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Association between microbiome and the development of adverse posttraumatic neuropsychiatric sequelae after traumatic stress exposure

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Patients exposed to trauma often experience high rates of adverse post-traumatic neuropsychiatric sequelae (APNS). The biological mechanisms promoting APNS are currently unknown, but the microbiota-qut-brain axis offers an avenue to understanding mechanisms as well as possibilities for intervention. Microbiome composition after trauma exposure has been poorly examined regarding neuropsychiatric outcomes. We aimed to determine whether the gut microbiomes of trauma-exposed emergency department patients who develop APNS have dysfunctional gut microbiome profiles and discover potential associated mechanisms. We performed metagenomic analysis on stool samples (n = 51) from a subset of adults enrolled in the Advancing Understanding of RecOvery afteR traumA (AURORA) study. Two-, eight- and twelve-week post-trauma outcomes for post-traumatic stress disorder (PTSD) (PTSD checklist for DSM-5), normalized depression scores (PROMIS Depression Short Form 8b) and somatic symptom counts were collected. Generalized linear models were created for each outcome using microbial abundances and relevant demographics. Mixed-effect random forest machine learning models were used to identify associations between APNS outcomes and microbial features and encoded metabolic pathways from stool metagenomics. Microbial species, including Flavonifractor plautii, Ruminococcus gnavus and, Bifidobacterium species, which are prevalent commensal gut microbes, were found to be important in predicting worse APNS outcomes from microbial abundance data. Notably, through APNS outcome modeling using microbial metabolic pathways, worse APNS outcomes were highly predicted by decreased L-arginine related pathway genes and increased citrulline and ornithine pathways. Common commensal microbial species are enriched in individuals who develop APNS. More notably, we identified a biological mechanism through which the gut microbiome reduces global arginine bioavailability, a metabolic change that has also been demonstrated in the plasma of patients with PTSD.

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INTRODUCTION

Adverse posttraumatic neuropsychiatric sequelae (APNS) such as posttraumatic stress disorder (PTSD), depression, and somatic symptoms are common after traumatic stress exposure [1–6]. Contemporary limitations in understanding the pathogenesis of APNS are a barrier to developing effective primary and secondary preventive interventions [1, 7]. Novel multidisciplinary approaches applying methodologic advances from other areas of neuroscience to the early post-trauma period may advance understanding of APNS pathogenesis.

One such area is the study of the microbiome-gut-brain axis. The microbiome-gut-brain axis has been demonstrated to have an important influence on brain function in neuropsychiatric disorders [8–10], and has been hypothesized to play an important

role in trauma-related neuropsychiatric disorders [11]. Gut microbes affect central nervous system function through the production of metabolites and neurochemicals [10]. For example, short-chain fatty acids such as butyrate support gut homeostasis by maintaining gut barrier integrity and inducing antiinflammatory factors which cross the blood-brain barrier [12, 13]. In contrast, *Enterobacteriaceae* expansion can augment neuroin-flammatory processes through the lipopolysaccharide (LPS) endotoxin-mediated activation of the Toll-like receptor 4 (TLR4) pathway, leading to the production of pro-inflammatory cytokines [14].

While specific mechanisms responsible for it remain incompletely understood, dysregulated immunity and elevated inflammation are risk factors for PTSD [15]. Given the above known

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influence of the gut-brain axis on neuroinflammation, these data support the hypothesis that variations in microbiome characteristics could influence APNS pathogenesis. This hypothesis is also supported by several small studies which identified broad phylum-level differences in microbiome characteristics among individuals with PTSD (n = 18) as compared to those without PTSD (n = 12) [16], and in veterans which found associations between lower diversity and a higher abundance of opportunistic microbes with PTSD symptoms and impaired cognition [17]. This hypothesis is also supported by animal studies in which trauma was found to cause alterations in the microbiome and lead to increased local and systemic inflammation [8].

In this nested study, we explored the association between the gut microbiome and APNS development in a subsample of individuals recruited from emergency departments (ED) in the immediate aftermath of trauma as part of the AURORA study (n = 74) and who also provided stool samples [1]. We employed shotgun metagenomic sequencing of stool samples from patients evaluated and discharged from the ED and machine learning (ML) predictive analytics to test the hypothesis that variability in microbiome species and pro- and anti-inflammatory metabolic pathways is associated with APNS outcomes.

PATIENTS AND METHODS

Study Population and Sample Collection

The AURORA study began in September 2017, we pre-registered a nested microbiome pilot study with the AURORA consortium and enrolled participants from November 2020 until February 2021 (Supplemental table 1) [18]. AURORA participants who completed their first outpatient remote assessment and had not received antibiotics within the previous 6 months were approached for enrollment into this nested study. Participants also could not have had any other influences that could cause major microbiome perturbations (e.g., infections). Of the 2,097 AURORA participants subjects approached, 106 provided consent to participate, and we received stool samples from n = 74 of these individuals (69.8%). Stool samples were self-collected at home using OMNIgene-GUT collection kits (DNAgenotek, catalog no. OMR-200). All stool samples were self-collected at least five days after the initial ED presentation (Supplemental figure 1). Upon receipt, samples were stored at -80 °C until DNA extraction and sequencing were performed. This study was approved by the University of North Carolina's institutional review board (IRB protocol #17-0703) and by the UMass Chan Medical School's institutional review board.

Outcome Measurements

After receiving written informed consent from eligible patients, study coordinators from each participating ED performed data collection for EDbased assessments, including baseline questionnaires. Follow-up evaluations were internet-based at the two-, eight-, and twelve-week intervals. We assessed posttraumatic stress symptoms using the Post Traumatic Stress Disorder (PTSD) checklist for DSM-5 (PCL-5) [19-22]. Per this instrument's guidelines, a PTSD diagnosis can be made by summing the self-reported responses to the PCL-5. We used total sum of 31 for a positive diagnosis of PTSD [23]. A normalized t-score for depression severity and the diagnosis was assessed using the PROMIS Depression Short Form 8b [24]. A t-score of 60 is one standard deviation lower than average and was used to designate a depression diagnosis. The Rivermead Post-concussive questionnaire (RPQ) was used to obtain a count of somatic symptoms. The twelve post-traumatic somatic symptoms included in the RPQ are headache, dizziness, nausea, noise sensitivity, fatigue, insomnia, poor concentration, taking longer to think, blurred vision, light sensitivity, double vision, and restlessness [25]. For each symptom, a yes/no variable was created. A positive response to a symptom would be counted as YES = 1 that it does exist, while a NO = 0 indicates that the symptom does not exist. The yes/no responses were summed to provide a simple count of somatic symptoms.

Sample Processing and DNA Sequencing

Prior to extraction, stool samples were heat inactivated at 65 °C-70 °C for 1 hour and stored at -80 °C. Approximately 250 mg of resulting sample was extracted using the QIAGEN DNeasy PowerSoil Pro Kits (QIAGEN, catalog no. 47016). Sequencing libraries were prepped using the Nextera XT DNA library prep kit and sequenced on a NextSeq 500 sequencer with 2×150 bp paired-end reads. Of the 74 stool samples received, 69 (93.2%) were successfully sequenced. Samples with a percentage of reads identified greater than 0.5%, and complete metadata were used for analysis (n = 51).

Shotgun metagenomic reads were trimmed and filtered for host contamination using the KneadData pipeline (https://github.com/biobakery/kneaddata). The resulting metagenomic data was profiled for microbial abundance and metabolic pathways using the MetaPhIAn3 [26] and HUMANN3 databases and tools [27]. The resulting relative microbial species abundance and metabolic pathway data was used for downstream analysis in R.

Microbiome Analysis and Statistics

Power Analysis. We performed a power analysis to estimate the study size according to a one sample Spearman correlation. Spearman correlation is a non-parametric test and is thus suitable for microbiome data. With a sample size of 51 a correlation of 0.5 (a medium effect size), the power was found to be 0.97. This analysis was performed in R using the pwr package (https://github.com/heliosdrm/pwr).

Linear mixed model-based analysis to determine the relative contribution of the microbiome. To determine how much of the variability in post-trauma APNS outcomes is explained by the gut microbiome, we first constructed linear mixed models (LMM) using the Ime4 R package [28] for each of the three APNS outcomes of interest (PTSD, Depression Scores and Somatic Symptoms) as a function of relevant clinical covariates (sex, age, body mass index, race/ethnicity) [29], and also of the arcsine square root-transformed abundance [30] of each microbial species independently. Features identified as significant (FDR corrected p-value <= 0.05) were combined and used to fit a global LMM. The contribution of each feature to the regression line fit by the model was determined by running analysis of variance (ANOVA) and examining the total sum of squares for the fixed effects. The resulting sum of squares ratio for each feature was graphed using WebR (https://github.com/cardiomoon/webr).

