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The Gut Microbiota and Dysbiosis in Autism Spectrum Disorders

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Abstract

Purpose of Review—There is a growing body of evidence indicating the gut microbiota influence neurodevelopment and behavior. The purposes of this review are to provide an overview of studies analyzing the microbiota and their metabolites in autism spectrum disorders (ASD) and to discuss the possible mechanisms of action involved in microbial influence on the brain and behavior.

Recent Findings—The microbiota-gut-brain (MGB) axis has been extensively studied in animal models, and it is clear that alterations in the composition of microbiota alter neurological and behavioral outcomes. However, findings in human studies are less abundant. Although there are several studies so far showing altered microbiota (dysbiosis) in ASD, the results are heterogeneous and often contradictory. Intervention studies such as fecal microbiota transplant therapies show promise and lend credence to the involvement of the microbiota in ASD.

Summary—A role for the microbiota in ASD is likely; however, further studies elucidating microbial or metabolomic signatures and mechanisms of action are needed. Future research should focus on intervention studies that can identify specific metabolites and immune mediators that improve with treatment to help identify etiologies and pathological mechanisms of ASD.

Keywords

Autism; Microbiota; Dysbiosis; Dysregulation; Neurodevelopment; Behavior

Introduction

Autism spectrum disorders (ASD) are a heterogeneous group of neurodevelopmental disorders characterized by deficits in communication, social interaction, and cognition [1]. ASD have increased significantly in prevalence since they were first identified to a current rate of 1:59, with a preponderance toward boys [1]. No single etiology of ASD has been identified, with current theories suggesting both genetic and environmental contributions. Genetic mechanisms account for approximately 10–20% of ASD cases [2], leaving us with questions about what might be driving etiology of these disorders. Environmental factors

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that have been implicated in increased risk for having a child with ASD include air pollution, pesticide exposure, maternal infections, and/or inflammatory conditions, or antibiotics during pregnancy [3–5]. Immune dysfunction and gastrointestinal (GI) inflammation are also common in individuals with ASD and contribute to severity of behaviors seen in the disorder [6–8]. New data on the role of the gut microbiome in neurodevelopmental disorders has prompted theories of the roles commensal bacteria may play in ASD. The studies discussed in this review have identified significant dysbiosis in children with ASD; however, it is unknown whether the GI dysfunction and dysbiosis is sequelae of the larger disorder or whether they directly contribute to causing ASD.

The gut microbiota consists of the collection of microbes present within the human GI tract, and their collective genome is the gut microbiome [9]. It is generally understood that initial colonization begins at birth through the acquisition of maternal microbiota during vaginal delivery; however, recent research suggests that there may be acquisition of maternal microbiota during gestation [10]. The infant microbiota are supported with breastmilk which is high in human oligosaccharides; however, the composition of early-life microbiota can be altered by delivery methods, hygiene, and feeding practices such as formula feeding [10]. The healthy infant gut is initially dominated by *Bifidobacterium* and *Lactobacillus*; however, it is unstable for the first several years of life, throughout weaning and food introduction, then stabilizing to a more “adult-like” composition around age 3 [10]. The dominant phyla of bacteria in the healthy adult gut are *Firmicutes* and *Bacteroidetes*, with a smaller portion of the microbiota made up of *Actinobacteria*, *Proteobacteria*, and *Verrucomicrobia* [9]. Within the last decade, our understanding of the microbiota and its importance to health has blossomed, and we now recognize that loss or alteration of microbiota may be leading to chronic diseases [11]. Disturbances of the microbiota can occur through antibiotic treatment, changes in diet, immune challenges, and stress [10], and this can upset the balance between beneficial commensals and potentially pathogenic microbes in the gut. This broken balance is termed *dysbiosis* [12].

Microbiota-Gut-Brain Axis

The gut microbiota play a critical role in the development of the intestinal architecture and mucosal immune system and are particularly important for regulation of the immune system in the gut [reviewed in [13, 14]]. In germ-free (GF) mice, the architecture of the mesenteric lymph nodes and Peyer's patches is smaller and disorganized, with reduced maturation of isolated lymphoid follicles and fewer numbers of immune cells present. These mice experience immune dysfunction and are more susceptible to infections than mice kept in conventional facilities [13, 15]. These deficits can be corrected by colonization with diverse commensal bacteria; however, this has been shown to be age-dependent, suggesting that both composition and timing of colonization are critical for the education of the immune system [16–18].

The microbiota also play important roles in digestion, nutrient assimilation, vitamin production, and metabolism [19, 20] and have recently been shown to have significant influence on the bi-directional signaling that takes place between the gut and the nervous system, termed the microbiota-gut-brain axis [21]. The gut microbiota can influence brain

function and lead to changes in behaviors, as shown extensively in animal models [22•]. For example, GF rodent studies have shown that microbiota can positively impact stress responses through the hypothalamus-pituitary-adrenal axis (Fig. 1) [23], such as decreased anxiety and increased exploratory behaviors [24–27]. Probiotics have also been shown to increase exploratory behavior in rodents under varied conditions [28–30] including probiotic treatment after dextran sodium sulfate (DSS)-induced colitis [31]. Conversely, stress in early life can induce dysbiosis and influence immune responses, as shown in maternal separation models of stress [32]. Deficits in working memory occur in GF mice [33], and emotional behavior and memory can be modulated through the administration of probiotics [34]. The introduction of probiotics influences gene expression in the central nervous system, specifically altering expression of γ -aminobutyric acid (GABA) receptors in the hippocampus, mediated through stimulation of the vagus nerve [34]. Microbiota have a profound influence over the synthesis and metabolism of serotonin [26, 35•]. Microbiota produce other neuroactive metabolites including GABA by *Lactobacillus* and *Bifidobacteria* [36], acetylcholine by *Lactobacillus*, dopamine by *Bacillus* and *Serratia*, and norepinephrine by *Escherichia* and *Saccharomyces*, which could enter circulation and directly affect neural processes throughout the body, including the brain (Fig. 1) [37].

The microbiota can also alter the integrity of the intestinal and the blood-brain barriers, specifically due to their production of short-chain fatty acids [38•, 39], and disrupted barrier function could facilitate the translocation of bacterial metabolites and immune mediators from the gut into circulation (Fig. 1), which could lead to activation of microglia as seen in an induced dysbiosis mouse model [40]. Through such animal studies, evidence is mounting that dysbiosis or altered microbiota composition has an impact on neurodevelopment and behavior.

