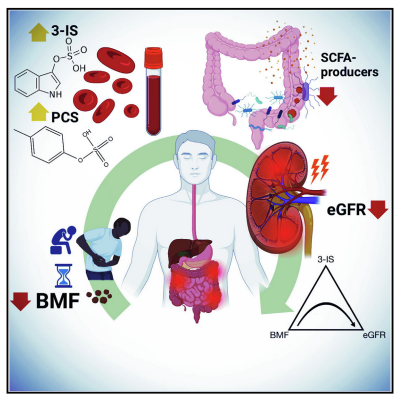
Article

Aberrant bowel movement frequencies coincide with increased microbe-derived blood metabolites associated with reduced organ function

Graphical abstract



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In brief

Johnson-Martínez et al. determine that bowel movement frequency (BMF) is significantly associated with variation in molecular phenotypes in a generally healthy human cohort. Constipation and diarrhea are negatively associated with liver and kidney function, respectively, with a microbial metabolite mediating the effect of BMF on kidney function.

Highlights

- BMF variation impacts the gut microbiota and blood metabolome in a healthy cohort
- Aberrant BMF associates with microbially derived toxins in blood and with inflammation
- Indoxyl sulfate linked to constipation and negatively associates with kidney function
- Diet, lifestyle, and cognitive factors are associated with BMF variation

Johnson-Martínez et al., 2024, Cell Reports Medicine 5, 101646 July 16, 2024 © 2024 The Author(s). Published by Elsevier Inc. https://doi.org/10.1016/j.xcrm.2024.101646



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Aberrant bowel movement frequencies coincide with increased microbe-derived blood metabolites associated with reduced organ function

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https://doi.org/10.1016/j.xcrm.2024.101646

SUMMARY

Bowel movement frequency (BMF) directly impacts the gut microbiota and is linked to diseases like chronic kidney disease or dementia. In particular, prior work has shown that constipation is associated with an ecosystem-wide switch from fiber fermentation and short-chain fatty acid production to more detrimental protein fermentation and toxin production. Here, we analyze multi-omic data from generally healthy adults to see how BMF affects their molecular phenotypes, in a pre-disease context. Results show differential abundances of gut microbial genera, blood metabolites, and variation in lifestyle factors across BMF categories. These differences relate to inflammation, heart health, liver function, and kidney function. Causal mediation analysis indicates that the association between lower BMF and reduced kidney function is partially mediated by the microbially derived toxin 3-indoxyl sulfate (3-IS). This result, in a generally healthy context, suggests that the accumulation of microbiota-derived toxins associated with abnormal BMF precede organ damage and may be drivers of chronic, aging-related diseases.

INTRODUCTION

The gut microbiome influences human health in a number of ways, from mediating early-life immune system development, ^{1,2} to determining personalized responses to nutritional interventions^{3,4} and influencing the central nervous system. ^{5,6} Bowel movement frequency (BMF), defined as the frequency of defecation, is a major determinant of the composition of the human gut microbiota and can be affected by diet, hydration, physical activity, host mucus production, microbe- and host-derived small molecules (e.g., short-chain fatty acids [SCFAs], bile acids, or neurotransmitters), and peristaltic smooth muscle contractions in the gastrointestinal tract.^{7–9} Aberrant BMFs have been implicated in the development of various chronic diseases.^{10–12}

Abnormally high BMF (e.g., diarrhea, defined as more than three watery stools per day) has been associated with lower gut microbiome alpha-diversity, inflammation, increased susceptibility to enteric pathogens, and poorer overall health.¹³⁻¹⁶ Abnormally low BMF (e.g., constipation, defined as fewer than three hard, dry stools per week) has been associated with higher gut microbiome alpha-diversity, an enrichment in microbially derived urinary metabolites known to be hepatotoxic or nephrotoxic, and an increased risk for several chronic medical conditions, including neurodegenerative disorders and chronic kidney disease (CKD).^{10,17–19} Indeed, constipation is a known risk factor for CKD severity and end-stage renal disease progression.^{20,21} In one study, up to 71% of dialysis patients suffered from constipation,²² while the prevalence of constipation in the general population was 14.5% in adults under 60 years old and 33.5% in those over 60.²³ A nationwide study of veterans found an incrementally higher risk for renal disease progression in those who reported increasingly severe constipation.²⁴ However, while it is clear that morbidity and mortality risk worsen with constipation in those with active CKD, potential connections between BMF



and the development of early-stage kidney disease are not yet established.

Both constipation and CKD associate with declines in gut microbiota-mediated SCFA production and a rise in the production of amino acid putrefaction byproducts, including several toxic microbe-host co-metabolites, such as 3-indoxyl sulfate (3-IS), *p*-cresol sulfate (PCS), *p*-cresol glucuronide (PCG), and phenylacetylglutamine (PAG), which all have been implicated in CKD progression.^{25–27} This is consistent with an established microbiota-wide transition from saccharolytic to proteolytic fermentation in constipated individuals due to the exhaustion of dietary fiber in stool.^{10,28} Thus, while the potential relationship between BMF and organ function in healthy populations is not fully understood, the gut metabolic phenotype associated with lower BMF suggests a connection.