Mixed-effect random forest analysis of microbiome permutated importance to outcomes at twelve weeks post-trauma. Microbiome data is non-linear and not normally distributed [31, 32]. Additionally, such data are characterized by many predictors, which, if combined in the same model (without prefiltering), can prohibit a traditional regression model to converge [33]. We have previously demonstrated that tree-based machine learning (ML) approaches such as random forest, which are non-parametric and perform intrinsic feature selection, are very powerful in finding a signal from static and time-resolved microbiome data [34-37]. To fully utilize the longitudinal clinical data collected by the AURORA (parent) study for this nested study [1], we assumed, barring perturbation, the gut microbiome is stable over time. Several high-resolution gut microbiome temporal studies have demonstrated that the microbiome is rather stable and displays only small random fluctuations [38-41]. The extent to which the microbiome is stable in individuals that have experienced trauma remains to be established. We built a mixed-effect random forest (MERF) regression model to predict either PTSD raw score, Depression t-score, or somatic symptoms count at twelve weeks post-trauma. We use either microbial abundance or metabolic pathway abundance and relevant clinical covariates (e.g., BMI, age, sex, and race) as variables in this modeling. The first step of our pipeline split our data into a training and test set. To predict twelve-week post-trauma outcomes, two-week and eight-week post-trauma outcomes were used to train the MERF model. The unseen twelve-week data was used for testing the model. For each APNS outcome, the pipeline was run ten times with ten different random seeds, and model performance, statistics, and outcomes were calculated for each seed. Model performance was evaluated by Root Mean Square Error (RMSE) and correlation of true versus predicted values, which illustrates the model's accuracy and fit for predicting outcome measurements at twelve weeks post-trauma (Supplemental figure 2). Permuted variable importance was calculated and used to evaluate models. Plots summarizing results were generated in R using the ggplot2.

RESULTS

Characteristics of Study Subjects

Among 2,097 individuals who were approached in the ED regarding this nested study, 106/2,097 (5.1%) agreed to participate, and 74/106

		Depression			PTSD		
Patient Characteristics	Overall, N = 51 ^ª	Depression-, N = 35 ^a	Depression $+$, N = 16 ^a	q-value (adjusted p-value ^b) ^c	PTSD-, N = 31 ^a	$PTSD + , N = 20^{a}$	q-value (adjusted p-value ^b) ^c
Demographics							
Age in years (at enrollment)	52 (36, 60)	52 (36, 59)	52 (37, 60)	>0.9	52 (36, 58)	50 (34, 61)	>0.9
Sex (Female)	26 (51%)	19 (54%)	7 (44%)	>0.9	19 (61%)	7 (35%)	0.7
BMI	29 (24, 33)	27 (24, 30)	33 (29, 40)	0.11	27 (23, 30)	31 (29, 36)	0.083
White	31 (61%)	21 (60%)	10 (62%)	>0.9	18 (58%)	13 (65%)	>0.9
Black	13 (25%)	11 (31%)	2 (12%)	>0.9	9 (29%)	4 (20%)	>0.9
Asian	1 (2.0%)	1 (2.9%)	0 (0%)	>0.9	1 (3.2%)	0 (0%)	>0.9
other	6 (12%)	2 (5.7%)	4 (25%)	0.8	3 (9.7%)	3 (15%)	>0.9
Hispanic	7 (14%)	4 (11%)	3 (19%)	>0.9	4 (13%)	3 (15%)	>0.9
U.S. Geographic Region				>0.9			>0.9
Midwest Region	10 (20%)	5 (14%)	5 (31%)		5 (16%)	5 (25%)	
Northeast Region	38 (75%)	27 (77%)	11 (69%)		24 (77%)	14 (70%)	
Southern Region	3 (5.9%)	3 (8.6%)	0 (0%)		2 (6.5%)	1 (5.0%)	
Trauma Event Type (Broad)				>0.9			>0.9
Animal-related	1 (2.0%)	1 (2.9%)	0 (0%)		1 (3.2%)	0 (0%)	
Fall < 10 feet or from unknown height	9 (18%)	7 (20%)	2 (12%)		6 (19%)	3 (15%)	
Fall ≥ 10 feet	4 (7.8%)	2 (5.7%)	2 (12%)		2 (6.5%)	2 (10%)	
Incident causing traumatic stress exposure to many people	1 (2.0%)	1 (2.9%)	0 (0%)		1 (3.2%)	0 (0%)	
Motor Vehicle Collision	28 (55%)	19 (54%)	9 (56%)		17 (55%)	11 (55%)	
Non-motorized Collision	3 (5.9%)	2 (5.7%)	1 (6.2%)		2 (6.5%)	1 (5.0%)	
other	5 (9.8%)	3 (8.6%)	2 (12%)		2 (6.5%)	3 (15%)	
Self-Reported Perceived Chance of Dving	6.0 (1.0, 8.5)	6.0 (1.0, 8.5)	5.5 (1.8, 8.2)	>0.9	6.0 (0.5, 8.5)	5.5 (1.8, 8.5)	>0.9

Table 1. Patient Characteristics by APNS outcome category.

^an (%); Median (IQR)

^bPearson's Chi-squared test; Fisher's exact test; Wilcoxon rank sum test

^cBonferroni correction for multiple testing

(69.8%) provided a stool sample. Of the stool samples received, 51(68.9%) passed all quality and metadata checks (see methods) and were profiled for microbial abundance and metabolic pathways. From these 51 participants (mean age 52 years; 26 (51%) female) stool samples were received a median of 45 days (range 5 to 182 days) (Supplemental figure 1) after ED visit for trauma evaluation that conferred entry to the AURORA (parent) study (Table 1). The most common types of trauma were motor vehicle collision (55%, n = 28), fall from height (n = 9, 18%), and other accidental or targeted/ involuntary events (9.8%, n = 5). Stool samples originated primarily from White (61%, n = 31) and Black (25%, n = 13) participants. Of the 31 white participants, 29 (94%) were non-Hispanic White. Microbiome diversity measures show that individuals do not stratify based on alpha (Simpson and Shannon) or beta diversity.

Gut Microbiome Features associated with APNS

Linear mixed effect models and ANOVA adjusting for sociodemographic characteristics were used to assess for associations between microbiome characteristics and APNS symptom severity. Gut microbiome characteristics accounted for 48%, 26%, and 44% of the variation in PTSD, depressive, and somatic symptoms after trauma, respectively (Fig. 1). The abundance of *Firmicutes bacterium CAG:555, Bifidobacterium adolescentis,* and the proinflammatory *Streptococcus infantis* were associated with PTSD symptom severity. The abundance of *B. adolescentis* was associated with depressive symptom severity (Fig. 1B).

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used to identify microbiome features associated with PTSD, depression, and somatic symptom burden at twelve weeks posttrauma (Supplemental figure 2). Our findings using this approach confirmed and expanded on findings from the linear mixed-effect models. Bacterial species identified in linear mixed effects modeling, specific species of the *Alistipes, Bifidobacterium*, and *Ruminococcus* genera were also found as important predictors by the MERF model (Fig. 2). The models also identified more species

Ruminococcus gnavus and Streptococcus parasanguinus were

While the above LMM approach is useful to assess total variance

accounted for by microbiome characteristics and is common in the

field, it has several limitations, including the inability to evaluate

microbial abundances in the context of the entire microbiome and

the limited ability to account for inter-individual microbiome

differences [33]. Therefore, we used a complementary machine

learning approach to train and test models examining associations

between gut microbiome characteristics and posttraumatic out-

comes. We have previously used this approach and have determined its optimality for inferring biologically relevant host-microbe inter-

actions from cross-sectional and longitudinal microbiome data (see

A Mixed-Effect Random Forest (MERF) regression model was

methods sections for further justification) [34-37, 42-44].

Post-trauma neuropsychiatric outcomes are predicted by

associated with somatic symptom severity (Fig. 1C).

microbial abundance



Fig. 1 Contribution of microbiome features significantly associated with neuropsychiatric outcomes. After metagenomic sequencing, microbial species abundances were combined with demographics and select clinical variables (other) in linear mixed effect models for the APNS outcomes at two-, eight-, and twelve weeks post trauma (A) PTSD raw score as determined by the DSM-5 PTSD checklist, (B) depression score as determined by the PROMIS Depression Short Form 8b and (C) somatic symptoms count (Yes/No) based on the Rivermead Post-Concussive Questionnaire. Individual microbial species, demographics, and clinical variables found as significant are displayed in the outer wheels. PTSD time point reflects the time at which the modeled score was taken (either two weeks, eight weeks, or twelve weeks). The length of time between trauma exposure and stool sample collection was included as days since trauma in all models.

as contributors to each outcome, with many similar species in the top 15 predictors identified by the three models. *B. adolescentis, B. longum*, and *Flavonifractor plautii* were among the top five predictors for all three APNS outcomes (Fig. 2). Across our models, both *Bifidobacterium* species and *F. plautii* showed similar trends of increased abundance being informative of higher scores (Supplemental figure 3). *Ruminococcus gnavus* was found to be among the top 15 predictors only for PTSD (ranked 7th) and depression (ranked 14th), with increases in abundance correlating with disease outcomes (Fig. 2, Table 2). Decreases in certain species were noted to cluster by individual outcomes for the most part (Fig. 2D).