Dysbiosis and impaired intestinal barrier function were seen in a maternal immune activation (MIA) model of autism, with improvements in ASD relevant behavior after treatment with *Bacteriodes fragilis* [41]. Recently, this model has been shown to be dependent on microbiota that can induce a T helper (T_H-) 17 response—dams that lacked certain microbiota could not mount this response, and their offspring did not display the ASD-like behaviors [42•, 43••]. Other mouse models of ASD such as the BTBR and Shank3 knock-out mice have shown intrinsic dysbiosis that correlated with the behavioral phenotypes seen in these models [44, 45]. The latter study saw differences specifically in males, with significant alterations in GABA receptor gene expression in the Shank3 KO mice that could be modulated with the introduction of the probiotic *Lactobacillus reuteri* [45]. In addition, dysbiosis introduced in conventionally housed adult mice also leads to abnormal behaviors, altered barrier functions, and activation of the microglia, the resident immune cells of the brain [40].

Microbiota in ASD

Evidence of microbial dysbiosis in ASD has been growing in recent years (Table 1). In addition to immune and GI dysfunction that may be linked to dysbiosis, there is some evidence that altering the microbiota in ASD can improve behaviors [46, 79•]. The earliest studies investigating the relationship of the microbiota and children with ASD proposed that

excessive antibiotic use led to an overgrowth of spore-forming *Clostridium*, which researchers hypothesized might be exposing these children to high levels of microbial metabolites that were neurotoxic. This hypothesis was based on an index case where regressive autism appeared to coincide with several rounds of antibiotic use, and subsequent vancomycin treatment significantly improved aspects of behavior. A small pilot study included 11 children with a regressive form of ASD and who had GI symptoms of diarrhea that were treated with oral vancomycin for 8 weeks, followed by 4 weeks of probiotics. Eight of these children (~73%) experienced significantly improved impairments in social behavior and communication. However, these improvements did not endure, with most children reverting to their starting behavioral impairments upon removal of the treatment [46]. Assessment of the microbiota in this study found increased *Clostridia* (supporting the hypothesis) as well as overgrowth of other spore-forming anaerobes, microaerophilic bacteria, and several *Clostridia* species within the gastric and duodenal secretions not seen in controls [47]. Further classification of the microbiota identified increased *Clostridia* clusters I/IX and *C. bolteae* (46-fold increase) [48].

These early findings, coupled with the noteworthy numbers of ASD children with gastrointestinal complaints and immune dysfunction, prompted additional research to determine if the microbiota were consistently altered in ASD. Using fluorescence in situ hybridization (FISH) techniques to identify bacteria present in stool samples, Parracho et al. confirmed increased *Clostridia*, and its presence highly correlated with GI symptoms in ASD children [49]. However, as studies designed to use more in-depth techniques were published, other bacterial species were identified as having differential abundance in ASD. Increased *Desulfovibrio* and *Bacteroides vulgatus* were identified in children with ASD and were correlated to autism severity [50]. *Desulfovibrio* was also elevated in a small cohort of Slovakian children with ASD and GI dysfunction compared to healthy controls [76]. *Desulfovibrio* could be an important contributor to GI inflammation, as its major metabolic byproduct—hydrogen sulfide—is cytotoxic to colonic epithelial cells [81]. Moreover, when given to rodents, *Desulfovibrio* decreased working memory [82].

Ileal and cecal biopsies from children with ASD and GI dysfunction showed increased *Firmicutes* and decreased *Bacteroidetes* [51]. As part of this study, the researchers looked at gene expression associated with carbohydrate digestion and transport and found impaired expression of these genes correlated with dysbiosis in ASD [51]. Dysfunctional carbohydrate digestion could alter the fermentation byproducts of the microbiota present, and undigested carbohydrate could preferentially select for certain bacteria [51]. Kang et al. found that children with ASD and GI dysfunction had decreased in commensals important for carbohydrate fermentation including *Prevotella*, *Coprococcus*, and unclassified *Veillonellaceae* [74]. A more recent biopsy study examined duodenal samples of ASD children with GI dysfunction to determine if disaccharidase activity was altered compared to controls, similar to the previous study. Although they did not see overall differences in diversity, they identified elevations in *Burkholderia* and reduced *Prevotella* and *Neisseria* in the duodenum of ASD children. Overall, they did not see the same reductions in disaccharidases; however, they found a correlation between disaccharidase activity with the presence of *Clostridium* [77].

Increased *Sutterella*, a mucosa-associated microbe, was found in significant numbers in intestinal biopsies of ASD children with GI dysfunction [56]. *Sutterella* was also increased in stool samples of ASD children, irrespective of GI issues [80]. Increased *Ruminococcus torques* [80] was also seen in ASD and is similar to dysbiosis noted in inflammatory bowel disorders (IBD) [71]. As well as increases in some species, reductions in important commensals *Bifidobacteria* and the mucin-degrading *Akkermansia muciniphila* [52] have been shown in ASD. One of the largest of recent studies that looked at 40 ASD (36 severe) and 40 typically developing control children found decreased abundance of the *Alistipes*, *Bilophila*, *Dialister*, *Parabacteroides*, and *Veillonella* families but increased *Collinsella*, *Corynebacterium*, *Dorea*, and *Lactobacillus*, suggesting major changes in the microbiota composition in ASD. The authors also found that constipation in ASD correlated with increases in *Escherichia/Shigella* and *Clostridium* cluster XVIII, a cluster known to produce exotoxins that are pro-inflammatory [78].

