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Brain structure and response to emotional stimuli as related to gut microbial profiles in healthy women

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Abstract

Objective—Brain-gut-microbiota interactions may play an important role in human health and behavior. However, while rodent models have demonstrated effects of the gut microbiota on emotional, nociceptive and social behaviors, there is little translational human evidence to date. In this study we identify brain and behavioral characteristics of healthy women clustered by gut microbiota profiles.

Methods—Forty women supplied fecal samples for 16s rRNA profiling. Microbial clusters were identified using Partitioning Around Medoids. Functional magnetic resonance imaging was acquired. Microbiota-based group differences were analyzed in response to affective images. Structural and diffusion tensor imaging provided gray matter metrics (volume, cortical thickness, mean curvature, surface area) as well as fiber density between regions. A sparse Partial Least Square-Discrimination Analysis was applied to discriminate microbiota-clusters using white and gray matter metrics.

Disclosures: The remaining authors disclose no conflicts.

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Results—Two bacterial genus-based clusters were identified, one with greater *Bacteroides* abundance ($n=33$), one with greater *Prevotella* abundance ($n=7$). The *Prevotella* group showed less hippocampal activity viewing negative valences images. White and gray matter imaging discriminated the two clusters, with accuracy of 66.7% and 87.2% respectively. The Prevotella cluster was associated with differences in emotional, attentional, and sensory processing regions. For gray matter, the *Bacteroides* cluster showed greater prominence in the cerebellum, frontal regions, and the hippocampus.

Conclusions—These results support the concept of brain-gut-microbiota interactions in healthy humans. Further examination of the interaction between gut microbes, brain and affect in humans is needed to inform preclinical reports that microbial modulation may affect mood and behavior.

INTRODUCTION

Comprised of trillions of organisms and responsible for numerous biologically important processes, the human microbiota has a role in health and disease that is increasingly evident (1, 2). One area of particular interest is the role of the gut microbiota within the gut-brain axis and their relationship to emotional processing (3, 4). While spatially separated from the brain by both the intestinal epithelial barrier and the blood brain barrier, there is bidirectional communication between the gut microbiota and the central nervous system (CNS) via the vagus nerve, the immune system, and neuroactive metabolites released into systemic circulation (5).

Evidence for the influence of the microbiota on the CNS has been plentiful in the preclinical literature and suggests that beyond brain development, the microbiota can influence behavior and affect (5, 6). Animals raised in germ-free environments exhibit altered brain chemistry as well as changes in behavior, with increased risk taking, reduced anxiety, and decreased sociability (3, 5, 7–10). Some of the biochemical changes resulting from germ-free status are irreversible, even after colonization of the animals with normal gut microbiota later in life. Other abnormalities, such as anxiety behavior, can be ameliorated after reconstitution of the gut microbiota (7).

Along with the preclinical evidence, emerging evidence in humans suggests that while some aspects of the microbiota's influence on the CNS are likely to be established early as traits, other aspects may be malleable and are vulnerable to environmental factors (11, 12). For example, in the only study to date showing interactions between brain structure and gut microbes, results suggested that obese patients had brain structural changes in the hypothalamus and caudate nucleus associated with specific microbiota profiles and changes in cognitive function. However, in this cross-sectional design, it was not clear whether these changes may have been a risk factor for, or a result of, obesity status (13). Small studies of probiotic interventions have shown modest effects on mood and variable effects on cognition, leading to speculations that gut bacteria may be evolutionarily programmed to improve our moods, thus making us more social and prone to activities that allow person-toperson transmission of the organisms (14–17). We have previously demonstrated in healthy women that 4-week ingestion of a fermented milk product with probiotics can shift functional brain responses to an emotional attention task towards lower reactivity in

viscerosensory, somatosensory, and affective regions, providing more direct evidence in humans that brain function can be affected by modulation of the gut microbiota (18).

In this study, using fecal samples, magnetic resonance imaging (MRI), and an emotion induction task obtained in conjunction with our previous study (18), we aimed to identify whether the gastrointestinal microbiota composition in healthy women was associated with characteristics of brain structure, brain structural white-matter connectivity, and brain function as measured by affective response to emotionally valenced images.