Metabolic Pathway Profiling Identifies Novel Gut-Brain-Axis Interface with APNS Outcomes

Species abundance, although informative of outcomes, does not directly provide context for the functional roles of associated microbes. By analyzing metabolic pathway abundances encoded in the metagenomic data, we can gain insight into the microbial products, metabolites, and functions that may associate with each outcome of interest, thus shedding light on possible mechanistic links. To this end, we employed the same MERF-based pipeline to examine associations between the abundance of microbiallyencoded metabolic pathways and PTSD, depressive, and somatic symptom severity twelve weeks after trauma (Supplemental figure 4). The Calvin-Benson-Bassham cycle, a common CO₂ fixation pathway in autotrophic bacteria, was identified as one of the top three predictors of PTSD and depressive symptom severity (i.e., cycle was reduced in samples with higher scores (Fig. 3A and B)). Amino acid biosynthesis pathways were a leading predictor for all three APNS outcomes, with the L-citrulline biosynthesis pathway identified as one of the top two predictors for all APNS outcomes. The super pathway of arginine and polyamine biosynthesis was identified as the top predictor for the PTSD model and as the third most important pathway for the depression model, correlating negatively with PTSD and depression scores (Fig. 4, Table 3). Furthermore, all three models identified pathways involving arginine, ornithine, and citrulline, three amino acids that are often interconverted (Fig. 5). De novo biosynthesis of L-ornithine, which can be interconverted from ornithine and urea to arginine and water, was identified by all models and appeared increased in patients with higher scores. Increased abundance of L-citrulline biosynthesis was also associated with higher scores of PTSD, depression, and somatic symptoms count (Fig. 4).

Given the strength of findings around pathways affecting global arginine levels, we next sought to identify microbes responsible for changes in these pathways. Focusing on ornithine, citrulline, and arginine, we examined the contribution of individual genera to each pathway (Tables 4, 5, 6) to PTSD and depression. Somatic symptoms were not further analyzed as the simple count nature of the variable was not appropriate for this analysis. We observed that in individuals with PTSD or depression, Escherichia had an increased contribution to all pathways involving arginine and ornithine biosynthesis. However, the contribution of E. coli to L-citrulline biosynthesis was decreased in those with PTSD and increased in those with depression. In patients with PTSD or depression, the Ruminococcus genus had a decreased contribution to L-arginine biosynthesis via both L-ornithine and the acetyl cycle. Alistipes, Flavonifractor and Faecalibacterium genera contributed more to the L-arginine biosynthesis via ornithine in individuals with diagnosed PTSD or depression.

DISCUSSION

We have demonstrated here that gut microbiome characteristics of trauma survivors were associated with the development of APNS, in this pilot longitudinal study of individuals enrolled in the ED after trauma exposure. Our analysis first suggested that overall inter-individual differences in gut microbiome taxonomy accounts for a substantial fraction (20-48%) of the differences in APNS outcomes. This finding is similar to what is attributable to known APNS outcome-associated clinical and demographic covariates (e.g., sex and race/ethnicity) [45]. We leveraged ML modeling to identify specific bacterial species and microbial encoded metabolic pathways with previously established proand anti-inflammatory properties that are predictive of APNS development. Our analysis highlights the most prevalent APNS outcome-associated gut microbiome-encoded pathways are those leading to the biosynthesis of arginine, ornithine, and citrulline. These pathways can possibly affect global arginine levels in the body, a biomarker that has been previously associated with PTSD [46]. To our knowledge, ours is the first study to longitudinally evaluate the link between the gut

Fig. 2 Mixed-Effect Random Forest (MERF) Regression Models Using Microbial Abundance and Clinical Covariates to Predict Neuropsychiatric Outcomes. MERF models combining microbial abundance data with clinical and demographic (highlighted in red) features demonstrate the importance of microbial features in predicting outcomes. Permutated importance analysis of model outcomes shows the top 15 features contributing to predictions of (A) PTSD raw score, (B) depression normalized score, and (C) somatic symptoms count (yes/no) are mostly microbial species. D Heat map showing correlation coefficients of the top 30 contributing species show common species that contribute to all three outcomes (increased abundance) and species which associate with individual outcomes (mostly decreased abundance).

microbiome and APNS outcomes while providing mechanistic links along the microbiome-gut-brain axis.

Our initial analyses by simple linear mixed-effect models demonstrated the significant contribution of microbiota to APNS outcomes in comparison to clinical covariates and demographic characteristics. The proportion of the gut microbiome accounting for variability in each APNS outcome (26-48%) is of the same order of magnitude as demographic characteristics (10-56%) that are already known to associate with each outcome [45]. However, this analysis required transformation of microbiome abundance variables and the use of a two-tiered approach to select

outcome-associated features, which is suboptimal. Although common, this approach neither accounts for the inter-personal variation of the microbiome nor evaluates the effect of microbial species together [47].

While there is evidence for dramatic changes in the gut microbiome after trauma within 72 hours [48], almost no research describing long-term changes in gut microbiota after acute trauma currently exists. One study on PTSD in frontline healthcare workers suggests that long-term gut microbiome dysbiosis induced by stress is sustained for months and predisposes individuals to recurring PTSD [49]. Thus, we built our pipeline

Table 2.	Spearman's	Correlation	of Im	portant Microbes	with	APNS Outcomes.
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Predictors	PTSD	Depression	Somatic Symptoms	Notes
Bifidobacterium adolescentis	0.286	0.216	0.0704	 Produces GABA in the gut [69] Induction of Th17 cells in mice [70] Decreased in Crohn's Disease [71] Lower abundance in children with autism spectrum disorder [72]
Bifidobacterium longum	0.34	0.259	0.251	 GABA production in vivo [73] Anxiolytic effect through the vagus nerve in mice [57] Lower abundance in children with autism spectrum disorder [72] Strain specific induction of cytokines in peripheral blood mononuclear cells [74]
Bifidobacterium bifidum	0.185	NA	NA	 Th17 inducing profile [75] Lower abundance in children with autism spectrum disorder [72]
Parabacteroides merdae	0.176	0.226	NA	 Indole negative [76] Rarely associated with infections[77] Branched-chain amino acid catabolism by <i>P. merdae</i> reduces atherosclerotic lesions [78]
Flavonifractor plautii	0.207	0.177	0.173	 In mice, suppresses Th2 immune response [79] [80] Genera found to be enriched in those with active major depressive disorder (MDD) [63]
Adlercreutzia equolifaciens	0.0195	NA	NA	Strain specific equal production [81]
Ruminococcus gnavus	0.441	0.335	NA	 Increased in Crohn's Disease [71] Depleted genera level in active MDD [63]
Alistipes finegoldii	0.198	NA	ΝΑ	 Can hydrolyze tryptophan to indole; Succinic acid producer in vitro with minor acetic and propionic acid; bile-resistant, esculin-negative; catalase-negative; nitrogen-reductase negative [82, 83] Enriched in colorectal cancer [84] Main cellular fatty acid 13-metyltetradecanoic acid (iso-C15:0) Genera Increased abundance in active MDD patients compared to healthy controls [63]
Eubacterium eligens	-0.216	NA	-0.14	 Decreased in COVID-19 health care workers with higher stress scores [49] Anti-inflammatory related [85]
Firmicutes bacterium CAG 110	-0.182	NA	NA	Identified in livestock [86]
Alistipes indistinctus	-0.0339	ΝΑ	ΝΑ	 Unable to hydrolyze tryptophan to indole; susceptible to bile; catalase-positive; urease and nitrogen reductase-negative; Succinic and acetic acid producer in vitro [83] Main cellular fatty acid 13-metyltetradecanoic acid (iso-C15:0) Genera Increased abundance in active MDD patients compared to healthy controls [63]
Bacteroides massiliensis	-0.0271	NA	NA	 Phocaeicola massiliensis Genera increased in health controls vs. active MDD patients [63]
Slackia isoflavoniconvertens	-0.221	NA	NA	Capable of equol production [87]
Desulfovibrio piger	NA	NA	-0.183	Sulfur-reducing bacteria associated with inflammatory bowel disease [88]
Blautia wexlerae	NA	NA	0.0584	 Inversely correlated with obesity and type 2 diabetes mellitus [89] Decreased abundance in progressive multiple sclerosis [90]
Intestinimonas butyriciproducens	NA	NA	-0.155	Butyrate production [91]
Bacteroides thetaiotaomicron	NA	NA	-0.172	• Genera increased in healthy controls vs. active MDD patients [63]
Parabacteroides goldsteinii	NA	NA	-0.155	• Anti-inflammatory and decreased in chronic inflammatory diseases including chronic obstructive pulmonary disease (COPD; negatively associates with severity) [92] and chronic kidney disease [93]