Two recent studies have provided evidence to support the hypothesis that children with ASD and GI dysfunction have elevated inflammatory immune responses and associated dysbiosis [57, 83]. Notably, the balance of inflammatory cytokines was skewed in children with ASD and GI dysfunction when compared with regulatory cytokines such as transforming growth factor (TGF) beta 1 [57]. However, whether it is the push towards an inflammatory environment that influences the microbiota composition in ASD or a bacterial composition that elicits inflammation needs further study. Inflammatory conditions in the GI tract are known to exacerbate dysbiosis, for example, during antibiotic-associated inflammation, increases in host-derived nitrate encourage *Proteobacteria* such as *Escherichia coli* to bloom [84, 85]. Differences in microbiota diversity, both at the family level and when comparing functional KEGG pathways, were also seen in children with ASD compared to typically developing controls. Interestingly, altered zonulin levels were seen in ASD children with GI dysfunction and may suggest increased intestinal permeability [57]. Luna et al. also showed that increased cytokines associated with mucosal immunity, including interleukin [IL]6, IL-1, IL-17A, and interferon (IFN)-gamma, were associated with abdominal pain and increased *Clostridiales* in ASD [83]. A recent clinical trial which involved fecal microbiota transplant (FMT) in 18 ASD children led to significantly improved GI symptoms and ASD-relevant behaviors, with increases in *Bifidobacterium*, *Prevotella*, and *Desulfovibrio*. These changes persisted at the 8-week follow-up [79]. Since immune dysfunction and cytokine dysregulation are so prominent in ASD [reviewed in [6, 8, 86], measuring immune responses and inflammatory cytokines in future probiotic and FMT clinical trials could help identify the role that dysbiosis plays in this dysfunction.

Overall, there does not seem to be consensus for differences in microbiota in ASD, and a meta-analysis of 15 studies could not amalgamate the often contradictory results [87]. Some studies found increased diversity of the microbiota in ASD [47, 64], whereas other studies show the opposite trend towards decreased diversity and richness [73, 74], or no differences in diversity [75, 77]. Several studies found an increased *Firmicutes/Bacteroidetes* ratio in both stool and biopsy samples, including one study that showed probiotic treatment reversed this trend [51, 76, 78]. However, other studies noted an opposite trend in stool [50, 64]. Tissue/specimen type and sample site may account for the differences in studies. For example, when fecal samples are analyzed, this will only detect what has moved through the

GI tract, not necessarily the dominant colonizers of the varied regions throughout the GI tract. This may skew results to those species that have difficulty adhering to the intestinal epithelium or species that are not surviving well within the GI tract and may explain some of the opposing trends seen when comparing studies using stools versus biopsy specimens [51]. Other technical issues may relate to comparing profiles to siblings who may also exhibit broader behavioral impairments rather than typically developing controls. Siblings are also more likely to have shared a common environment that would alter the microbiota [55, 75]. Additional sources of error include differing techniques; for example, sequencing can vary in error profile, and results can vary based on read length and sequencing depth [88]. Heterogeneity in age of subjects and sex may also reduce comparability across studies. Future studies should investigate microbiota profiles at earlier time points pre-diagnoses of ASD to help determine if there are differences between microbiota and how that relates to ASD outcome or broader neurodevelopmental concerns such as anxiety and cognition. Note that the composition of microbiota is very sensitive to changes diet, fiber composition, etc., and can also be altered by stress and other compounding factors [10]. Children with ASD are known to have feeding problems, food sensitivities/aversions, and extremely restricted diets [89], which could also be contributing to differences in microbiota but not to an ASD diagnosis. A preliminary study showed that despite no differences in the composition of microbiota, early-life probiotic treatment and presence of *Bifidobacterium* during infancy reduced the risk of neurodevelopmental disorders [90]. This early period of colonization is an area needing further research with larger studies and advanced technological tools.

Bacteria are not the only microbes to make up the microbiota. New techniques are identifying the fungal microbiota and its role in human health [91]. GI overgrowth of fungal species such as *Candida* may have deleterious immune consequences and have been linked to IBD and celiac disease [92–94]. Fungal commensals can bloom after antibiotic administration, and the presence of *Candida* can interfere with reassembly of the microbiota after antibiotic perturbation, contributing to dysbiosis [95]. The immune system responds to fungal infections with a T_H-17 response, producing IL-17, a cytokine recently implicated in ASD etiology in the MIA mouse model [42•, 43••]. So far, fungal microbiota studies in ASD are scant; however, one culture-based study showed significant presence of *Candida* species in the feces of children with ASD, the majority of which were *Candida albicans*. Fluconazole-resistant species were also found in significant numbers of cultures, including *C. krusei* and *C. glabrata* [96]. A separate study found *Candida* present in nearly 60% of ASD samples, with none present in controls. They also identified hyphae formation, suggesting that the dimorphic yeast had switched to its invasive and adhesive form [97]. However, one culture-based study looked for the presence of yeast in ASD children and did not see an over-representation compared to control samples [53]. Newer studies that utilize sequencing techniques may be more reliable and have shown that *Candida* is the most abundant taxa of mycobiota seen in children with ASD, found at nearly twice the rate of typically developed children [78]. Elevated ratios of the urinary metabolites D-arabinitol/L-arabinitol (DA/LA) have been used for early identification of invasive candidiasis, as d-arabinitol is a major metabolite of pathogenic fungal species [98]. A preliminary study of 22 children with ASD and GI dysfunction found that DA concentrations and DA/LA ratios were significantly reduced with daily administration of probiotics, and improvements were

seen in certain ASD behaviors including ability to concentrate [62]. Elevated DA (listed as D-arabitol) was also found in a 2014 study of 21 Italian children with ASD [67]. This group also saw elevated glycolic acid, which may also be associated with overgrowth of yeast in the GI tract [67].

Microbial Metabolites in ASD

Due to the many limitations of microbiota analysis, some researchers have turned to metabolomic tools to identify how byproducts of microbial fermentation and metabolism might be interacting with human health or influencing disease. These tools identify altered patterns of metabolites within urine, stool, and blood samples that may provide a biochemical signature of ASD and supportive evidence of dysbiotic gut microbiota [99]. A summary of metabolomics studies specific to the microbiota and ASD are listed in Table 2.

Altered patterns of bacterial metabolites were seen in a large number of children with ASD compared to healthy control children. These metabolites included elevated 3-(3-hydroxyphenyl)-3-hydroxypropionic acid (HPPHA), a catabolic byproduct of *Clostridia* [59]. Increased HPPHA was found more recently in study of Italian children with ASD [67], and a 2016 intervention study showed that Vancomycin treatment could reverse increases of HPPHA and two associated metabolites [69]. Reduced hippurate, phenylacetylglutamine, and p-cresylsulfate were seen in ASD children, indicating alterations in gut microbiota [58]. P-cresol, another microbial metabolite produced by *Clostridium* species, was significantly elevated in children with ASD, especially females with severe autism [60]. Further studies showed that elevations of urinary p-cresol and other derivative metabolites of *Clostridia* were associated with repetitive behaviors [65] and constipation in ASD [70]. In animal models of ASD, elevations of microbial metabolite 4-ethylphenylsulfate (4EPS) correlated with altered behaviors in mice, and probiotic therapy reduced its concentration and improved behaviors in MIA model of ASD [41]. 4EPS is chemically related to p-cresol, the uremic toxin seen elevated in ASD children [41, 60, 65].