METHODS

Subjects

Healthy, non-obese women aged 18–55 were recruited by advertisement. A medical history, physical exam, and standardized psychiatric screening exam (Mini International Neuropsychiatric Interview Plus 5.0)(19) were performed. Exclusion criteria included: use of antibiotics or probiotics in the month prior, active pain disorder, active medical or psychiatric illness, tobacco dependence, pregnancy or lactation, exercise >8 hours/week, metallic implants, claustrophobia, body mass index >30 or <18. Subjects taking central nervous system (antidepressants including serotonin reuptake inhibitors and serotonin/ norepinephrine reuptake inhibitors, sedatives or anxiolytics) medications or use of opioid analgesics were excluded. Informed consent was obtained from all subjects. Data acquisition was performed between January 2009 and December 2012. All procedures complied with the principles of the Declaration of Helsinki and were approved by the Institutional Review Board at UCLA.

Behavioral and Clinical Measures

Questionnaires were completed before MRI scans. A 2-week electronic diary was used to assess gastrointestinal symptoms prior to MRI. Subjects were excluded if they reported abdominal pain or discomfort on 2 or more days, abnormal stool form (Bristol stool scale 1, 6, or 7) or stool frequency of less than 3 bowel movements per week or greater than 3 bowel movements per day (20). The Positive Affect Negative Affect Schedule (PANAS) was used to measure positive and negative affect (21, 22). The Hospital Anxiety and Depression Scale (23) was used to assess active mood symptoms and the Spielberger Trait Anxiety Inventory (24) was collected.

Stool collection and 16S rRNA pyrosequencing

Subjects were provided stool collection kits with a sterile collection container, ice packs and a freezer bag for transport. They were instructed to collect the stool on the day prior to their MRI visit, freeze it in their home freezer and then transport it on ice packs to the Research Center where the stool was frozen at −80 degrees until analysis. Samples were shipped frozen to the laboratory facilities at Danone Research (Palaiseau, France) where it underwent 16S rRNA profiling using 454 pyrosequencing (V5–V6 variable regions). V5 and V6 hypervariable 16S ribosomal RNA (rRNA) regions were amplified using primers 784F and 1061R (25). Sequencing was performed by DNAVision SA (Charleroi, Belgium) on a 454 Life Sciences Genome Sequencer FLX instrument (Roche) using titanium chemistry

and primer A. Analyses were performed using Quantitative Insights Into Microbial Ecology (QIIME) v1.7 (26). Reads were filtered according to the following quality criteria: size between 150 and 500nt, quality above 25 over a 50 base pairs window, no mismatch authorized in primers and barcode sequences, and absence of polymers larger than 6nt. Remaining sequences were denoised using default parameters and clustered into Operational Taxonomic Units (OTUs) defined at 97% identity using CD-HIT (27). Representative sequences for each OTU were aligned and taxonomically assigned using Greengenes (August 2011). ChimeraSlayer was used to discard potential chimeric sequences (28) leading to 5943 +/− 3100 (mean +/− SD) reads per sample. Normalization was performed using relative abundances. Beta diversity was performed on weighted Unifrac distances using rarefaction of 1770 sequences per sample. To cluster the samples based on weighted distance matrix, we used the Partitioning Around Medoids (PAM) algorithm in the R package 'Cluster'. The optimal number of clusters was two, as it had the highest Rousseeuw's Silhouette internal cluster quality index (SI=.36), calculated using the R package 'Cluster' (29). The resulting microbial clusters showed one with greater Prevotella abundance (*Prevotella*-high, n=7) and one with greater *Bacteroides* abundance (*Bacteroides*high, n=32) (Figure 1), SI=.36. The variance accounted for by the two components that defined the microbiota clusters totaled 57.2% (Component 1=39.5%, Component 2= 17.7%). These two clusters were utilized for the analyses described below. To identify demographic factors which may confound the cluster designations, t-tests were performed for age, body mass index, anxiety and depression (HAD score), and positive and negative affect using Statistical Package for the Social Sciences (SPSS) software (version 19).