Table 2. Continued				
Predictors	PTSD	Depression	Somatic Symptoms	Notes
Pseudoflavonifractor sp An184	NA	NA	-0.215	 Positively associated with weight loss [94]
Streptococcus thermophilus	NA	NA	0.0866	
Clostridium symbiosum	NA	0.204	0.163	 Increased abundance in early colorectal cancer [95]
Lactobacillus acidophilus	NA	NA	-0.21	
Bacteroides vulgatus	NA	0.115	NA	• Genera increased in healthy controls vs. active MDD patients [63]
Anaerostipes hadrus	NA	-0.0635	NA	
Dialister invisus	NA	0.0954	NA	 [80] Genera increased in healthy controls vs. active MDD patients [63]
Bacteroides ovatus	NA	0.194	NA	
Bilophila wadsworthia	NA	0.274	NA	
Eubacterium ramulus	NA	-0.096	NA	
Roseburia hominis	NA	0.102	NA	
Lawsonibacter asaccharolyticus	NA	-0.0814	NA	

Fig. 3 Mixed-Effect Random Forest (MERF) Regression Models Using Microbial Metabolic Pathways and Clinical Covariates to Predict Neuropsychiatric Outcomes. MERF models combining microbial metabolic pathway abundance data with clinical and demographic (highlighted in red) features demonstrate the importance of microbial pathways in predicting outcomes for (A) PTSD raw score, (B) depression normalized score, and (C) somatic symptoms count (yes/no). The top 15 features from analysis of permutated importance on model outcomes are displayed for PTSD and depression. Our model found only seven significant features for somatic symptoms.

assuming the microbiome at our sampling point is representative of the patient's microbiome at the longitudinal post-trauma outcome measurement points, which we hypothesize associates with target outcomes of developing APNS in prolonged periods after trauma.

ML models such as Mixed-Effect Random Forest (MERF) models are capable of accounting for interpersonal variation, evaluating microbial abundances in the context of the entire microbiome, and, importantly, because no scaling is required, it can be used with different data types (e.g., categorical, numerical, proportion, etc.) [33]Our MERF models identified more species as significant contributors to our outcomes than the linear models and microbes previously associated with neuropsychiatric disorders, specifically *Bifidobacterium*, *Alistipes*, and *Flavonifractor* species. Further, the model identified *F. plautii, E. eligens, E. ramulus, R. homins, C. sybiosum*, and *Blautia* species which are members of the *Clostridium* Cluster IV and XIVa, two clusters of known beneficial bacteria for inflammatory bowel disease and gut health [50–52]. Several of these are short-chain fatty acid producers and have been associated with reduced inflammation in humans [53, 54]. Additionally, they are known to promote several antiinflammatory immune signatures, such as regulatory T-cell expansion in vivo [55, 56].

Our models based on species abundance identified *Bifidobacterium* species and *Flavonifractor plautii* as the most important factors for predicting all three APNS outcomes. Specifically, we found higher abundances of *B. adolescentis, B. longum*, and *B. bifidum* were associated with higher PTSD and depression scores, which contradicts some literature regarding positive relationships between *Bifidobacterium* spp. and improved neuropsychiatric outcomes [57–59]. Contrary to this, many other microbiome-based studies of neuropsychiatric disorders, including major depressive disorder (MDD), schizophrenia, and anxiety, have failed to find similar associations [60–62]. Indeed, for MDD, various

Tahla 2

continued

Fig. 4 Important metabolic pathways for PTSD and depression involve arginine, citrulline and ornithine. Heat map (A) of correlation coefficients for the top 15 metabolic pathways contributing to PTSD, depression, and somatic symptoms from MERF analysis show significant contributions by amino acid and polyamine biosynthesis pathways. **B** Violin plots showing differences in contributions of arginine, citrulline, and ornithine biosynthesis pathways for depression (top graphs) and PTSD (bottom graphs). Blue violins indicate no diagnosis (PTSD raw score \leq 31; Depression T-Score < 60 indicating none to mild depression) red indicates PTSD or depression diagnosis (PTSD RS > 31; Depression T-Score \geq 60 indicating moderate to severe depression).

investigators have found both increased and decreased abundances of *Bifidobacterium* spp. to be associated with clinical disease suggesting possibly strain specific effects [50, 51, 62]. Our analysis identifying increased abundances of *Bifidobacterium* spp. as being associated with APNS outcomes adds more contradiction to the literature but highlights the importance of looking beyond species abundances in microbiome studies.

Alone, species-based associative studies can be confounding and limit clearer investigations into biological mechanisms. We have previously shown that also analyzing microbial metabolic pathways can reveal valuable insight into possible biologic impacts of the gut microbiome on clinical outcomes [35, 42]. As with our previous work, our combined analysis approach here, leveraging metagenomic sequencing and ML-based pipelines on both species' abundance and metabolic pathways, identified a microbially-based signature within the biosynthesis of arginine/ citrulline/ornithine as highly informative of the development of all three APNS outcomes. This is the first study to support the existence of microbially produced metabolites acting through the microbiome-gut-brain axis with APNS attributable outcomes.

PTSD patients have been shown to have decreased levels of arginine and increased levels of ornithine and citrulline in

Table 3.	Spearman's	Correlation of	Important	Metabolic	Pathways	with	Outcomes
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Predictors	PTSD	Depression	Somatic Symptoms	Notes
ARG-POLYAMINE-SYN: superpathway of arginine and polyamine biosynthesis	-0.251	-0.229	NA	• Enriched patients with Crohn's Disease (CD) in remission [96]
CITRULBIO-PWY: L-citrulline biosynthesis	0.346	0.298	0.261	 Enriched patients with CD in remission [96]
CALVIN-PWY: Calvin Benson Bassham cycle	-0.269	-0.315	NA	 Enriched patients with CD in remission [96]
ARGININE-SYN4-PWY: L-ornithine de novo biosynthesis	0.201	0.165	0.139	
CENTFERM-PWY: pyruvate fermentation to butanoate	-0.322	-0.26	-0.168	
BIOTIN-BIOSYNTHESIS-PWY: biotin biosynthesis I	0.181	0.127	NA	
BRANCHED-CHAIN AA-SYN-PWY: superpathway of branched amino acid biosynthesis	-0.127	-0.00818	NA	
ANAEROFRUCAT-PWY: homolactic fermentation	-0.0199	NA	NA	
ASPASN-PWY: superpathway of L-aspartate and L-asparagine biosynthesis	-0.0404	0.0571	0.037	
COA-PWY-1: coenzyme A biosynthesis II	-0.0454	NA	-0.053	
ARGSYNBSUB-PWY: L-arginine biosynthesis II (acetyl cycle)	-0.0768	-0.131	NA	
ARGSYN-PWY: L-arginine biosynthesis I (via L-ornithine)	-0.0579	-0.105	NA	
1CMET2-PWY: N ¹⁰ formyl tetrahydrofolate biosynthesis	NA	-0.0885	NA	Depleted in Ulcerative Colitis (UC) in remission [96]
ANAGLYCOLYSIS-PWY: glycolysis III (from glucose)	NA	-0.048	NA	
ARGORNPROST-PWY: arginine, ornithine and proline interconversion	NA	0.0264	NA	

Fig. 5 Arginine is converted into citrulline and ornithine commonly in the host and the microbiome. The ratio of arginine to citrulline and ornithine in PTSD patients has previously been found to negatively correlated with PTSD. We find that this negative correlation in the metabolic pathways encoded for in the microbiome of trauma survivors.

peripheral blood [46]. The ratio of arginine to its two main catabolic products, ornithine, and citrulline, is used as a readout of nitric oxide (NO) capacity and is referred to as the global arginine bioavailability ratio (GABR) [52]. Bersani et al. sought to examine NO production in PTSD patients via the GABR as arginine is the sole nitrogen source of NO synthesis. They found that GABR was negatively correlated with PTSD, which they reasoned was indicative of dysfunctional NO synthase. The directionality observed by Bersani and authors mirrors the directionality of arginine-related metabolic pathways in our analysis. Altered arginine metabolism has also been implicated in other neuropsychiatric disorders, such as schizophrenia and MDD [53, 54]. The influence of altered arginine metabolism is likely multidimensional and spans multiple mechanisms of action, including through arginine vasopressin or NO [55, 56]. Identifying microbial sources of these potential mechanisms in specific neuropsychiatric conditions is an important first step towards enabling research on interventions or therapies.

In our species trained MERF pipeline, *Flavonifractor plautii*, a flavonoid-degrading bacterium often found in the human gut microbiome, was the only species among the top 5 contributors to all three APNS outcomes that affect GABR. This genus contributes to the L-arginine biosynthesis pathway via ornithine in those with PTSD and via acetyl in those with PTSD or depression. From our metabolic pathway models, the contribution of *Ruminococcus*, *Alistipes* and *Flavonifractor* genera to L-arginine biosynthesis via ornithine was identified as increased in individuals with PTSD or depression. Although the contribution of these three species were not ranked as highly as *Bifidobacterium* spp. in our species trained MERF model, by analyzing microbial metabolic pathways, we see that their roles in PTSD and depression may be far more significant.