In this study, we focus on categories of self-reported BMF in a large population of generally healthy individuals with a wide range of molecular phenotypic data in order to quantify the phenotypic impact of BMF on blood plasma metabolites, blood proteins, clinical chemistries, and gut microbiome composition in a pre-disease context. By exploring the molecular phenotypic consequences of BMF variation in a generally healthy cohort, along with BMF-associated demographic, dietary, lifestyle, and psychological factors, we aimed to identify early-stage biomarkers and potential therapeutic targets for the monitoring and prevention of certain chronic, non-communicable diseases, like CKD.

RESULTS

A cohort of generally healthy individuals

3,955 Arivale Scientific Wellness program participants with BMF data were initially considered in this analysis. Arivale, Inc. (USA) was a consumer scientific wellness company that operated from 2015 until 2019. Briefly, participants consented to having their health, diet, and lifestyle surveyed through an extensive questionnaire, along with blood and stool sampling for multi-omic and blood plasma chemistries data generation (Figure 1). Any respondents that indicated "true" or affirmatively to any of the following questionnaire features were excluded from the analysis (i.e., they were not considered "generally healthy"): taking blood pressure, cholesterol, or laxative medication or having self or family history of bladder or kidney disease (i.e., kidney cancer, bladder infections, polycystic kidney disease or PKD, kidney stones, kidney failure, or kidney disease), inflammatory bowel disease (IBD; both Crohn's disease and ulcerative colitis), irritable bowel syndrome, celiac disease, diverticulosis, gastroesophageal reflux disease, or peptic ulcers (i.e., these individuals were not considered "generally healthy"; see supplemental information and Table S1). There were 1,425 participants who met these exclusion criteria and had necessary covariate data. Across all Arivale participants that had available demographic and survey information, 82.8% of those individuals identified as "White" (N = 2,562), 8.5% identified as "Asian" (N = 262), 3.2% identified as "Black or African American" (N = 98), 0.2% identified as "American Indian or Alaska Native" (N = 9), 0.65% identified as "Native Hawaiian or other Pacific Islander" (N = 20), and 4.7% identified as "other" (N = 144). 93.6% of these individuals iden-

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tified as "non-Hispanic" (N = 2,897) and 6.4% identified as "Hispanic" (N = 198, 55.6% of which self-identify as "White"). Respondents were in the United States, predominantly from the Pacific West, and their ages ranged from 19 to 89 years old. 65.1% were female with a mean ± SD body mass index of 27.15 ± 5.89 (Table S1). 1,062 of these individuals had gut microbiome data, 486 had blood metabolomics data, 823 had proteomics data, 1,425 had clinical chemistries data, and 1,420 had survey data (derived from questionnaires). Self-reported BMF values (responses to typical number of bowel movements per week) were grouped into four categories (Figure 1), which we labeled as "constipation" (≤2 bowel movements per week), "low-normal" (3-6 bowel movements per week), "high-normal" (1-3 bowel movements per day), and "diarrhea" (4 or more bowel movements per day). We first looked at potential associations between BMF and relevant covariates: gender, age, BMI, estimated glomerular filtration rate (eGFR), low-density lipoprotein blood plasma levels (LDL), C-reactive protein blood plasma levels (CRP), hemoglobin A1c blood plasma levels (A1C), and the first three principal components of genetic ancestry (PC1, PC2, and PC3) (N = 1,425; Figure 2). When BMF was coded as an ordinal dependent variable and regressed using ordered proportional odds logistic regression²⁹ (POLR), only BMI (POLR, false discovery rate (FDR)-corrected p = 1.82E-3), age (POLR, FDR-corrected, p = 2.07E-3), gender (POLR, FDR-corrected p = 3.68E - 16), and the first three principal components of genetic ancestry (PC1, PC2, and PC3; POLR, FDR-corrected p < 0.0001) showed significant, independent associations with BMF, with females, older individuals, and individuals with lower BMIs tending to report lower BMFs (Figure 2). All covariates listed earlier were included in downstream regressions, regardless of whether or not they showed an independent association with BMF. The high-normal BMF group was chosen as the reference for all downstream regressions where BMF was encoded as a categorical variable. eGFR was also regressed against BMF and the other covariates to determine which were significantly associated with eGFR, and the covariates with significant p values included gender, age, BMI, LDL, A1C, PC1, PC2, and PC3 (generalized linear modeling [GLM], p < 0.05).

Gut microbiome structure and composition across BMF categories

We looked at a subcohort of individuals that met our health exclusion criteria with 16S amplicon sequencing data from stool (N = 1,062). Amplicon sequence variant richness (GLM, p = 2.85E-3, linear $\beta_{BMF} = -65.9E-3$) and Shannon diversity (GLM, p = 1.07E-3, linear $\beta_{BMF} = -3.25E-1$) were negatively associated with BMF, independent of the covariates listed earlier, and with BMF encoded as an ordinal variable with a linear coefficient (Figures 3 and S1). Pielou's evenness, on the other hand, was positively associated with BMF (GLM, p = 8.5E-3, linear $\beta_{BMF} = 2.6E-3$), independent of covariates (Figure 3).

