's disease gut microbiome suggests alterations linked to intestinal inflammation. NPJ Parkinsons Dis 2021;7(1):27.
  [153] Sampson TR, Debelius JW, Thron T, Janssen S, Shastri GG, Ilhan ZE, et al. Gut microbiota regulate motor deficits and neuroinflammation in a model of Parkinson's disease. Cell 2016;167(6):1469-80.e12.
  [154] Klann E, Tagliamonte MS, Ukhanova M, Mai V, Vedam-Mai V. Gut microbiota dynamics in parkinsonian mice. ACS Chem Neurosci 2020;11(20):3267-76.

- [155] Ioannides ZA, Ngo ST, Henderson RD, McCombe PA, Steyn FJ. Altered metabolic homeostasis in amyotrophic lateral sclerosis: mechanisms of energy imbalance and contribution to disease progression. Neurodegener Dis
- 2016;16(5-6):382-97.
  [156] Evrensel A, Ceylan ME. Fecal microbiota transplantation and its usage in neuropsychiatric disorders. Clin Psychopharm Neurosci 2016;14(3):231-7.