Short-chain fatty acids (SCFA) are microbial metabolites of fiber fermentation and can be found in high concentrations in the colon. SCFA metabolites of commensal microbiota are generally beneficial to the host; however, some SCFAs such as propionic acid (PPA) can be neurotoxic in higher concentrations and have been shown to cause behavioral abnormalities in rodent models [100, 101] (Fig. 1). SCFA influence gene expression epigenetically through histone deacetylase (HDAC) inhibition. For example, butyrate-producing bacteria promote peripheral regulatory T cell (T_{regs}) expansion, altering gut immunity by promoting tolerance. This occurs through HDAC inhibition leading to increased acetylation at the Foxp3 promoter and expansion of $CD4^+ Foxp3^+ T_{regs}$ [102••]. Therefore, shifts in SCFA production could potentially contribute to altered immune regulation in the gut and lead to peripheral inflammation (Fig. 1). Several studies found lower levels of fecal SCFA, including butyrate in feces of ASD children [53, 64], while a 2012 study found increased ammonia and SCFA including acetic, butyric, isobutyric, valeric, and isovaleric acids in ASD [61]. Lower SCFA occurred alongside elevated phenol, 4-(1,1-dimethylethyl)-phenol and p-cresol, and correlated with increased *Bacteroides* and *Clostridia* and decreased

Ruminococcoceae [64]. These children also had elevations in free amino acids, indicating increases in proteolytic bacteria [64].

Tryptophan (Trp) metabolites directly influence host physiology including immune and gut homeostasis [reviewed in: [103]]. The gut microbiota are critical for regulating Trp metabolism, either directly by degrading Trp to indole-derivatives or indirectly through mammalian kynurenine and serotonin pathways. GF mice have increased levels of circulating Trp, adding evidence to the importance of the microbiota for Trp metabolism [26, 104]. Metabolome studies have repeatedly shown evidence of alterations in tryptophan, kynurenine, and serotonin pathways in children with ASD, including reports of increased urinary excretion of tryptophan and associated metabolites [63, 67, 68, 72, 105]. The primary metabolic pathway of tryptophan is the kynurenine pathway, leading to production of kynurenic acid. Increases in an alternative branch of the kynurenine pathway led to reduced kynurenic acid and elevated quinolinic acid (QA), a compound known to be an excitotoxic N-methyl-D-aspartate (NMDA) receptor agonist, in children with ASD [68] (Fig. 1). Immune activation may be responsible for this elevation, as activated macrophages and microglia are the main producers of QA [106]. Bacterial degradation of tryptophan, yielding increased indolyl 3-acetic acid, indolyl lactate, and other indole derivatives were also shown in ASD and these shifts were associated with reduced urinary melatonin, downstream of serotonin production that may be the result of bacterial metabolism of available tryptophan [68]. In 2014, a meta-analysis of 22 studies that measured alterations in blood serotonin concluded that there was significantly elevated blood and platelet-rich plasma 5-HT in ASD individuals compared to controls [66]. Serotonergic metabolites were also increased with a decrease in Trp in mucosal biopsies of children with ASD and functional GI disorders, and these findings strongly correlated with several specific bacteria (Table 2) [83]. Spore-forming gut microbiota mediate the production of peripheral serotonin (5-HT) from Trp through the production of metabolites that increase expression of tryptophan hydroxylase (Tph)1 in enterochromaffin (EC) cells of the gut which in turn increases colonic and circulating levels of 5-HT [35]. Serotonin expressed in enteric neurons early in development can also contribute to the 5-HT pool and motility of the GI tract [107]. GF mouse models have shown that the serotonergic system in the brain is also influenced by the microbiota [24, 26], whereby excess peripheral Trp may cross the BBB to influence the rate serotonin synthesis in the brain [108]. Serotonin plays important neurotrophic roles during early development, and elevations of 5-HT during critical time periods can alter cognition and sensory processing [109].

Microbial metabolites and inflammatory mediators are capable of signaling through the vagus nerve [110] (Fig. 1). As the major nerve of the parasympathetic nervous system, it has a bidirectional role in enervating organs throughout the body including the gut. The majority of the fibers of the vagus nerve are afferent sensory fibers, delivering sensory information from the periphery, including the GI tract, to the brain [110]. It is understood that neural communication between the brain and the microbiota may be occurring indirectly through hormones or neurotransmitter release by gut endocrine cells in presence of the microbiota and metabolites, or perhaps through direct pattern recognition sensing due to Toll-like receptor expression on afferent fibers [110]. Vagal activation by the microbiota has been shown in several animal models: vagotomy caused reversal of probiotic effects on memory

and hippocampal GABA receptor gene expression [34], and also attenuated anxiety in mice with DSS-induced colitis. Administration of *Bifidobacterium longum* also reduced anxiety; however, vagotomy prevented this probiotic effect [111]. Although vagal involvement has been identified in some studies, others show independence of vagal communication. For example, alteration of the microbiota increased exploratory behavior in mice and increased brain-derived neurotrophic factor in the hippocampus. These effects were not influenced by vagotomy [112]. More research is needed to elucidate this mechanism further.

Conclusion

Why do children with ASD have dysbiosis? Is this inherent to the disorder, or perhaps causal? These significant questions still remain to be determined. Immune system dysfunction is a well-known issue in ASD, possibly driving the dysbiotic microbiota, or alternatively created by it. The relationship between dysbiosis and the high incidence of comorbid GI dysfunction in ASD is not well elucidated; however, these studies reviewed here indicate that a relationship exists. Reports that behaviors improve after modification of the microbiota support the hypothesis that dysbiotic microbiota, their influence on the immune system, and their metabolic byproducts contribute directly to the development of these disorders. More research to clarify mechanism(s) of the influence of dysbiosis on brain and immune function and behavior are needed.