Differential abundance analysis of commensal gut bacterial genera across BMF categories was conducted using beta-binomial regression³⁰ (count regression for correlated observations with the beta-binomial [CORNCOB]) with BMF encoded as a categorical variable with the high-normal group as the reference category. Of the 135 genera that passed our prevalence filter

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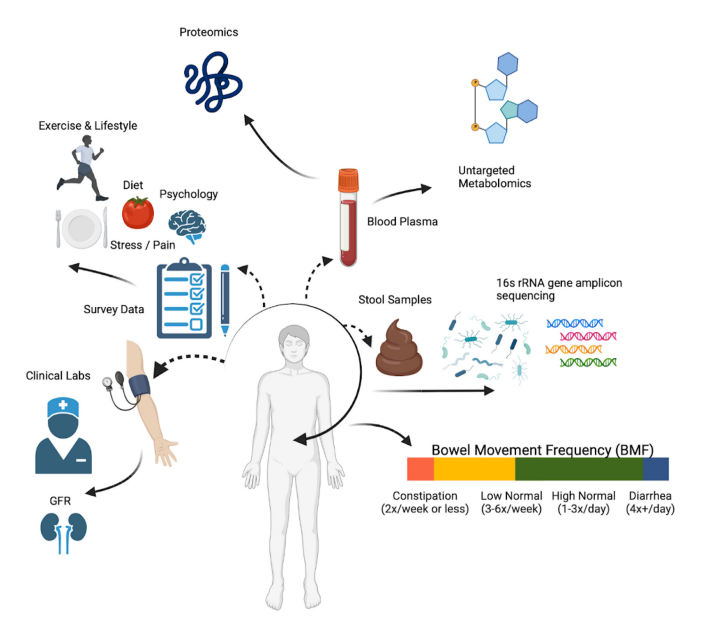


Figure 1. Data collection strategy

Arivale participants were sampled for blood plasma and stool, in addition to filling out extensive diet, health, and lifestyle questionnaires. Clinical chemistries, untargeted metabolomics, and proteomics data were generated from blood plasma samples. Gut microbiome 16S rRNA amplicon sequencing data were generated from stool samples collected using at-home kits. BMF data were extracted from the questionnaire data as self-reported frequencies per week or day.

(i.e., detection across \geq 30% of individuals), 59 were significantly associated with BMF (49 of which had genus-level taxonomic annotations; see Table S1 for detailed list of β -coefficients and p values), independent of covariates and following an FDR correction for multiple tests on the likelihood ratio test p values (FDR-corrected p < 0.05). We Z score normalized the centered log-ratio (CLR) abundances of the 49 annotated genera across all samples and then plotted the average Z score within each BMF bin for each taxon as a heatmap (Figure 4). We also provide supplemental boxplots, showing CLR abundances across BMF categories, of the top 10 most abundant taxa and 10 taxa with the smallest p values from the 49 mentioned earlier (Figures S1 and S2). In order of descending abundance, the following taxa were significantly enriched in constipated individuals, compared to the high-normal BMF category (Wald test, FDR-corrected $\beta_{BMF} p < 0.05$): Ruminiclostridium_9, Ruminococcaceae_UCG-005, Ruminococcaceae_NK41214_group, Family_XIII_AD3011_group, Romboutsia, Ruminocacceae_UCG-004, UBA1819, Negativibacillus, DTU089, GCA-900066225, Candidatus_Soleaferrea, Anaerotruncus, Defluviitaleaeceae_UCG-011, Eisenbergiella, Pygmalobacter, Peptococcus, Hydrogenoanaerobacterium, Anaerofustis, and DNF00809. Lachnospiraceae_ND3007_group and Lachnospiraceae_UCG-004 were significantly depleted in constipated individuals. Several more were associated with