Our findings also provide an added layer of contextual insight into seemingly contradictory findings from prior research on this topic. In addition to being increased in Crohn's disease, the Ruminococcus genera have been identified as decreased in those with MDD in multiple studies [59, 63]. Although we find R. gnavus abundance increases with depression score, the contribution of the Ruminococcus genera to arginine biosynthesis through the acetyl cycle (ARGSYNBSUB-PWY) and via ornithine (ARGSYN-PWY) was lower in those with depression compared to those without, implying species specific effects may also be at work. Likewise, Alistipes have been found to be both increased [63] and decreased in patients with MDD [59, 64] as compared to healthy controls, suggesting genus-based examination of Alistipes may not be sufficient. A. finegoldii and A. indistinctus, were both identified by our microbial model as informative of PTSD score yet with opposite directions. Furthermore, A. finegoldii abundance was positively associated with PTSD scores and specifically had a

Table 4.	Microbial genus	contribution to	arginine-related	metabolic pathwa	ys stratified b	y PTSD and de	epression diagnoses.
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Pathway	Genus	PTSD Raw Score < 31	PTSD Raw Score≥31	Depression t- score < 60	Depression t- score ≥ 60
ARG-POLYAMINE-SYN:	Escherichia	53.64	69.95	42.86	100.00
Superpathway of arginine and polyamine biosynthesis	Klebsiella	46.36	30.05	57.14	0.00
ARGSYNBSUB-PWY: L-arginine	Akkermansia	7.72	2.36	7.61	1.20
biosynthesis II (acetyl cycle)	Anaerostipes	2.49	0.67	2.29	0.68
	Anaerotignum	0.36	1.14	0.48	1.03
	Bifidobacterium	1.07	1.47	1.01	1.72
	Bilophila	0.52	1.61	0.58	1.75
	Blautia	4.12	4.00	4.46	3.11
	Escherichia	0.97	3.81	1.04	4.37
	Faecalibacterium	33.79	41.04	32.37	46.62
	Flavonifractor	0.79	3.32	1.55	2.06
	Fusicatenibacter	8.76	4.57	8.00	5.40
	Gemmiger	10.68	7.95	10.19	8.46
	Klebsiella	0.96	1.74	1.73	0.00
	Lachnospiraceae unclassified	9.40	8.91	9.14	9.46
	Roseburia	1.42	1.15	1.58	0.68
	Ruminococcaceae unclassified	2.43	1.16	2.55	0.50
	Ruminococcus	4.14	3.45	4.09	3.39
	Tyzzerella	0.00	1.40	0.33	0.94

Table 5. Microbial genus contribution to ornithine-related metabolic pathways stratified by PTSD and depression diagnoses.

ARGININE-SYN4-PWY: L-ornithine de novo biosynthesis Bacteroides 38.78 29.26 37.40 28.63 Catenibacterium 13.22 0.14 7.15 4.90 Escherichia 12.28 25.78 12.20 29.42 Klebsiella 5.21 4.50 8.41 0.00 Parabacteroides 28.82 35.58 33.49 31.06 Paraprevotella 1.68 4.75 1.35 5.99	Pathway	Genus	PTSD Raw Score < 31	PTSD Raw Score≥31	Depression t- score < 60	Depression t- score ≥ 60
L-ornithine de novo biosynthesis Catenibacterium 13.22 0.14 7.15 4.90 Escherichia 12.28 25.78 12.20 29.42 Klebsiella 5.21 4.50 8.41 0.00 Parabacteroides 28.82 35.58 33.49 31.06 Paraprevotella 1.68 4.75 1.35 5.99	ARGININE-SYN4-PWY:	Bacteroides	38.78	29.26	37.40	28.63
Escherichia 12.28 25.78 12.20 29.42 Klebsiella 5.21 4.50 8.41 0.00 Parabacteroides 28.82 35.58 33.49 31.06 Paraprevotella 1.68 4.75 1.35 5.99	L-ornithine de novo	Catenibacterium	13.22	0.14	7.15	4.90
Klebsiella 5.21 4.50 8.41 0.00 Parabacteroides 28.82 35.58 33.49 31.06 Paraprevotella 1.68 4.75 1.35 5.99	biosynthesis	Escherichia	12.28	25.78	12.20	29.42
Parabacteroides 28.82 35.58 33.49 31.06 Paraprevotella 1.68 4.75 1.35 5.99		Klebsiella	PTSD Raw Score < 31 PTSD Raw Score ≥ 31 Depression t- score < 60 Depression t- score < 60 38.78 29.26 37.40 28.63 13.22 0.14 7.15 4.90 12.28 25.78 12.20 29.42 5.21 4.50 8.41 0.00 28.82 35.58 33.49 31.06 1.68 4.75 1.35 5.99 3.21 16.69 3.92 18.18 2.60 0.38 2.40 0.31 1.28 1.47 1.21 1.65 3.95 3.29 4.29 2.46 1.36 3.40 1.45 3.65 1.36 3.40 1.45 3.65 1.36 3.40 1.45 3.65 1.38 3.56 8.30 4.12 1.38 7.32 10.89 7.41 0.55 0.77 0.96 0.00 2 2.06 1.20 2.30 0.55	0.00		
Paraprevotella 1.68 4.75 1.35 5.99		Parabacteroides	28.82	35.58	33.49	31.06
		Paraprevotella	1.68	4.75	1.35	5.99
ARGSYN-PWY: L-arginine Alistipes 3.21 16.69 3.92 18.18	ARGSYN-PWY: L-arginine	Alistipes	3.21	16.69	3.92	18.18
biosynthesis I (via L- Anaerostipes 2.60 0.38 2.40 0.31	biosynthesis I (via L-	Anaerostipes	2.60	0.38	2.40	0.31
Bifidobacterium 1.28 1.47 1.21 1.65	officiality)	Bifidobacterium	1.28	1.47	1.21	1.65
Blautia 3.95 3.29 4.29 2.46		Blautia	3.95	3.29	2.40 0.31 1.21 1.65 4.29 2.46 1.45 3.65 37.33 42.5	2.46
<i>Escherichia</i> 1.36 3.40 1.45 3.65		Escherichia	1.36	3.40	1.45	3.65
Faecalibacterium 38.95 39.19 37.33 42.50		Faecalibacterium	38.95	39.19	37.33	42.50
Flavonifractor 0.72 2.79 1.56 1.56		Flavonifractor	0.72	2.79	1.56	1.56
Fusicatenibacter 9.18 3.56 8.30 4.12		Fusicatenibacter	9.18	3.56	8.30	4.12
Gemmiger 11.38 7.32 10.89 7.41		Gemmiger	11.38	7.32	10.89	7.41
Klebsiella 0.55 0.77 0.96 0.00		Klebsiella	0.55	0.77	0.96	0.00
Lachnospiraceae 9.42 7.48 9.13 7.64 unclassified		Lachnospiraceae unclassified	9.42	7.48	9.13	7.64
Roseburia 2.06 1.20 2.30 0.55		Roseburia	2.06	1.20	2.30	0.55
Ruminococcaceae 2.32 1.02 2.49 0.41 unclassified		Ruminococcaceae unclassified	2.32	1.02	2.49	0.41
Ruminococcus 4.51 3.05 4.43 2.89		Ruminococcus	4.51	3.05	4.43	2.89

higher contribution to L-citrulline biosynthesis in those with PTSD or depression. We also found that the Alistipes genera contributed more to ornithine-related pathways in those with PTSD and depression. Thus, by combining analyses of species abundance

with microbial metabolic pathways, we can better dissect the functional contributions of the microbiota.

The emerging role of polyamines in neuropsychiatric disorders opens a door to a better understanding of the complex

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Table 6.	Microbial genus contribution to	citrulline-related	metabolic pathways s	stratified by PTSD a	and depression diagnoses.
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Pathway	Species	PTSD Raw Score < 31	PTSD Raw Score≥31	Depression t-score < 60	Depression t-score ≥ 60
CITRULBIO-PWY: L-citrulline	Alistipes finegoldii	61.07	70.93	54.58	76.92
biosynthesis	Coprococcus catus	2.81	0.00	1.92	0.00
	Escherichia coli	17.97	15.87	15.87	16.77
	Flavonifractor plautii	11.61	11.18	19.29	6.31
	Klebsiella pneumoniae	6.53	2.02	8.34	0.00

pathophysiology of these disorders. The ability of microbiota to produce polyamines and known associations between polyamineproducing microbes and neuropsychiatric disorders highlight the importance of the microbiome-gut-brain axis in human health. Our novel finding of the gut microbiome contributing to alterations in GABR pathways in our studied outcomes is the first direct mechanistic link between the gut microbiome and APNS. This result may provide some indirect evidence of a biological link for APNS along the microbiome-gut-brain axis via microbially generated metabolites.