MRI Data Acquisition

Subjects were scanned on a 3-Tesla Siemens Trio after a sagittal scout was used to position the head. Structural scans were acquired using a high-resolution 3-dimensional T1-weighted, sagittal magnetization-prepared rapid gradient echo (MP-RAGE) protocol: Repetition Time $(TR) = 20$ ms, echo time $(TE) = 3.00$ ms, flip angle $(FA) = 25^\circ$, field of view $(FOV) = 256$ mm, acquisition matrix = 256×256 , slice thickness 1mm, voxel size $1 \times 1 \times 1$ mm³. Functional scans were acquired in transverse orientation, interleaved, with $TR = 2500$ ms, $TE = 26$ ms, $FA = 90^\circ$, $FOV = 200 \text{mm}$, acquisition matrix of $64 \times 64 \text{mm}$, slice thickness 3.0mm, and voxel size $3.1 \times 3.1 \times 3.0$ mm³. Diffusion weighted MRIs (DTI) were acquired in 64 non-collinear directions with $b = 1000$ s/mm² images with the following protocol: TR = 7000ms, TE = 93ms, and FOV = 190mm with an acquisition matrix of 96×96 , and a slice thickness of 2mm to produce $2 \times 2 \times 2$ mm³ isotropic voxels.

Functional MRI (fMRI) and emotion induction task

Functional MRI was performed in three runs. During each run the subjects viewed mood inducing negative, positive or neutral valence pictures presented in four blocks of the same valence. Negative and positive runs were counterbalanced with the neutral run in the second position. Each run contained 24 pictures of the same valence presented in 4 blocks. Pictures were presented for 5 seconds each, and a 30 second interblock interval (during which a crosshair was presented) separated each block. The images were selected from the International Affective Picture Set to have moderate arousal and valence (9 point scale)

based on the mean scores for women.(30). Arousal and valence scores are presented in Table 1. They were converted to black and white and were luminosity adjusted.

The first 2 volumes were discarded to allow for stabilization of the magnetic field. The remaining functional images were slice-time and motion corrected, spatially normalized to the Montreal Neurologic Institute template, and spatially smoothed with an 8mm^3 Gaussian kernel using SPM8 (Welcome Department of Cognitive Neurology, London, UK). The experiment was analyzed in a block design. A first-level fixed effects general linear model (GLM) was applied in SPM8 to determine blood oxygen level dependent (BOLD) activity by specifying as regressors, the 30 second baseline and the three valence conditions (negative, positive, neutral). At the subject-level, regressors for the conditions were convolved with a canonical hemodynamic response function to estimate brain activity. Individual brain responses were determined by subtracting estimated brain activity during the neutral valence image condition from estimates of brain activity during the positive and negative affect conditions. To determine microbiota-based cluster group differences, a GLM was implemented at the second-level using a region of interest (ROI) approach. ROIs defined by the Destrieux atlas (31) included the extended emotional arousal network (amygdala, anterior cingulate and hippocampus). An initial cluster defining threshold of p<. 001 was applied and cluster based extent threshold significance was p<.05 after controlling for family wise error as implemented in SPM (32).

Structural MRI Analysis

Data pre-processing workflows for the MRI data were designed and created in collaboration with the University of Southern California, Laboratory of Neuroimaging (LONI) Pipeline [\(pipeline.loni.usc.edu](http://pipeline.loni.usc.edu)). T1-image segmentation and regional parcellation were conducted using FreeSurfer (33, 34) following the nomenclature described in (31). Based on the Destrieux and Harvard-Oxford atlases, for each cerebral hemisphere, a set of 74 cortical structures were labeled in addition to 7 subcortical structures, the cerebellum and the brain stem, resulting in a complete set of 165 parcellations for the entire brain. Four representative morphological measures were computed for each cortical parcellation: gray matter volume (GMV), surface area (SA), mean cortical thickness (CT), and mean curvature (MC) (35, 36). For subcortical regions only volume was computed.