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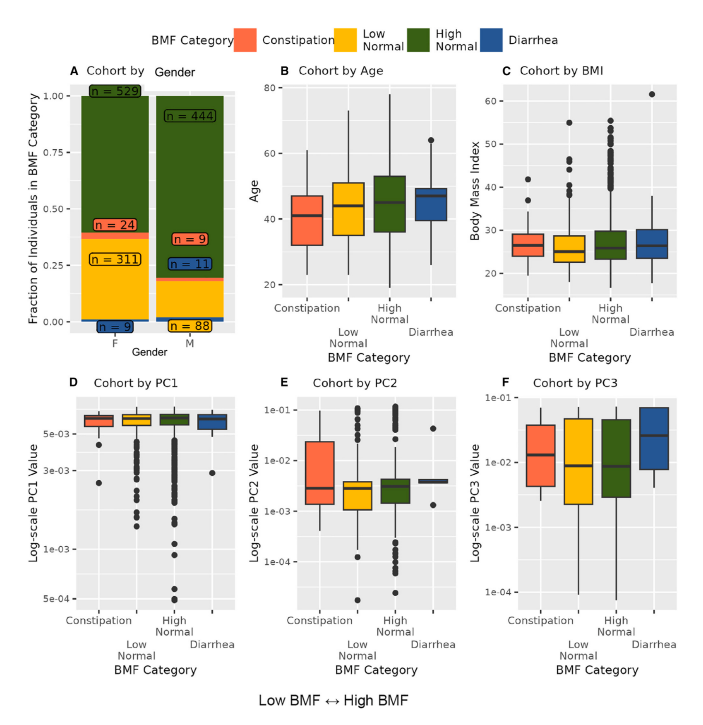
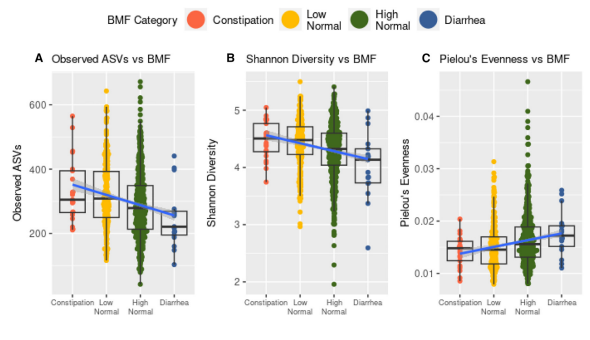


Figure 2. Exploring covariate associations with bowel movement frequency (BMF)

Plotting covariates that showed a significant association with BMF: gender, age, BMI, and the first three principal components of genetic ancestry (PC1-PC3) (A–F). POLR was used to regress BMF against the covariates (gender, age, BMI, eGFR, LDL, CRP, A1C, plus the first three principal components of genetic ancestry in the cohort, PC1, PC2, and PC3). The result was that gender (p = 3.68E-16), BMI (p = 1.82E-3), age (p = 2.075E-3), and PC3 1–3 (p < 0.00001, respectively) were significantly associated with BMF. In panels (B)–(E), center lines on boxplots represent the median, the top and bottom edges of the box represent the interquartile range, the black dots show points more than 1.5 times the interquartile range from the ends of the box, and the whiskers show the smallest or largest value within 1.5 times the 25th or 75th quartile, respectively.

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Linear Ordinal Coefficient P-Values:

P value = 0.00285

P value = 0.00107

P value = 0.0085

Figure 3. Associations between gut microbiome alpha-diversity and BMF

(A) Richness of amplicon sequence variants (ASVs) across BMF categories (ordinal BMF variable, linear regression, p = 2.85E-3).

(B) Shannon diversity across BMF categories (ordinal BMF variable, linear regression, p = 1.07E-3).

(C) Pielou's evenness across BMF categories (ordinal BMF variable, linear regression, p = 8.5E-2).

Center lines in the boxplots show the median, the ends of the boxes show the interquartile range, and whiskers show the span of points within 1.5 times the interquartile range from the ends of the box.

enrichment or depletion in the low-normal BMF category, compared to the reference category (Figure 4; see supplemental information). There was no significant difference between the high-normal and diarrhea categories for any of the genera, which could be due to low sample size in the diarrhea category (i.e., we were likely underpowered to detect those associations).

Variation in blood metabolites across BMF categories

Blood metabolite vs. BMF regression analyses were run using a GLM framework in LIMMA (linear models for microarray and RNA-seq data), with BMF as a categorical independent variable, along with the same set of covariates mentioned earlier. Of the metabolites that passed our abundance and prevalence filters (N = 956, see method details), 9 separate metabolites were significantly associated with BMF (all 9 showed differential abundance between lownormal and high-normal categories, which is the comparison we were most powered for), independent of covariates and following an FDR correction for multiple tests (GLM, FDR-corrected p < 0.05; Figures 5 and S3). The annotated metabolites tended to show a decreasing trend with increasing BMF, while the unannotated metabolites and 3-IS showed more varied relationships (e.g., monotonic and non-monotonic) with BMF (Figures 5 and S3). PCS, PAG, PCG, and 3-IS were significantly enriched in the low-normal BMF category, compared to the reference category (Figures 5 and S3). 75 different metabolites were significantly associated with eGFR, independent of covariates and following the same FDR correction for multiple tests (linear regression, FDR-corrected p < 0.05; Figures 5 and S3). Only one of these eGFR-associated metabolites overlapped with any of the BMF-associated metabolites: 3-IS.

Blood plasma chemistries across BMF categories

Of the 55 blood plasma chemistries filtered for prevalence (see method details), 21 were significantly associated with diarrhea (e.g., omega-6 fatty acid, homocysteine, total protein, and bilirubin) and one (omega-6/omega-3 ratio in the blood) was associated with the low-normal BMF category, relative to the reference category, after adjusting for all covariates and for multiple testing (Figures 6 and S4; N = 1,425, GLM, FDR-corrected p < 0.05).

Blood proteomics across BMF categories

None of the 274 blood proteins that passed our prevalence filter (see method details) showed significant associations with BMF after adjusting for all covariates and for multiple testing (N = 823, GLM, FDR-corrected p < 0.05).