Strengths and Limitations

We are the first to delve into the predictiveness of the microbiome to the core components of APNS. Although the sample size of our study was limited, our employment of ML methods and metagenomic profiling enabled us to maximize the utility and richness of data from the samples we had. Furthermore, the AURORA (parent) study ensured a comprehensive, thorough, and standardized system of scoring APNS outcomes longitudinally. Our sub-study involved a single time point collection for every individual and the assumption that the microbiome remains stable over the 2-12 weeks' time-period post trauma. We acknowledge that this sampling strategy is not ideal and that we would have gained better resolution with having samples at each visit. However, the microbiome tends to stabilize within two weeks event post major perturbations including infection, diet, and antibiotics [65-67]. Therefore we feel that a month after trauma exposure the microbiome have reached a stable yet trauma-shaped state and that the within individual variability that could occur among the different time points would be due to dayto-day fluctuations to be a fair assumption. Additionally, we encountered logistical difficulties with recruitment and sample collection due to COVID-19 restrictions resulting in only 106 of the approach 2,097 participants participating in this sub-study. This challenge necessitated the working assumption that the microbiome is stable following trauma. Further, compared to the parent study, our cohort was generally older, male, and non-Hispanic white (Supplemental figure 1). Despite an established high comorbidity between trauma and substance use [68], we did not find any correlation between our outcomes and substance uses in our cohort, but this should be reevaluated in a larger study as usage may affect the gut microbiome. Future work will aim to expand on our preliminary findings with a larger cohort and more sampling time points, including sampling closer to time of trauma.

Although we approached these APNS diagnoses as discrete outcomes, it is well known that there is much overlap between them. Traditional APNS classification evolved from the realms of specific medical specialties and thus are not indexed to specific biological processes or basis [1]. This biologic overlap may be responsible for the overlap in some features identified by our modeling. A study with a larger sample size may be able to tease apart the overlap of these outcomes. Finally, while our study points to a possible role of the microbiome in mediating APNS outcomes, animal models with selective microbiota modulation will be necessary to determine directionality of this phenotype.

CONCLUSIONS

APNS can have devastating long-term consequences for patients who have already suffered trauma, but APNS may be preventable. Our nested study, using a subset of the AURORA cohort, demonstrated the importance of the microbiome in influencing APNS development. While more work is needed, we are the first to describe a possible biologic link between the gut microbiome and post-trauma outcomes through arginine metabolism and global arginine pathways, which have already been associated with PTSD and other neuropsychiatric disorders. This discovery opens avenues for investigating prevention and treatment strategies through both targeted therapies and microbiome-based interventions. Our findings provide some evidence of a biological link for APNS along the microbiome-gut-brain axis via microbially generated metabolites.

DATA AVAILABILITY

Microbiome sequencing data is deposited in the Short Read Archive (SRA) under BioProject accession no. PRJNA940405. All data and scripts produced in the present study are also available upon reasonable request to the authors.

REFERENCES

- McLean SA, Ressler K, Koenen KC, Neylan T, Germine L, Jovanovic T, et al. The AURORA Study: a longitudinal, multimodal library of brain biology and function after traumatic stress exposure. Mol Psychiatr. 2020;25:283–96.
- Roberts AL, Gilman SE, Breslau J, Breslau N, Koenen KC. Race/ethnic differences in exposure to traumatic events, development of post-traumatic stress disorder, and treatment-seeking for post-traumatic stress disorder in the United States. Psychol Med. 2011;41:71–83.
- 3. Kessler RC. Posttraumatic stress disorder: the burden to the individual and to society. J Clin Psychiatry. 2000;61:4–12.
- Kessler RC, Berglund P, Chiu WT, Demler O, Heeringa S, Hiripi E, et al. The US National Comorbidity Survey Replication (NCS-R): design and field procedures. Int J Method Psych. 2004;13:69–92.
- Santiago PN, Ursano RJ, Gray CL, Pynoos RS, Spiegel D, Lewis-Fernandez R, et al. A Systematic Review of PTSD Prevalence and Trajectories in DSM-5 Defined Trauma Exposed Populations: Intentional and Non-Intentional Traumatic Events. Plos One. 2013;8:e59236.
- Koenen KC, Ratanatharathorn A, Ng L, McLaughlin KA, Bromet EJ, Stein DJ, et al. Posttraumatic stress disorder in the World Mental Health Surveys. Psychol Med. 2017;47:2260–74.
- Beaudoin FL, An X, Basu A, Ji Y, Liu M, Kessler RC, et al. Use of serial smartphonebased assessments to characterize diverse neuropsychiatric symptom trajectories in a large trauma survivor cohort. Transl Psychiat. 2023;13:4.
- Warner BB. The contribution of the gut microbiome to neurodevelopment and neuropsychiatric disorders. Pediatr Res. 2019;85:216–24.
- Rogers GB, Keating DJ, Young RL, Wong M-L, Licinio J, Wesselingh S. From gut dysbiosis to altered brain function and mental illness: mechanisms and pathways. Mol Psychiatr. 2016;21:738–48.
- Socała K, Doboszewska U, Szopa A, Serefko A, Włodarczyk M, Zielińska A, et al. The role of microbiota–gut–brain axis in neuropsychiatric and neurological disorders. Pharm Res. 2021;172:105840.
- Malan-Muller S, Valles-Colomer M, Raes J, Lowry CA, Seedat S, Hemmings SMJ. The Gut Microbiome and Mental Health: Implications for Anxiety- and Trauma-Related Disorders. Omics J Integr Biol. 2018;22:90–107.
- Hoyles L, Snelling T, Umlai U-K, Nicholson JK, Carding SR, Glen RC, et al. Microbiome–host systems interactions: protective effects of propionate upon the blood–brain barrier. Microbiome. 2018;6:55.