Diffusion tensors were computed and rotationally re-oriented at each voxel. Tensor-valued images were linearly realigned based on tri-linear interpolation of log-transformed tensors as described in (37) and resampled to an isotropic voxel resolution ($2 \times 2 \times 2$ mm³). White matter connectivity for each subject was estimated between the 165 brain regions using DTI fiber tractography performed via the Fiber Assignment by Continuous Tracking algorithm (38) using TrackVis software. The final estimate of white matter connectivity between each of the brain regions was determined based on the number of fiber tracts intersecting each region, normalized by the total number of fiber tracts within the entire brain (39–42).

Sparse partial least squares Discriminant analysis (sPLS-DA)

A sPLS-DA was used to determine if the brain signatures (gray matter morphology and white-matter connections) could discriminate microbial cluster membership. sPLS-DA

simultaneously performs feature selection and modelling and achieves sparsity using lasso penalization (43). sPLS-DA operates using a supervised framework to find orthogonal components, linear combinations of a limited set of variables (brain features) that predict class membership. We refer to each component as discriminatory "brain signature". sPLS-DA was performed using the R package mixOmics version 5.1([http://www.R-project.org\)](http://www.R-project.org).

Features entered in the sPLS-DA models—The predictive power of brain morphometry and DTI white-matter connectivity were assessed separately. Covariates for all models included age and total GMV. To investigate morphometry, GMV, SA, CT, and MC estimates for the 165 brain regions were entered as predictors. For DTI white-matter connectivity data, subject-specific matrices indexing relative fiber density between the 165 regions were transformed to 1 dimensional matrices containing 13,530 unique connectivities (upper triangle from the initial matrix). These matrices were then concatenated across subjects and entered into the sPLS-DA. As an initial data reduction step, near zero variance predictors were dropped resulting in 2751 predictors for the anatomical based discrimination.

sPLS-DA model specifications and development—For each model, the number of components to identify was fixed at 2 (42, 44). To select the optimal number of features for each component we estimated the 5-fold classification error with respect to a range of number of features (10 to 200 by units of 10). This process was repeated 50 times and the results averaged. This "tuning" procedure indicated that two components comprised of 10 brain features each would be optimal for both the DTI and Morphometric models.

Model summary indices—The discriminatory brain signatures were summarized using variable loadings and VIP coefficients. Each variable has an associated "loading" indexing the relative importance of that variable in the brain signature for group discrimination (42). Variable importance in projection (VIP) scores is a standardized measure that represents contribution of each feature relative to the variance explained by all selected brain signatures (44). Usually, predictors with VIP coefficients greater than one are considered particularly important for discrimination (42). Graphical displays illustrated the discriminative abilities of the algorithms (44). The accuracy of the final models were assessed using leave-one-out cross-validation. This overall error rate or accuracy reflects the number of correct predictions from all the predictions made. During this cross-validation procedure, we calculate the stability of the selected variables comprising a component if a training set is altered (44). Stability is computed by calculating the frequency of selected variables across the cross validation runs. During this cross validation step, it was noted that the classification accuracy for of both models decreased with 2 components, therefore the second component for each model was dropped. PLS-DA is discovery-based method that enables generation of novel hypotheses. Post hoc exploratory evaluation of the specific role of *Prevotella* and Bacteriodes on specific brain regions and affect was performed by correlating bacterial abundances with brain metrics and PANAS scores (See Tables S1–S3, Supplemental Digital Content 1).

RESULTS

Cluster Characteristics

The total sample of 40 healthy females included 7 subjects in the Prevotella-high group and 33 subjects in the *Bacteroides*-high group (Figure 1). The mean age of the subjects was 28.89 years, SD 9.87; body mass index (BMI) was normal with a mean of 23.33, SD = 2.69; anxiety and depression symptoms were within normal ranges (mean 3.48 , SD = 2.44 and 1.2 , $SD = 1.81$, respectively), and normalized trait anxiety was in the mild to moderate range with a mean of 43.03 , $SD = 8.87$. No differences between the microbiota clusters were identified for age (t = 1.58, p = .12), BMI (t = 1.11, p = .27), anxiety (t = 1.08, p = .29), depression (t = .09, p = .93), or trait anxiety (t = .61, p = .54). Mean PANAS scores for positive and negative affect at baseline and after viewing positive and negative images are shown in Table 2. No difference in baseline positive or negative affect on the PANAS was seen between groups (positive affect t=−.84, p=.41; negative affect t=.16, p=.87). The PANAS questionnaire data was missing for one subject in the *Bacteroides*-high group so that subject was excluded from the analysis of affect. One subject in the *Bacteroides*-high group had imaging data that failed quality control measures and was excluded, leaving a total of 39 subjects analyzed for neuroimaging.