				Normalized	ł
	**			Bacteroides Z-SCOre	
	**			Blautia	0.6
	*			Lachnoclostridium	0.4
	**			Ruminococcus_1	0.2
	***			Butyricicoccus	0
				Ruminococcaceae_UCG-013	-0.2
	**			Ruminococcaceae_UCG-002	-0.4
	***			Ruminiclostridium_5	-0.6
*	*			Ruminiclostridium_9	
	***			Christensenellaceae_R-7_group	
***	***			Ruminococcaceae_UCG-005	
				Bifidobacterium	
*				Lachnospiraceae_ND3007_group	
*	***			Lachnospiraceae_UCG-004	
*	*			Ruminococcaceae_NK4A214_group	
***	***			Family_XIII_AD3011_group	
	**			Ruminococcaceae_UCG-003	
	***			Marvinbryantia	
*	**			Romboutsia	
*	***			Ruminococcaceae_UCG-004	
				Akkermansia	
***	**			UBA1819	
***	***			Negativibacillus Adiercreutzia	
	***			Adlercreutzia	
	**			Ruminococcaceae_UCG-010	
	*			Family_XIII_UCG-001	
**	***			DTU089	
	*			Turicibacter	
*	***			GCA-900066225	
*	***			Candidatus_Soleaferrea	
**	**			Anaerotruncus	
*	***			Defluviitaleaceae_UCG-011	
	**			Caproiciproducens	
	*			Shuttleworthia	
	*			Veillonella	
	**			Lactococcus	
***	**			Eisenbergiella	
	**			Actinomyces	
	***			Desulfovibrio	
***	**			Pygmaiobacter	
	**			Angelakisella	
*	*			Peptococcus	
	*			Methanobrevibacter	
	***			Merdibacter	
	*			Anaerofilum	
*	**			Hydrogenoanaerobacterium	
	*			Ruminiclostridium	
***	***			Anaerofustis	
***	*			DNF00809	
Constipation	Low	High	Diarrhea		
	Normal	Normal			

 $\mathsf{Low}\;\mathsf{BMF}\leftrightarrow\mathsf{High}\;\mathsf{BMF}$

Self-reported diet, lifestyle, and anxiety and depression histories associated with BMF categories and demographic covariates

99 survey questions (see supplemental information; questions with sparse data were filtered out) on health, diet, and lifestyle were examined from 1,420 generally healthy individuals from the Arivale cohort in order to identify covariate-independent associations with BMF. Tests were run using the "polr" package in R (ordinal regression),²⁹ including the same set of covariates from the prior analyses, and with BMF coded as a categorical variable with high-normal BMF as the reference group (Figure 7A). Response categories for each question ascended ordinally in value or intensity (i.e., low to high), so that a positive association represented an increase in that variable. Across the 99 questions, the top results with significant odds ratios related to BMF categories were displayed relative to high-normal BMF (Figure 7A), colored by the variable category ("Diet/Lifestyle" or "Health/Digestion"). BMI, age, gender, and other covariates were also associated with many of these questionnaire-derived features, independent of BMF (Figure 7A). In particular, females tended to eat more vegetables and fruits in a week and had a higher diarrhea frequency. Males, on the other hand, showed higher weekly snack intake and easier bowel movements (Figure 7A). Unsurprisingly, constipation (lowest BMF range) was negatively associated with reported ease of bowel movement, and diarrhea was positively associated with self-reported diarrhea frequency (i.e., these were separate questions on the questionnaire) (Figure 7A). Those with higher weekly snack intake were more likely to be in the low-normal BMF category, and those with higher weekly vegetable intake, weekly fruit intake, greater ease of bowel movements, and higher self-reported diarrhea frequency were more likely to be in the high-normal BMF category (Figure 7A). Higher diarrhea frequency was significantly associated with having a higher BMI and with being younger relative to the rest of the cohort, while being older made one more likely to report having greater ease of bowel movement (Figure 7A). Finally, those with low LDL values (better cholesterol health) were more likely to report higher fruit intake, and those with low CRP (low inflammation) values were more likely to report higher vegetable intake (Figure 7A). These findings showcase a variety of common-sense dietary and lifestyle factors that could be leveraged to manage BMF, cardiometabolic, and immune health.

A subset of participants self-reported their history of depression and anxiety, including "self-current," "self-past," and "family" history of depression and anxiety (N = 2,096, see supplemental information; 11 questions related to anxiety and 23 related to depression). After logistic regression, 3 "true or false"-response questions related to history of depression in self and family history appeared marginally significant (logistic regression, FDR-cor-



rected p < 0.1), with a self-reported "true" response to a "family history of depression" showing a marginal association with constipation (logistic regression, FDR-corrected <0.1), a self-reported "true" response to a "sibling history of depression" showing a significant association with diarrhea (logistic regression, FDR-corrected <0.05), and a self-reported "true" response to "recent ailments in self-history of depression" showing a marginal association with low-normal BMF (logistic regression, FDR-corrected <0.1). Similarly, the same approach yielded a single marginal association between a "true" response to "self-past history of anxiety disorder" and low-normal BMF (logistic regression, FDR-corrected <0.1). Each of these associations was relative to the high-normal BMF reference category.