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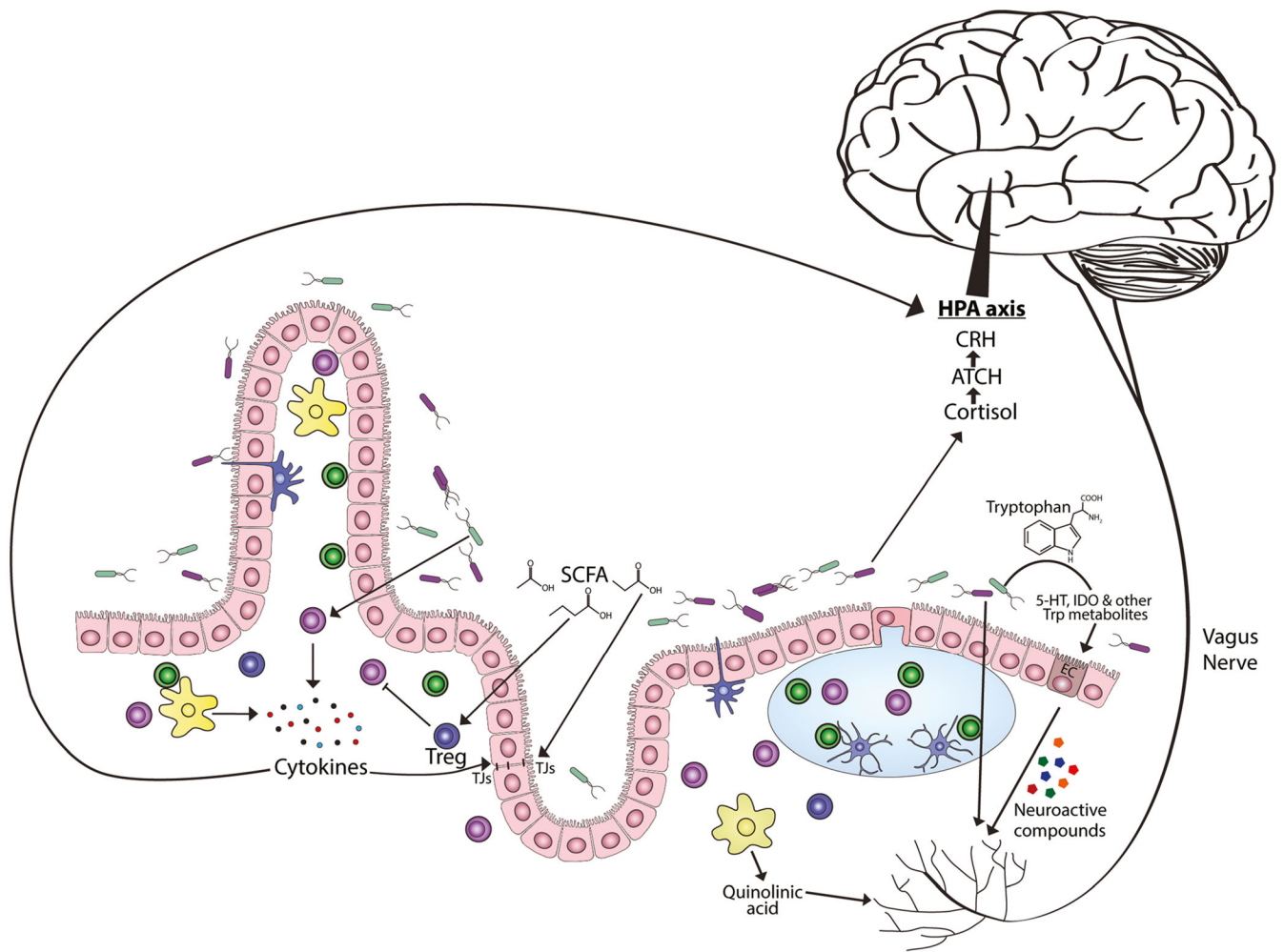


Fig. 1. Mechanisms of the microbiota-gut-brain axis. Possible mechanisms of the MGB-axis are being actively investigated and include neuroimmune pathways, neural communication through the vagus nerve, influence of metabolites produced by the microbiota, microbial-derived neurotransmitters, and the significant influence the microbiota have on tryptophan, kynurenine, and serotonin metabolism. Short-chain fatty acids (SCFA) can promote peripheral T regulatory (Treg) cell expansion as well as influence tight junction (TJ) proteins and intestinal barrier function. Microbiota regulate tryptophan (Trp) metabolites by degrading Trp to indole-derivatives or through kynurenine and serotonin pathways, such as increasing expression of tryptophan hydroxylase (Tph)1 in enterochromaffin (EC) cells. Dysbiosis can promote activation of immune cells, including macrophages that produce quinolinic acid (QA) through an alternative kynurenine pathway, a known excitotoxic N-methyl-D-aspartate (NMDA) receptor agonist. Activated immune cells also produce proinflammatory cytokines which can further disrupt microflora and impact intestinal barrier function. Neural communication can also occur through the vagus nerve via signaling from hormones and neurotransmitters release by gut endocrine cells and immune cells. Breach of

the intestinal barrier would also allow direct pattern recognition sensing due to Toll-like receptor expression on afferent fibers