Microbiota cluster-related group differences in emotional response

Changes in positive and negative affect scores (PANAS) obtained at baseline and after each valence block confirmed appropriate affect response to the positive and negative valence blocks. The Prevotella-high cluster group had higher negative affect after viewing the negative valence picture block, p=.012 (21.86, SD=10.2 vs. 15.9, SD=3.8). There was no difference between groups after viewing positive pictures, p=.36 (29.57, SD=8.3 vs. 25.93, SD=9.6). A significantly lower BOLD activity in the right hippocampus during negative stimuli was observed in the Prevotella-high group compared to the Bacteroides-high group $(p = .041,$ familywise error corrected; Z=4.4, cluster size =16). No differences were seen in the other regions during negative emotion condition and no regions of interest were different in the positive emotion condition.

Anatomical white-matter signatures associated with microbiota group clusters

One brain signature comprised of 10 white matter connectivities was able to discriminate between the *Bacteroides*-high and *Prevotella*-high clusters. Figure 2 demonstrates this discrimination by plotting individuals with respect to their scores on the signature. This anatomical signature accounted for 85% of the variance in group differences. Overall classification accuracy was 66.7%. This anatomical signature is comprised of 10 brain white-matter connectivities with all connections showing less fiber density in the Bacteroides-high cluster (Table 3). Based on the VIP coefficients, the connectivities having the most explanatory power were between emotional regulation (amygdala, anterior cingulate cortex) and basal ganglia regions as well as attentional (right middle frontal gyrus) and sensory (the right central sulcus) regions. Other sensory connectivities included thalamic connections with the pericallosal sulci and the temporal pole, and connectivity between the left central sulcus and the left posterior midcingulate cortex. Several connections with the temporal gyrus were also major contributors in the prediction. The stability analysis

indicated that 7 of 10 selected brain features were quite reliable with >90% stability (See Table 3).

Morphological gray matter signatures associated with microbiota group clusters

Three of the four gray matter characteristics (GMV, CT, MC) discriminated the two clusters. One signature comprised of 10 gray matter metrics demonstrated good discriminative ability. Figure 3 plots the individual scores on the brain signature. Overall classification accuracy was 87.2%. Table 4 contains the list of selected gray matter metrics for each brain signature along with their loadings, VIP coefficients and stability indices. The first brain signature explained 52.8% and the second explained 21.8% of the variability in microbiota cluster discrimination. The variables with the most explanatory power included volume of the cerebellum and the hippocampus and the cortical thickness of the frontomarginal gyrus and the anterior insula, which were larger in the *Bacteroides*-high cluster. In the *Prevotella*-high cluster the nucleus accumbens had a greater volume, and two regions, the subparietal sulcus and the superior occipital gyrus, showed greater mean curvature. High reliability was observed for 8 of 10 brain morphometric features contributing to the model.

DISCUSSION

To date, most of our understanding of the interaction between the microbiota and the brain has come from rodent models, in which the gut microbiota is linked to brain signaling mechanisms and affective behavioral phenotypes, such as anxiety or depression-like behavior. The relationship between gut microbial community structure, brain structure and emotional processing observed in this study is consistent with some of these preclinical findings.