BMF-associated blood metabolites also associated with kidney function in a generally healthy cohort

Using the nine BMF-associated metabolites (ordered in ascending p value: PCS, X-23997, PAG, X-11850, PCG, X-12216, 3-IS, X-11843, and X-21310), an analysis was performed on all of the generally healthy Arivale participants with paired BMF, eGFR, and blood metabolomic data (N = 572). Using ordinary least squares (OLS), eGFR was regressed against BMF (encoded as a numerical variable between 1, 2, 3, and 4, with 1 being constipation, 2 being low-normal, 3 being highnormal, and 4 being diarrhea) and the nine BMF-related metabolites, which yielded a significant overall model (Figure S6B; OLS, $R^2 = 0.082$, p = 2.42E-7). Two of the BMF-associated metabolites showed significant beta-coefficients in the model: X-12216 and 3-IS (Figure S6B; OLS, $\beta_{X-12216} = -1.98$, p =1.20E-2 and $\beta_{3-1S} = -9.69$, p = 1.96E-5, respectively). These negative coefficients indicated that higher baseline levels of these blood metabolites were associated with lower kidney function.

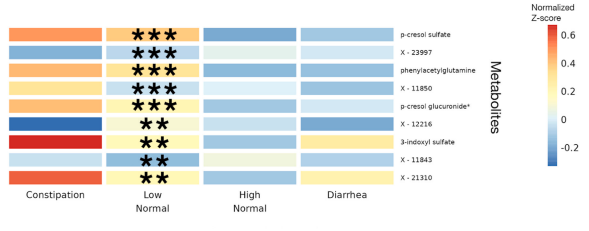
Finally, given that microbially derived 3-IS was independently associated with both eGFR and BMF, we hypothesized that 3-IS may be mediating, in part, the impact of BMF on eGFR. To test this hypothesis, we ran a causal mediation analysis (using the mediation library in R³¹; see STAR Methods) on the generally healthy Arivale individuals with BMF, eGFR, and the blood metabolomics data (N = 572; Figures 7B and S7). BMF categories were merged into "Low" (low-normal BMF and constipation) and "High" categories (high-normal BMF and diarrhea participants) in order to consolidate the BMF categories with very small Ns (i.e., constipation and diarrhea). The total effect of the overall model did not quite pass our significance threshold of alpha <0.05 (total effect, p = 0.064), but we saw a significant average direct effect (ADE) of BMF on eGFR (ADE = -4.458, p = 0.012) and a highly significant average causal mediation effect (ACME) of BMF via 3-IS on eGFR (ACME = 1.343, p < 2E-16; Figure 7B).

Figure 4. Heatmap of average Z scored CLR abundances within each BMF category for all annotated genera significantly associated with BMF

46 significant taxa, in order of decreasing average relative abundance, with their *Z* scored, CLR-transformed abundances averaged within each BMF category plotted as a heatmap. Covariates included gender, age, BMI, eGFR, LDL, CRP, A1C, and PCs 1–3. Asterisks denote the individual FDR-corrected significance threshold for the Wald test *p* value of the β_{BMF} coefficient for each BMF category, relative to the high-normal reference category. Rows without asterisks showed a significant overall model (FDR *p* value <0.05), despite a lack of significance for the individual coefficients. (***): *p* < 0.0001, (*): 0.001 < *p* < 0.01, (*): 0.001 < *p* < 0.05.



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Low BMF ↔ High BMF

Figure 5. Heatmap of average Z scored blood plasma metabolites levels within each BMF category for all metabolites significantly associated with BMF

11 significant blood plasma metabolites, with average *Z* scores within each BMF category plotted as a heatmap. Significant associations were identified using LIMMA, with FDR-corrected *p* values of the ratio test between the main model and the null model. Here, the covariates included gender, age, BMI, eGFR, LDL, CRP, A1C, and PCs 1–3. Asterisks denote metabolites with significant β_{BMF} coefficient(s) in the linear regression model after FDR correction. (***): *p* < 0.0001, (**): 0.0001 < *p* < 0.01, (*): 0.01 < *p* < 0.05.

DISCUSSION

In this study, we delve into the multi-omic fingerprint of crosssectional BMF variation in a large, generally healthy population (Figure 1). We find that aberrant BMFs were associated with variation in the ecological composition of the gut microbiota, plasma metabolite levels, clinical chemistries, diet, lifestyle, and psychological factors (Figures 4, 5, 6, and 7A). Overall, we observe an enrichment of microbially derived uremic toxins in the blood resulting from protein fermentation in the guts of individuals with lower BMFs. These toxins have been implicated in disease progression and mortality in CKD,^{21,32} and many of the same metabolites have been associated with other chronic diseases, like neurodegeneration.^{33,34}