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ASD microbiota studies

Table 1

Subject details	Methods	Findings	References
11 with regressive ASD (10M/1F); 3.5–7 years	Antibiotic Intervention Study Oral vancomycin 500 mg/day for 8 weeks, followed by 2 weeks daily probiotics (40 Billion CFU).	ASD behaviors improved after treatment with oral vancomycin. Gains were lost after treatment ended.	Sandler et al., 2000 [46]
13 with regressive ASD 8 TD; 3.5–7 years Gender not specified	Fecal, gastric/jejunum/duodenal juice samples: sPCR identification after culturing	Increased <i>Clostridia</i> in ASD. Spore formers found in upper GI of ASD, not present in TD	Finegold et al., 2002 [47]
15 ASD, 8 TD. Age, gender not specified	Fecal samples: RT-qPCR with species or cluster specific TaqMan primers/probes	Increased <i>Clostridia</i> clusters I (9-fold increase) and IX (3.5-fold increase), increased <i>C. bolteae</i> (46-fold increase)	Song et al., 2004 [48]
58 ASD (48M/10F; 53 with GI) 12 SIB (7M/5F; 3 with GI) 10 TD (6M, 4F; no GI) 3–16 years	Fecal samples: fluorescence in situ hybridization (FISH) for bacterial identification	Increased <i>Clostridia</i> associated with worsening GI symptoms and probiotic use. Intermediate levels seen in siblings.	Parracho et al., 2005 [49]
33 ASD with GI (24M/9F) 7 SIB no GI (2M/5F) 8 TD no GI 5M/3F 2–13 years	Fecal samples: bacterial tag encoded FLX amplicon pyrosequencing (bTEFAP)	Increased <i>Bacteroidetes</i> to <i>Firmicutes</i> ratio. Increased <i>Desulfovibrio</i> and <i>Bacteroides vulgatus</i> in severe ASD.	Finegold et al., 2010 [50]
15 ASD with GI 7 TD with GI All male 4–4.5 years	Ileal and cecal biopsies: RT-qPCR for expression of enzymes and transporters associated with carbohydrate digestion and transport. 16S rRNA pyrosequencing of cecal and ileal tissues for bacterial communities	Reduced mRNA transcripts for disaccharidases and carbohydrate transporters, associated with decreased <i>Bacteroidetes</i> increased <i>Firmicutes</i> and <i>beta-Proteobacteria</i>	Williams et al., 2011 [51]
23 ASD (21M/2F) 22 SIB (11M/11F) 9 TD (4M/5F) 3–18.5 years	Fecal samples: qPCR with bacteria specific primers	Reduced <i>Bifidobacteria</i> and <i>Akkermansia muciniphila</i> in feces of ASD children.	Wang et al., 2011 [52]
58 ASD (50M/8F) 39 TD (18M/21F) 3–12 years	Fecal samples: culture technique for bacterial measurement. Stool chemistry testing via ELISA. SCFA measured via gas chromatography.	GI scores strongly correlated with severity of ASD Reduced <i>Bifidobacteria</i> , increased <i>Bacillus</i> and <i>Lactobacillus</i> . Reduced levels of SCFA including propionate, especially in those taking a daily probiotic. Reduced lysozyme	Adams et al., 2011 [53]
41 ASD (32M/9F) 10 TD (5M/5F) 3–18 years	Fecal samples: culture technique for <i>Clostridia</i> measurement. ELISA for enterotoxin and lactoferrin analysis	No enterotoxins found. Elevated FLA in ASD boys only, with GI issues occurring 3–4 times per year. <i>Clostridia</i> found in both ASD and TD, the majority in ASD were <i>C. perfringens</i> . Strains varied from previous <i>Clostridia</i> studies, may be a factor of geographic location (Poland).	Martirosian et al., 2011 [54]
51 ASD, 28 with GI (42M/9F) 53 TD siblings (19M/34F) 2–12 years	Fecal samples: bTEFAP pyrosequencing of 16S rDNA	No significant differences at phylum and genus levels for ASD compared to TD siblings. Comparisons also done based on severity of ASD and GI symptoms, no differences detected.	Gondalia et al., 2012 [55]

Subject details	Methods	Findings	References
23 ASD with GI 9 TD with GI All male 4–4.5 years	Ileal and cecal biopsies - 16 s rRNA pyrosequencing and <i>Sutterella</i> -specific PCR, qPCR. Immunoblot for IgM/IgG reactivity	12/23 ASD positive for <i>Sutterella</i> , mainly <i>S. wadsworthensis</i> or <i>S. stercoricanis</i> . Immune response detected – IgG/IgM reactivity to <i>S. wadsworthensis</i> in 11 ASD children seen via Western blot. No positive controls.	Williams et al., 2012 [56]
23 ASD (21M/2F) 22 SIB (11M/11F) 9 TD (4M/5F) 3–18.5 years	Fecal samples: qPCR with bacteria specific primers	Elevated <i>Sutterella</i> in ASD, elevated <i>Ruminococcus torque</i> in ASD with GI, and associated with reduced <i>Akkermansia muciniphila</i> found previously.	Wang et al., 2013 [80]
10 ASD 10 PDD-NOS 10 TD 14M/16F; Dx not specified by sex 4–10 years	Fecal samples: bTEFAP pyrosequencing of 16S rDNA and rRNA. Metabolome measured by GC-MS/SPME.	Significant differences at the phylum levels among ASD vs. PDD-NOS vs. TD, highest diversity in ASD children. <i>Faecalibacterium</i> and <i>Ruminococcus highest</i> in PDD-NOS and TD; ASD high in <i>Caloramator</i> , <i>Sarcina</i> and <i>Clostridium</i> in ASD. <i>Bacteroidetes</i> higher in ASD and PDD-NOS, <i>Sutterellaceae</i> and <i>Enterobacteriaceae</i> highest in ASD. <i>Alistipes</i> and <i>Akkermansia</i> also higher in PDD-NOS and ASD. Differences in metabolites in ASD vs. TD (refer to Table 2 for details).	DeAngelis et al., 2013 [64]
20 ASD with GI (18M/2F) 20 TD no GI (17M/3F) ASD 6.7 (±2.7) years TD 8.3 (±4.4) years	Fecal samples: bTEFAP pyrosequencing of 16S rDNA	Decreased overall diversity and richness in ASD, correlating with ASD severity. Reduced carbohydrate fermenters Prevotellia, Coprococcus and unclassified Veillonellaceae.	Kang et al., 2013 [74]
59 ASD, 25 with GI (52M/7F) 44 SIB, 13 with GI (21M/23F) 7–14 years	Fecal samples: qPCR with bacteria specific primers, 16S rRNA, gene sequencing	No composition or diversity differences between ASD and NT siblings. Several low abundance taxa at genus level associated with ASD and/or ASD with GI.	Son et al., 2015 [75]
10 ASD (9M/1F) SIB (7M/2F) TD (all male) 2–17 years	Probiotic intervention study: Oral probiotics give 3x/day for 4 months. Fecal samples: RT-PCR with primers specific for main phyla and several species. No behavior measurement post-treatment.	Higher GI scores in ASD and siblings, ASD severity associated with GI symptoms. Reduced ratio of <i>Bacteroidetes</i> to <i>Firmicutes</i> in ASD, increased <i>Lactobacillus</i> . Increased <i>Desulfovibrio</i> associated with severity of ASD. Daily probiotic supplementation increased <i>Bacteroidetes</i> to <i>Firmicutes</i> ratio and reduced <i>Desulfovibrio</i> and <i>Bifidobacterium</i> in ASD, and <i>Clostridia</i> trended lower.	Tomova et al., 2015 [76]
21 ASD with GI (19M/2F; 14.4 ± 1.1 years) 19 TD with GI (10M/9F; 16.1 ± 1.3 years)	Duodenal biopsies; 16S rRNA gene sequencing. Disaccharidase activity measured via Dahlqvist method	No differences in disaccharidase activity. No differences in diversity/species richness and evenness. Increased <i>Burkholderia</i> and decreased <i>Neisseria</i> , <i>Bacteroides</i> species and <i>Escherichia coli</i> in ASD. <i>Clostridium</i> species correlated with disaccharidase activity in ASD.	Kushak et al., 2017 [77]
14 ASD with GI (all male) 15 TD with GI (12M/3F) 6 TD no GI (all male) 3–18 years	Rectal biopsies: Biopsy supernatant for 16S rRNA gene sequencing for microbiota characterization, 5-HT metabolites measured via HPLC. Blood and biopsy supernatant for cytokines via Luminex.	ASD with GI had increased <i>Clostridiales</i> , decreased <i>Dorea</i> , <i>Blautia</i> , and <i>Sutterella</i> . Abdominal pain correlated significantly with IL-6, IL-1, IL-17A, and IFN-γ. IL-6 and tryptophan highest in ASD with GI pain, serotonin metabolites were increased in all with GI. Inflammatory cytokines, tryptophan and serotonin correlated with <i>Clostridiales</i> in ASD.	Luna et al., 2017 [83]
40 ASD (31M/9F) 40 TD (28M/12F) 3.5–17 years	Fecal samples: Calprotectin levels measured by ELISA. 454 pyrosequencing	No differences in Calprotectin levels in feces. Reduced ratio of <i>Bacteroidetes</i> to <i>Firmicutes</i> and decreased <i>Alistipes</i> , <i>Bifidobacteria</i> , <i>Dialister</i> , <i>Parabacteroides</i> , and <i>Veillonella</i> , increased <i>Collinsella</i> , <i>Corynebacterium</i> , <i>Dorea</i> , and <i>Lactobacillus</i> in ASD. Greater than twice the abundance of <i>Candida</i> in ASD. High levels of <i>Escherichia</i> , <i>Shigella</i> and <i>Clostridium</i> cluster XVIII associated with constipation in ASD.	Strati et al., 2017 [78]
18 ASD (16M/2F) 20 TD (18M/2F) 7–16 years	FMT Intervention Study 2-week tx with oral vancomycin followed by bowel cleanse and extended FMT: high initial dose, then daily maintenance doses for 7–8	Fecal microbiota transplant improved behavior and GI symptoms in ASD, increased levels of <i>Bifidobacteria</i> , <i>Prevotella</i> and <i>Desulfovibrio</i> TD controls were monitored, but not treated.	Kang et al., 2017 [79]