- Silva YP, Bernardi A, Frozza RL. The Role of Short-Chain Fatty Acids From Gut Microbiota in Gut-Brain Communication. Front Endocrinol. 2020;11:25.
- 14. Xu K, Gao X, Xia G, Chen M, Zeng N, Wang S, et al. Rapid gut dysbiosis induced by stroke exacerbates brain infarction in turn. Gut. 2021;70:1486–94.
- Neigh GN, Ali FF. Co-morbidity of PTSD and immune system dysfunction: opportunities for treatment. Curr Opin Pharm. 2016;29:104–10.
- Hemmings SMJ, Malan-Müller S, Heuvel LL, van den, Demmitt BA, Stanislawski MA, Smith DG, et al. The Microbiome in Posttraumatic Stress Disorder and Trauma-Exposed Controls. Psychosom Med. 2017;79:936–46.
- Bajaj JS, Sikaroodi M, Fagan A, Heuman D, Gilles H, Gavis EA, et al. Posttraumatic stress disorder is associated with altered gut microbiota that modulates cognitive performance in veterans with cirrhosis. Am J Physiol-gastr L. 2019;317:G661–G669.
- Haran JP The intestinal microbiome's roles in the development of stress-related disorders. 2019 Analysis Proposal Submitted to AURORA Executive Committee, University North Carolina on 2019-10-16.
- Stein DJ, McLaughlin KA, Koenen KC, Atwoli L, Friedman MJ, Hill ED, et al. DSM-5 AND ICD-11 DEFINITIONS OF POSTTRAUMATIC STRESS DISORDER: INVESTIGAT-ING "NARROW" AND "BROAD" APPROACHES. Depress Anxiety. 2014;31:494–505.
- Weathers FW, Litz BT, Keane TM, Palmieri PA, Marx BP, Schnurr PP The PTSD Checklist for DSM-5 (PCL-5). 2013. https://www.ptsd.va.gov/professional/ assessment/adult-sr/ptsd-checklist.asp (accessed 16 Sep2022).
- Blevins CA, Weathers FW, Davis MT, Witte TK, Domino JL. The Posttraumatic Stress Disorder Checklist for DSM-5 (PCL-5): Development and Initial Psychometric Evaluation. J Trauma Stress. 2015;28:489–98.
- Lang AJ, Stein MB. An abbreviated PTSD checklist for use as a screening instrument in primary care. Behav Res Ther. 2005;43:585–94.
- Bovin MJ, Marx BP, Weathers FW, Gallagher MW, Rodriguez P, Schnurr PP, et al. Psychometric Properties of the PTSD Checklist for Diagnostic and Statistical Manual of Mental Disorders–Fifth Edition (PCL-5) in Veterans. Psychol Assess. 2016;28:1379–91.
- Amtmann D, Kim J, Chung H, Bamer AM, Askew RL, Wu S, et al. Comparing CESD-10, PHQ-9, and PROMIS Depression Instruments in Individuals With Multiple Sclerosis. Rehabil Psychol. 2014;59:220–9.
- Auvergne L, Bortsov AV, Ulirsch JC, Peak DA, Jones JS, Swor RA, et al. Association of Epidemiologic Factors and Genetic Variants Influencing Hypothalamic-Pituitary-Adrenocortical Axis Function With Postconcussive Symptoms After Minor Motor Vehicle Collision. Psychosom Med. 2016;78:68–78.
- Beghini F, McIver LJ, Blanco-Míguez 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.
- Franzosa EA, McIver LJ, Rahnavard G, Thompson LR, Schirmer M, Weingart G, et al. Species-level functional profiling of metagenomes and metatranscriptomes. Nat Methods. 2018;15:962–8.
- Bates D, Mächler M, Bolker B, Walker S Fitting Linear Mixed-Effects Models Using Ime4. J Stat Softw 2015; 67. https://doi.org/10.18637/jss.v067.i01.
- Nievergelt CM, Maihofer AX, Klengel T, Atkinson EG, Chen C-Y, Choi KW, et al. International meta-analysis of PTSD genome-wide association studies identifies sex- and ancestry-specific genetic risk loci. Nat Commun. 2019;10:4558.
- Morgan XC, Kabakchiev B, Waldron L, Tyler AD, Tickle TL, Milgrom R, et al. Associations between host gene expression, the mucosal microbiome, and clinical outcome in the pelvic pouch of patients with inflammatory bowel disease. Genome Biol. 2015;16:67.
- Lahti L, Salojärvi J, Salonen A, Scheffer M, Vos WMde. Tipping elements in the human intestinal ecosystem. Nat Commun. 2014;5:4344.
- Thompson J, Johansen R, Dunbar J, Munsky B. Machine learning to predict microbial community functions: An analysis of dissolved organic carbon from litter decomposition. Plos One. 2019;14:e0215502.
- Topçuoğlu BD, Lesniak NA, Ruffin MT, Wiens J, Schloss PD. A Framework for Effective Application of Machine Learning to Microbiome-Based Classification Problems. Mbio. 2020;11:e00434–20.
- Haran JP, Bhattarai SK, Foley SE, Dutta P, Ward DV, Bucci V et al. Alzheimer's Disease Microbiome Is Associated with Dysregulation of the Anti-Inflammatory P-Glycoprotein Pathway. *Mbio* 2019; 10. https://doi.org/10.1128/mbio.00632-19.
- Haran JP, Bradley E, Zeamer AL, Cincotta L, Salive M-C, Dutta P et al. Inflammation-type dysbiosis of the oral microbiome associates with the duration of COVID-19 symptoms and long-COVID. *Jci Insight* 2021; 6. https://doi.org/ 10.1172/jci.insight.152346.
- Wipperman MF, Bhattarai SK, Vorkas CK, Maringati VS, Taur Y, Mathurin L, et al. Gastrointestinal microbiota composition predicts peripheral inflammatory state during treatment of human tuberculosis. Nat Commun. 2021;12:1141.
- Olendzki B, Bucci V, Cawley C, Maserati R, McManus M, Olednzki E, et al. Dietary manipulation of the gut microbiome in inflammatory bowel disease patients: Pilot study. Gut Microbes. 2022;14:2046244.

- Martínez I, Muller CE, Walter J. Long-Term Temporal Analysis of the Human Fecal Microbiota Revealed a Stable Core of Dominant Bacterial Species. Plos One. 2013;8:e69621.
- 39. Caporaso JG, Lauber CL, Costello EK, Berg-Lyons D, Gonzalez A, Stombaugh J, et al. Moving pictures of the human microbiome. Genome Biol. 2011;12:R50.
- David LA, Materna AC, Friedman J, Campos-Baptista MI, Blackburn MC, Perrotta A, et al. Host lifestyle affects human microbiota on daily timescales. Genome Biol. 2014;15:R89.
- Bucci V, Bradde S, Biroli G, Xavier JB. Social Interaction, Noise and Antibiotic-Mediated Switches in the Intestinal Microbiota. Plos Comput Biol. 2012;8:e1002497.
- Bradley ES, Zeamer AL, Bucci V, Cincotta L, Salive M-C, Dutta P, et al. Oropharyngeal microbiome profiled at admission is predictive of the need for respiratory support among COVID-19 patients. Front Microbiol. 2022;13:1009440.
- Haran JP, Zeamer A, Ward DV, Dutta P, Bucci V, McCormick BA The Nursing Home Older Adult Gut Microbiome Composition Shows Time-dependent Dysbiosis and is Influenced by Medication Exposures, Age, Environment, and Frailty. *Journals Gerontology Ser* 2021. https://doi.org/10.1093/gerona/glab167.
- Bradley ES, Schell B, Ward DV, Bucci V, Zeamer A, Haran JP The Urinary Microbiome of Older Adults Residing in a Nursing Home Varies With Duration of Residence and Shows Increases in Potential Pathogens. *Journals Gerontology Ser* 2021. https://doi.org/10.1093/gerona/glab345.
- 45. Prince C, Bruhns ME. Evaluation and Treatment of Mild Traumatic Brain Injury: The Role of Neuropsychology. Brain Sci. 2017;7:105.
- 46. Bersani FS, Wolkowitz OM, Lindqvist D, Yehuda R, Flory J, Bierer LM, et al. Global arginine bioavailability, a marker of nitric oxide synthetic capacity, is decreased in PTSD and correlated with symptom severity and markers of inflammation. Brain Behav Immun. 2016;52:153–60.
- Segata N, Izard J, Waldron L, Gevers D, Miropolsky L, Garrett WS, et al. Metagenomic biomarker discovery and explanation. Genome Biol. 2011;12:R60.
- Howard BM, Kornblith LZ, Christie SA, Conroy AS, Nelson MF, Campion EM, et al. Characterizing the gut microbiome in trauma: significant changes in microbial diversity occur early after severe injury. Trauma Surg Acute Care Open. 2017;2:e000108.
- Gao F, Guo R, Ma Q, Li Y, Wang W, Fan Y, et al. Stressful events induce longterm gut microbiota dysbiosis and associated post-traumatic stress symptoms in healthcare workers fighting against COVID-19. J Affect Disord. 2022;303:187–95.
- Rong H, Xie X, Zhao J, Lai W, Wang M, Xu D, et al. Similarly in depression, nuances of gut microbiota: Evidences from a shotgun metagenomics sequencing study on major depressive disorder versus bipolar disorder with current major depressive episode patients. J Psychiatr Res. 2019;113:90–99.
- Lai W, Deng W, Xu S, Zhao J, Xu D, Liu Y, et al. Shotgun metagenomics reveals both taxonomic and tryptophan pathway differences of gut microbiota in major depressive disorder patients. Psychol Med. 2021;51:90–101.
- Tang WHW, Wang Z, Cho L, Brennan DM, Hazen SL. Diminished Global Arginine Bioavailability and Increased Arginine Catabolism as Metabolic Profile of Increased Cardiovascular Risk. J Am Coll Cardiol. 2009;53:2061–7.
- Liu P, Jing Y, Collie ND, Dean B, Bilkey DK, Zhang H. Altered brain arginine metabolism in schizophrenia. Transl Psychiat. 2016;6:e871–e871.
- Ozden A, Angelos H, Feyza A, Elizabeth W, John P. Altered plasma levels of arginine metabolites in depression. J Psychiatr Res. 2020;120:21–28.
- 55. Viñuela-Berni V, Gómez-González B, Quintanar-Stephano A. Blockade of Arginine Vasopressin receptors prevents blood-brain barrier breakdown in Experimental Autoimmune Encephalomyelitis. Sci Rep.-uk. 2020;10:467.
- Fiori LM, Turecki G. Implication of the polyamine system in mental disorders. J Psychiatry Neurosci Jpn. 2008;33:102–10.
- Bercik P, Park AJ, Sinclair D, Khoshdel A, Lu J, Huang X, et al. The anxiolytic effect of Bifidobacterium longum NCC3001 involves vagal pathways for gut–brain communication. Neurogastroenterol Motil. 2011;23:1132–9.
- Kelly JR, Borre Y, Brien CO, Patterson E, Aidy SE, Deane J, et al. Transferring the blues: Depression-associated gut microbiota induces neurobehavioural changes in the rat. J Psychiatr Res. 2016;82:109–18.
- Liu Y, Zhang L, Wang X, Wang Z, Zhang J, Jiang R, et al. Similar Fecal Microbiota Signatures in Patients With Diarrhea-Predominant Irritable Bowel Syndrome and Patients With Depression. Clin Gastroenterol H. 2016;14:1602–1611.e5.
- Savignac HM, Tramullas M, Kiely B, Dinan TG, Cryan JF. Bifidobacteria modulate cognitive processes in an anxious mouse strain. Behav Brain Res. 2015;287:59–72.
- Nikolova VL, Hall MRB, Hall LJ, Cleare AJ, Stone JM, Young AH. Perturbations in Gut Microbiota Composition in Psychiatric Disorders. Jama Psychiat. 2021;78:1–12.
- 62. Liu RT, Rowan-Nash AD, Sheehan AE, Walsh RFL, Sanzari CM, Korry BJ, et al. Reductions in anti-inflammatory gut bacteria are associated with depression in a sample of young adults. Brain Behav Immun. 2020;88:308–24.