Of the core set of covariates used in our regression analyses, only age, gender, BMI, and genetic ancestry PCs 1–3 were independently associated with BMF, with females, individuals with lower BMIs, and younger individuals showing lower average BMFs (Figure 2). Consistent with these results, women are known to be at higher risk of constipation and kidney dysfunction.^{35,36} In a prior study, individuals with lower BMIs were shown to produce less motilin (i.e., a hormone involved in gut motility) and were more likely to suffer from constipation.³⁷ Lower BMFs have also been linked to inflammation, oxidative stress, and cardiovascular disease risk.^{38,39} The associations between BMF and the first three principal components of genetic ancestry indicate a relationship between host genetics and BMF variation, which is further supported by a prior genome-wide association study.⁴⁰

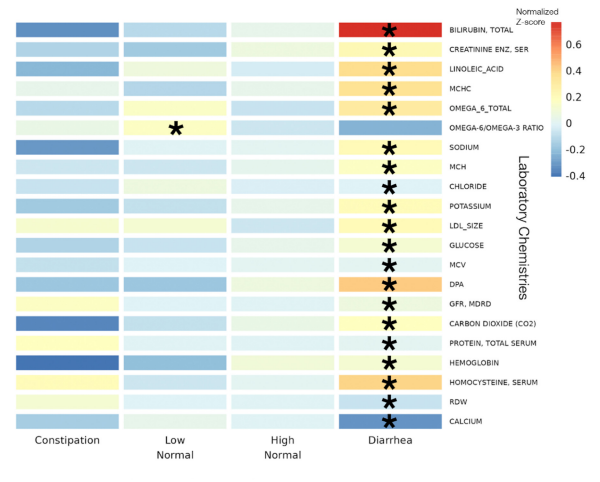
Independent of these covariates, several gut bacterial genera enriched in individuals with lower BMFs (CORNCOB, *p* < 0.001), such as *Christensenellaceae_R-7_group*, *Anaerotruncus*, *Blautia*, *Family_XIII_AD3011_group* (Anaerovoracaceae family), and *Methanobrevibacter*, were previously found to be enriched in patients with Parkinson's disease (PD), who often suffer from chronic constipation⁴¹. *Desulfovibrio*, which has been shown to be enriched in several disease states,⁴² was elevated at lower BMF (Figure 4). Another set of genera were depleted in lower BMF categories, such as *Bacteroides*, *Lachnoclostridium*, *Lachnospiraceae_ND3007_group*, *Lachnospiraceae_UCG-004*, and *Veillonella*, which are all important contributors to SCFA production.^{43–46} This reduction in SCFA producers is consistent with the switch away from saccharolytic fermentation toward proteolytic fermentation in the case of constipation.¹⁰ Reduced SCFA production is known to weaken smooth muscle contractions that drive peristalsis,^{47–49} acting as a positive feedback on constipation.

Consistent with our microbiome results, we found gut microbiome-derived protein fermentation byproducts, like PCS, PAG, and 3-IS, were enriched in the blood of individuals with lower BMFs (Figure 5).^{50–52} PCS has been associated with deteriorating kidney function and damage to nephrons, as well as cognitive decline and neuroinflammation.^{53,54} 3-IS has been associated with vascular disease and mortality in patients with CKD.⁵⁵ PAG has been associated with CKD progression and mortality.^{26,27,50,51} Ultimately, we see an enrichment in microbially derived uremic toxins in the blood of generally healthy individuals with lower BMFs.

Most of the clinical chemistry-BMF associations showed relative enrichment in the higher BMF category, and these features tended to reflect hepatic and nephrotic function. For example, high bilirubin can indicate liver disease from the overactive breakdown of red blood cells, but interestingly, higher bilirubin levels in serum coincide with a lower risk for CKD development and progression, which coincides with our observation that the lowest BMF categories had higher levels of uremic toxins but

Article





Low BMF \leftrightarrow High BMF



lower bilirubin levels.⁵⁶ Other metrics, like creatinine levels and linoleic acid levels, correlate positively with BMF and negatively with kidney function.^{57–59} In fact, most of the laboratory values, such as the mean corpuscular hemoglobin concentration, which measures the concentration of blood cells, can indicate kidney or liver disease.⁶⁰ It is interesting to note that biomarkers indicating kidney disease risk and progression were enriched at lower BMFs and biomarkers indicating liver disease risk and progression were enriched at higher BMFs in a generally healthy population, showing how aberrant BMF in either direction may increase chronic disease risk.

In addition to demographic factors associated with BMF, the questionnaire results indicate dietary and lifestyle factors that are known to influence BMF, like fruit and vegetable intake (i.e., sources of dietary fiber and polyphenols).^{36,38} We observed a lower fruit and vegetable intake and an increased likelihood of

snacking in the low-normal BMF category compared to the highnormal BMF category.^{23,36} We also found that constipation and diarrhea were marginally (and in one case, significantly) associated with self-reported measures of depression and anxiety, which aligns with prior work showing higher prevalence of anxiety and depression (between 22% and 33%) on the Hospital Anxiety and Depression Scale and the Mini International Neuropsychiatric Interview in patients with chronic constipation.⁶¹