Subject details	Methods	Findings	References
21 ASD with GI (25M/4F) 29 ASD no GI (17M/12F) 7 TD with GI (6M/1F) 34 TD no GI (32M/2F) 3–12 years	weeks. Periodic fecal samples and swabs: 16 s rRNA gene sequencing. Pre-, during and post behavioral assessments: PGI-III, CARS, ABC, SRS and VABS-II Fecal and blood samples: PBMC stimulation and cytokine analysis of supernatant via Luminex. Fecal 16S rRNA gene sequencing.	Elevated inflammatory immune responses in ASD with GI. Balance of inflammatory vs regulatory cytokines skewed. ASD microbiota clustered differently than TD samples, increased <i>Bacteroidaceae</i> , <i>Lachnospiraceae</i> , <i>Prevotellaceae</i> and <i>Ruminococcaceae</i> in ASD with GI vs TD with GI. Differences seen in functional KEGG pathways in ASD compared to TD. Altered zonulin levels in ASD with GI, may suggest increased intestinal permeability.	Rose et al., 2018 [57]
23 ASD (22M/1F) 21 TD (15M/6F) 4–17 years	Fecal samples: 16S rRNA gene sequencing	Reduced diversity, reduced <i>Prevotella copri</i> , <i>Faecalibacterium prausnitzii</i> and <i>Haemophilus parainfluenzae</i> . Altered microbial metabolites (refer to Table 2 for details).	Kang et al., 2018 [73]

ASD autism spectrum disorder, *regressive ASD* child developed typically but lost skills and speech, usually between 15 and 30 months, *TD* unrelated typically developing control, *SIB* typically developing sibling, *ASPA* sparger's disorder, *GI (group)* gastrointestinal symptoms, *PDD-NOS* pervasive development disorder-not otherwise specified, *Dx* diagnosis, *GC* gas chromatography, *MS* mass spectrometry, *SPME* solid-phase microextraction, *FMT* fecal microbiota transplant, *Tx* treatment, *FLA* fecal lactoferrin