- 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.
- 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 Psychiatr. 2016;21:786–96.
- Neto CC, Mortzfeld BM, Turbitt JR, Bhattarai SK, Yeliseyev V, DiBenedetto N, et al. Proanthocyanidin-enriched cranberry extract induces resilient bacterial community dynamics in a gnotobiotic mouse model. Micro Cell. 2021;8:131.
- 66. Letourneau J, Holmes ZC, Dallow EP, Durand HK, Jiang S, Carrion VM, et al. Ecological memory of prior nutrient exposure in the human gut microbiome. ISME J. 2022;16:2479–90.
- 67. Leo S, Lazarevic V, Dach E, von, Kaiser L, Prendki V, Schrenzel J, et al. Effects of antibiotic duration on the intestinal microbiota and resistome: The PIRATE RESISTANCE project, a cohort study nested within a randomized trial. EBioMedicine. 2021;71:103566.
- Ullman SE, Relyea M, Peter-Hagene L, Vasquez AL. Trauma histories, substance use coping, PTSD, and problem substance use among sexual assault victims. Addict Behav. 2013;38:2219–23.
- Duranti S, Ruiz L, Lugli GA, Tames H, Milani C, Mancabelli L, et al. Bifidobacterium adolescentis as a key member of the human gut microbiota in the production of GABA. Sci Rep.-uk. 2020;10:14112.
- Tan TG, Sefik E, Geva-Zatorsky N, Kua L, Naskar D, Teng F, et al. Identifying species of symbiont bacteria from the human gut that, alone, can induce intestinal Th17 cells in mice. Proc Natl Acad Sci. 2016;113:E8141–E8150.
- Joossens M, Huys G, Cnockaert M, Preter VD, Verbeke K, Rutgeerts P, et al. Dysbiosis of the faecal microbiota in patients with Crohn's disease and their unaffected relatives. Gut. 2011;60:631.
- Wang L, Christophersen CT, Sorich MJ, Gerber JP, Angley MT, Conlon MA. Low Relative Abundances of the Mucolytic Bacterium Akkermansia muciniphila and Bifidobacterium spp. in Feces of Children with Autism. Appl Environ Micro. 2011;77:6718–21.
- Barrett E, Ross RP, O'Toole PW, Fitzgerald GF, Stanton C. γ-Aminobutyric acid production by culturable bacteria from the human intestine. J Appl Microbiol. 2012;113:411–7.
- Sela DA, Mills DA. Nursing our microbiota: molecular linkages between bifidobacteria and milk oligosaccharides. Trends Microbiol. 2010;18:298–307.
- López P, González-Rodríguez I, Gueimonde M, Margolles A, Suárez A. Immune Response to Bifidobacterium bifidum Strains Support Treg/Th17 Plasticity. Plos One. 2011;6:e24776.
- Bellali S, Lo CI, Naud S, Fonkou MDM, Armstrong N, Raoult D, et al. Parabacteroides massiliensis sp. nov., a new bacterium isolated from a fresh human stool specimen. N. Microbes N. Infect. 2019;32:100602.
- Bartlett A, Padfield D, Lear L, Bendall R, Vos M A comprehensive list of bacterial pathogens infecting humans. *Microbiology*+ 2022; 168. https://doi.org/10.1099/ mic.0.001269.
- Qiao S, Liu C, Sun L, Wang T, Dai H, Wang K, et al. Gut Parabacteroides merdae protects against cardiovascular damage by enhancing branched-chain amino acid catabolism. Nat Metab. 2022;4:1271–86.
- Ogita T, Yamamoto Y, Mikami A, Shigemori S, Sato T, Shimosato T. Oral Administration of Flavonifractor plautii Strongly Suppresses Th2 Immune Responses in Mice. Front Immunol. 2020;11:379.
- Valles-Colomer M, Falony G, Darzi Y, Tigchelaar EF, Wang J, Tito RY, et al. The neuroactive potential of the human gut microbiota in quality of life and depression. Nat Microbiol. 2019;4:623–32.
- Vázquez L, Flórez AB, Rodríguez J, Mayo B. Heterologous expression of equal biosynthesis genes from Adlercreutzia equalifaciens. Fems Microbiol Lett. 2021;368:fnab082.
- Hannele J-S Wadsworth-KTL anaerobic bacteriology manual. 6th ed. Star Pub.: Belmont, CA, 2002.
- Parker BJ, Wearsch PA, Veloo ACM, Rodriguez-Palacios A. The Genus Alistipes: Gut Bacteria With Emerging Implications to Inflammation, Cancer, and Mental Health. Front Immunol. 2020;11:906.
- Dai Z, Coker OO, Nakatsu G, Wu WKK, Zhao L, Chen Z, et al. Multi-cohort analysis of colorectal cancer metagenome identified altered bacteria across populations and universal bacterial markers. Microbiome. 2018;6:70.
- Chung WSF, Meijerink M, Zeuner B, Holck J, Louis P, Meyer AS et al. Prebiotic potential of pectin and pectic oligosaccharides to promote anti-inflammatory commensal bacteria in the human colon. *Fems Microbiol Ecol* 2017; **93**. https:// doi.org/10.1093/femsec/fix127.
- Quan J, Wu Z, Ye Y, Peng L, Wu J, Ruan D, et al. Metagenomic Characterization of Intestinal Regions in Pigs With Contrasting Feed Efficiency. Front Microbiol. 2020;11:32.
- 87. Schröder C, Matthies A, Engst W, Blaut M, Braune A. Identification and Expression of Genes Involved in the Conversion of Daidzein and Genistein by the Equol-

Forming Bacterium Slackia isoflavoniconvertens. Appl Environ Micro. 2013;79:3494–502.

- Loubinoux J, Bronowicki J, Pereira IAC, Mougenel J, Faou AE. Sulfate-reducing bacteria in human feces and their association with inflammatory bowel diseases. Fems Microbiol Ecol. 2002;40:107–12.
- Hosomi K, Saito M, Park J, Murakami H, Shibata N, Ando M, et al. Oral administration of Blautia wexlerae ameliorates obesity and type 2 diabetes via metabolic remodeling of the gut microbiota. Nat Commun. 2022;13:4477.
- Cox LM, Maghzi AH, Liu S, Tankou SK, Dhang FH, Willocq V, et al. Gut Microbiome in Progressive Multiple Sclerosis. Ann Neurol. 2021;89:1195–211.
- Bui TPN, Shetty SA, Lagkouvardos I, Ritari J, Chamlagain B, Douillard FP, et al. Comparative genomics and physiology of the butyrate-producing bacterium Intestinimonas butyriciproducens. Env Microbiol Rep. 2016;8:1024–37.
- Lai H-C, Lin T-L, Chen T-W, Kuo Y-L, Chang C-J, Wu T-R, et al. Gut microbiota modulates COPD pathogenesis: role of anti-inflammatory Parabacteroides goldsteinii lipopolysaccharide. Gut. 2022;71:309–21.
- Wu I-W, Lin C-Y, Chang L-C, Lee C-C, Chiu C-Y, Hsu H-J, et al. Gut Microbiota as Diagnostic Tools for Mirroring Disease Progression and Circulating Nephrotoxin Levels in Chronic Kidney Disease: Discovery and Validation Study. Int J Biol Sci. 2020;16:420–34.
- Louis S, Tappu R-M, Damms-Machado A, Huson DH, Bischoff SC. Characterization of the Gut Microbial Community of Obese Patients Following a Weight-Loss Intervention Using Whole Metagenome Shotgun Sequencing. Plos One. 2016;11:e0149564.
- Xie Y-H, Gao Q-Y, Cai G-X, Sun X-M, Zou T-H, Chen H-M, et al. Fecal Clostridium symbiosum for Noninvasive Detection of Early and Advanced Colorectal Cancer: Test and Validation Studies. Ebiomedicine. 2017;25:32–40.
- Ananthakrishnan AN, Luo C, Yajnik V, Khalili H, Garber JJ, Stevens BW, et al. Gut Microbiome Function Predicts Response to Anti-integrin Biologic Therapy in Inflammatory Bowel Diseases. Cell Host Microbe. 2017;21:603–610.e3.

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AUTHOR CONTRIBUTIONS

All authors certify that they meet authorship criteria. A.L.Z. wrote the first draft and incorporated feedback from coauthors, who had also contributed to conduct of the study. The analysis was planned by A.L.Z, V.B. and J.P.H., with input from other coauthors. Sequencing data was generated and processed by A.L.Z. Clinical data were prepared by MC.S. and A.X. and analyzed by A.L.Z., who had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

COMPETING INTERESTS

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