When healthy women were clustered by their stool microbiota composition into two groups, the groups showed differential response to negatively valenced images, with heightened increases associated with negative affect in the cluster with greater abundance of Prevotella. This tendency towards greater behavioral responses to negative valenced stimuli was associated with both functional and structural differences in the hippocampus. The hippocampus, a brain region involved in emotion regulation, had lower volume and showed less BOLD response during negative image viewing. Reduced hippocampal engagement to negative imagery may be associated with increased emotional arousal. Such changes have been suggested to result in less specificity of encoding the contextual details of incoming stimuli, a deficit seen in the setting of several psychiatric disorders, including depression, post traumatic stress disorder, and borderline personality disorder (45–47). While the subjects in this study were healthy, it is possible that the patterns which emerge from the microbial clustering represent vulnerability factors.

The interaction between behavior, functional response, and microbiota was supported by the structural brain signatures, which describe the microbial clusters. The observed lower hippocampal volume and higher volume of the nucleus accumbens in the *Prevotella*-high group are consistent with greater reactivity to an affective stimulus, with similar volumetric changes described in some studies of mood disorder (48, 49). The *Prevotella*-high group was associated with greater white matter connectivity in limbic-cortical-striatal-pallidal-thalamic

circuitry (e.g., anterior cingulate cortex to pallidum, amygdala to caudate), and a smaller hippocampal volume compared to the *Bacteriodes*-high group, both findings also seen in depression (50). This group also showed differences in regions associated with attention and sensory processing. Specifically, they showed more connectivity between the central sulcus, which borders the primary somatosensory and motor cortices, and the middle frontal gyrus, a region at the convergence of the dorsal and ventral attentional networks (51); more dense white matter tracts were seen between the splenium and thalamus, which along with less cortical thickness in the anterior insula, suggests altered sensory processing. Higher connectivity in the Prevotella group was also noted between multiple regions of the temporal lobe associated with visual conceptual/semantic processing (e.g., fusiform gyrus, inferior temporal gyrus, collateral sulcus, temporal pole) (52, 53).

To our knowledge, this is the first report of behavioral and neurobiological differences related to microbial composition in healthy humans. Although these groups were identified using an unsupervised approach based on microbial composition, the identified two clusters of subjects defined by the genera Bacteriodes and Prevotella are similar to clusters previously identified across diverse populations (54–56). These clusters have been described variously as enterotypes or enterogradients, though this nomenclature has been controversial both because it implies a fixed characteristic of a individual or group and because not all of the clusters are reproducible across studies and analytic techniques (57). Of the reported "enterotypes" the most robust groupings appear to be those based on Bacteroides and Prevotella with frequency in the Western population being dominated by the Bacteroides group, as is seen in the current sample (54–56). In previous studies these groupings have been independent of age, sex, body mass index or nationality and are shared between monozygotic twins at a high rate (58). Based on a diet questionnaire recalling food intake over the past year (Food Frequency Questionnaire) (59) but not a short term recall, a diet more plentiful in saturated fat and animal protein was associated with the *Bacteroides* group and a diet rich in plant based fiber with the *Prevotella* group (56) suggesting an important role for long term diet. Further, in a short term study using a controlled dietary intervention with either high fiber/low fat or high fat/low fiber, individuals failed to change from one "enterotype" to another, despite rapid changes in individual bacterial species (56). Beyond these descriptions, the relative abundance of *Bacteroides* and *Prevotella* has not been firmly associated with any specific health or physiologic phenotype and the functional pathways associated with the microbiota clusters are not clearly defined. While "enterotypes" have been described to be fairly stable, there has been a published report of a single healthy individual moving between "enterotypes" over time, without a specific intervention to induce such change (60). Such changes could certainly be mirrored or even induced by changes in brain function and by affective responses, but changes in structure would not be expected on this temporal scale. Therefore, if an interaction between diet and the brain signatures described in the current study exists, it could be based on long term diet, or potentially on dietary factors in early life, while the brain is developing the cortico-limbic and sensory circuitry featured in the brain signatures.