Table 2

ASD microbial metabolome studies

Subject details	Methods	Findings	References
39 ASD (35M/4F) 28 SIB (14M, 14F) 34 TD (17M, 17F) 3-9 years	Urine samples: ¹ H NMR spectroscopy	Altered pattern of microbial co-metabolites dimethylamine, hippurate, and phenylacetylglutamine. Elevated Glutamate and Taurine also found in ASD Urine.	Yap et al., 2010 [58]
262 ASD (211M/51F) 60 TD (30M/30F) 2-13 years 12 Schizophrenia (4M/8F) 19 Adult controls (11F/8M)	Urine samples: GC/MS	Elevated 3-(3-hydroxyphenyl)-3-hydroxypropionic acid (HPHPA) from <i>Clostridium</i> species in ASD. Also extremely elevated in a schizophrenia patient during acute psychosis, reduced significantly after oral vancomycin treatment.	Shaw 2010 [59]
59 ASD (44M/15F) 59 TD (44M/15F) 2-18 years	Urine samples: HPLC, UV-DAD	Elevated urinary microbial metabolite p-cresol in ASD <8 years old, especially girls. Also elevated in severe ASD regardless of sex.	Altieri et al., 2011 [60]
23 ASD (21M/2F) 22 SIB (11M/11F) 9 TD (4M/5F) 3-18.5 years	Fecal samples: C18R method, GC, HPLC	Elevated ammonia and SCFA (acetic, butyric, isobutyric, valeric, isovaleric acids), likely of microbial origin. Reduced caproic acid in ASD. No differences in phenol and p-cresol levels.	Wang et al., 2012 [61]
22 ASD with GI (20M/2F) 4-10 years	Urine samples: GC/MS	Elevated D-arabinitol (DA) produced by <i>Candida</i> spp. in ASD urine, elevated DA/L-arabinitol ratio. Significant decrease in DA and DA/LA ratio with probiotic <i>L. actophilus</i> . Improvements also seen in concentration and ability to follow orders after probiotic treatment.	Kaluzna-Czaplinska et al., 2012 [62]
48 ASD, 29 with GI (36M/12F) 53 TD (34M/19F) Median ages: 10.7/10.2 years	Urine samples: UPLC/MS/MS, GC/MS	Microbial co-metabolites significantly altered in ASD with GI issues including significantly increased levels of 2-(4-hydroxyphenyl)propionate and taurocholate sulfate and significantly decreased 3-(3-hydroxyphenyl)propionate and 5-aminovaleate. ASD also had altered amino acid metabolism and decreased antioxidants, including increased glutaroylamine derived from tryptophan and lysine metabolism.	Ming et al., 2012 [63]
10 ASD 10 PDD-NOS 10 TD 14M/16F; Dx not specified by sex 4-10 years	Fecal samples: GC-MS/SPME	Lower levels of fecal SCFA in ASD (except PPA), correlated with reduced <i>Escherichia coli</i> and <i>Bifidobacterium</i> . Elevated phenol, 4-(1,1-dimethylethyl)-phenol, and p-cresol. Elevated free amino acids indicating increases in proteolytic bacteria in ASD.	De Angelis et al. 2013 [64]
33 ASD 33 TD 4-16 years	Urine samples: HPLC-FLD	Elevated urinary microbial metabolites p-cresol, p-cresylsulfate and p-cresylglucuronate in ASD 8 years old. Not associated with severity of ASD, however increases correlated with repetitive behaviors.	Gabriele et al., 2014 [65]
Review of 22 studies identifying levels of 5-HT in individuals with ASD compared to controls	Systematic review and meta-analysis	Significantly higher 5-HT in ASD, both in whole blood and platelet-rich plasma. Platelet-poor plasma yielded no differences. HPLC and fluorometric assays yielded similar cumulative effect sizes, HPLC had lowest variability. Analysis may be limited by lack of normalization by platelet number in most studies.	Gabriele et al., 2014 [66]
21 ASD 21 SIB 4-17 years Sex not specified	Urine samples: GC-MS	Increased 3-(3-hydroxyphenyl)-3-hydroxypropanoic acid, 3,4-dihydroxybutyric acid. Increased glycolic acid, ribose, arabinol, xylitol and threitol, some with possible fungal origin. Increased tryptophan, glycine, cis-aconitic acid; phenylalanine, tyrosine, p-hydroxyphenylacetic acid, and homovanillic acid in ASD	Noto et al., 2014 [67]
30 ASD (22M/8F) 30 TD (22M/8F)	Urine samples: HILIC-UHPLC, MS	Urinary metabolites of tryptophan and purine metabolism significantly altered in ASD, specifically tryptophan metabolized through an alternative kynurenine pathway yielding increased	Gevi et al., 2016 [68]

Subject details	Methods	Findings	References
2–7 years		xanthurenic and quilolinic acids. Increased indoxyl sulfate, indoxyl 3-acetic acid and indoxyl lactate, created through microbial degradation of tryptophan. Significant decrease in melatonin its catabolite N-acetyl-5-methoxy-tryptamine.	Xiong et al., 2016 [69]
62 ASD (48M/14F) 62 TD (48M/14F) 1.5–7 years	Urine samples: GC/MS Vancomycin treatment in 50 HPHPA+ ASD children for 30 days, followed by <i>Bifidobacterium</i> BB-12 supplementation.	HPHPA, 3HPA, and 3HHA concentrations were significantly higher in Asd compared to TD. Vancomycin significantly decreased HPHPA, 3HPA, and 3HHA (eliminated in 35/50, 6/9, and 12/17 cases respectively). Levels rebounded to initial levels in 3 patients within 6 months, others had lesser but measurable recurrence consistent with likely <i>Clostridia</i> germination after antibiotics.	Gabriele et al., 2016 [70]
53 ASD (44M/9F) 59 TD 1–20 years	Urine samples: HPLC Fecal cultures to measure <i>Clostridium</i>	p-cresol significantly higher in ASD than previously assessed controls. Elevated urinary p-cresol associated with constipation in ASD, but not with <i>Clostridium</i> species. All ASD tested negative for <i>C. difficile</i> .	Luna et al., 2017 [83]
14 ASD with GI (14M) 15 TD with GI (12M/3F) 6 TD no GI (6M) 3–18 years	Rectal biopsies: HPLC of mucosal supernatant	Decreased Tryptophan, increased serotonergic metabolites in ASD children with GI compared to TD, including 5-HIAA which was associated with abdominal pain. <i>Erysipe lotrichaceae</i> , <i>Clostridium lituseburense</i> , and <i>Terrisporobacter</i> correlated with tryptophan in ASD with GI. <i>Lachnoclostridium bolleae</i> , <i>Lachnoclostridium hathewayi</i> , and <i>Faecalibacterium plautii</i> correlated with serotonin.	Lussu et al., 2017 [72]
21 ASD (18M/3F) 21 SIB (10M/11F) 4–16 years	Urine samples: ¹ H NMR spectroscopy	Increased tryptophan and hippurate, dysregulated amino acid metabolism compared to siblings.	Kang et al., 2018 [73]
21 ASD (15M/6F) 23 TD (22M/1F) 4–17 years	Fecal samples: ¹ H NMR spectroscopy 16s rRNA gene sequencing	Increased isopropanol, p-cresol. Trending decrease in GABA, TD had higher <i>Streptococcus thermophilus</i> which may contribute to normal GABA production. No differences in butyrate or PPA	

ASD autism spectrum disorders. TD unrelated typically developing control. SIB typically developing sibling. GC gas chromatography. MS mass spectrometry. HPLC high-performance liquid chromatography. UV-DAD ultraviolet ultraviolet diode array detection. UPLC/MS/MS ultrahigh performance liquid chromatography/tandem mass spectrometry. SPME solid-phase microextraction, C1CR catalyzed indophenol colorimetric reaction, SCFA short-chain fatty acids, SPME solid-phase microextraction, FLD fluorescence detector, 5-HT 5-hydroxytryptamine (serotonin), PRP platelet-rich plasma, HILIC hydrophilic interaction chromatography. PPA propionic acid, GABA γ -aminobutyric acid