While the findings of this study are consistent with reports from rodent models of brain-gutmicrobiota interactions, caution should be used in the interpretation of this proof of concept study, as it is based on a small sample of physically and psychologically healthy women, and

the results may or may not be generalizable to other populations. Due to the preponderance of individuals in the Western population with lower Prevotella abundance, supported in the distribution seen in this sample, the *Prevotella* group analyzed is very small. Conclusions drawn from this sample will require further validation in larger samples. Further, it is unclear whether the propensity towards developing a more negative affect during this paradigm is a marker of personality traits, a risk factor for developing clinically relevant negative mood states, or just a healthy variant. Although sPLS-DA was chosen specifically because of its ability to deal with small samples, biases due to the unbalanced nature of the samples may exist. For the fMRI task analysis, non-parametric testing for cluster-based inference may have resulted in alternative p-values and better control over family-wise error rate. Importantly, relatively few brain biomarkers were able to predict microbial subgroup membership better than chance and the variables selected for the solutions make biological sense in terms of previous preclinical data on neurobiological-microbiota associations. Ultimately, future studies are needed to test the generalizability of the solution and the estimates of overall model accuracy to other samples.

Much work remains in defining the relationship between the microbiota and the brain, and several key questions are raised by the work presented here. Of primary interest is the directionality of the interaction between the brain and gut microbes in healthy humans. The concept that the gut microbiota can drive affective response is supported by preclinical work showing affect changes in rodents after manipulation of gut microbes using antibiotics, probiotics, or fecal transplants, and is also suggested by the changes exhibited in human brain networks after chronic probiotic ingestion (18, 61–63). However, the human gut microbiota is frequently affected by the intake of antibiotics, dietary changes, and other interventions without noting changes in mood or affective response. The clinical studies which have shown that such interventions affect mood symptoms in humans have shown very modest changes (14, 64–66). The alternate, more plausible hypothesis is that the brain influences gut microbiota via the autonomic nervous system, stress hormones, and centrally mediated immune modulation. Therefore, the same central drivers that create negative affect may also modulate the gastrointestinal milieu which secondarily shapes microbial composition. Whether the microbiota-brain connections we see in this study reflect the modulation of the gut microbiota structure by the brain, or the influence of the microbiota on the brain and its affective responses, cannot be answered in this cross-sectional view. Careful longitudinal studies, including assessment of gut microbial community structure and microbial metabolomics, in conjunction with neuroimaging and behavioral testing is required to establish directionality and causality.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations

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Figure 1. Identification of bacterial clusters

Plot of the relative abundance of Bacteroides and Prevotella are shown for each cluster, Bacteroides-high (N=32) and Prevotella-high (N=7).

Anatomical Brain Signature

Figure 2. Classification of the white matter connections based on the microbiota group clusters Depicts the discriminative abilities of the white-matter classifier based on the bacterial clusters with an overall accuracy of 66.7%. The samples from each group are connected based on the centroid.

Morphometric Brain Signature

Figure 3. Classification of the gray matter connections based on the microbiota group clusters Depicts the discriminative abilities of the gray-matter classifier based on the bacterial clusters with an overall accuracy of 87.2%. The samples from each group are connected based on the centroid.

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Mean valence and arousal of IAPS pictures based on means from adult women. Mean valence and arousal of IAPS pictures based on means from adult women.

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Table 2

Mean and standard deviation (SD) PANAS scores for microbiota based clusters are shown at baseline, after viewing positive IAPS images and after Mean and standard deviation (SD) PANAS scores for microbiota based clusters are shown at baseline, after viewing positive IAPS images and after viewing negative IAPS images. viewing negative IAPS images.

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Table 3

Brain signature regions: white matter **Brain signature regions: white matter**

Brain region connections with positive loading scores are stronger in the Prevotella-high group. Brain region connections with positive loading scores are stronger in the Prevotella-high group. This table shows the white-matter connections differentiating the microbiota-based clusters. This table shows the white-matter connections differentiating the microbiota-based clusters.

Table 4

Brain signature regions: gray matter **Brain signature regions: gray matter**

This table shows the gray-matter regions differentiating the microbiota-based clusters. This table shows the gray-matter regions differentiating the microbiota-based clusters. Brain regions with positive loading scores are stronger in the Prevotella-high group. Brain regions with positive loading scores are stronger in the *Prevotella*-high group.