Blood levels of 3-IS were independently associated with both BMF and eGFR, which led us to the hypothesis that 3-IS may mediate the potential influence of BMF on eGFR. Indeed, we observed a significant ADE of BMF on eGFR (p = 0.012) and a highly significant ACME for 3-IS (p < 2E-16; Figure 7B). Together, these results indicate that aberrant BMF-associated increases in 3-IS are associated with declining kidney function in a generally healthy cohort, which is consistent with similar



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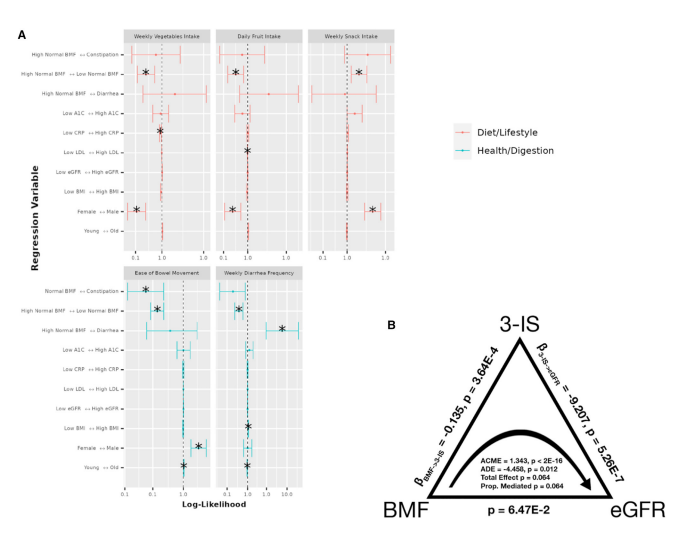


Figure 7. Ordinal regression odds ratio for health, diet, and lifestyle survey data vs. BMF and covariates, and causal mediation analysis, with BMF as the treatment variable, 3-IS as the mediator variable, and eGFR as the response variable

(A) BMF categories are shown by question type (diet/lifestyle or health/digestion). "High-normal" BMF (7-21/week) is the reference. Vertical ticks show directional associations in likelihood between variables across the horizontal axis. The center line (x = 1.0) indicates equal likelihood of increased values on either side. Confidence intervals not crossing the line are significant associations (FDR p < 0.05).

(B) BMF affects eGFR directly (average direct effect, ADE) and indirectly (average causal mediated effect, ACME) through 3-IS (a metabolite). Both effects are significant (N = 572); the total effect was not significant (N = 572; ADE = -4.458, p = 0.012; ACME = 1.343, p < 2E-16). The total effect and the proportionmediated terms did not pass our significance threshold of a = 0.05.

associations that have been observed between 3-IS and poorer outcomes in patients with CKD.55

Bowel movement abnormalities, such as constipation or diarrhea, have been linked to diseases ranging from enteric infections¹⁶ to many chronic diseases like CKD and IBD and neurodegenerative conditions like Alzheimer's and PD.33,62,63 Indeed, even in the context of our generally healthy cohort, we see the build-up of microbially derived uremic toxins in the blood of individuals with lower BMFs. Perhaps, most concerning was our observation that aberrant BMF-associated microbial metabolite 3-IS was also associated with lower eGFR values. Our results underscore common-sense dietary and lifestyle changes, like increasing intake of fruits and vegetables, which may help to normalize BMF and perhaps reduce BMF-associated risk. We suggest that chronic constipation or diarrhea may be underappreciated drivers of organ damage and chronic disease, even in healthy populations, although additional work is required to rigorously quantify how BMF impacts disease risk.

Limitations of the study

There are some important limitations to consider when interpreting the results of this study. The generally healthy cohort studied here was overwhelmingly "White," predominantly female, and from the West Coast of the USA, which limits the generalizability of our results. In addition, the diet, lifestyle, and mood data were self-reported and subject to biases and errors, although BMF was binned into four coarse-grained categories in an attempt to mitigate self-reporting bias. BMF is not synonymous with transit time through the gut, which can be measured through means like the "blue dye method,"⁷ although BMF still appears



to be a useful and interesting measure of self-reported bowel habit differences in this study. We had limited representation in the constipation and diarrhea categories, which reflects the "generally healthy" nature of this cohort, but this also limited our statistical power for detecting associations in these groups. Detailed renal assessments beyond eGFR and clinical laboratory chemistries were also not available in this study. Additionally, the dietary variables that were associated with better BMF outcomes (i.e., increased dietary fiber intake, in the form of fruits and vegetables) are not devoid of clinical risk and may not be appropriate for everyone. For example, high-fiber diets can sometimes lead to bloating and inflammation in patients with IBD.⁶⁴ Furthermore, patients with CKD are often coached to limit their intake of fiber-rich foods because they can contain high levels of potassium and phosphorus.⁶⁵ However, low-fiber diets may act as a positive feedback on constipation and inflammation. This highlights the importance of intervening at the prodromal stage, before disease manifests, when a healthy, largely plant-based diet is well tolerated. While we find some evidence for microbially derived, BMF-associated uremic toxins in blood influencing kidney function in a generally healthy cohort, more work is needed to establish a link between longer-term BMF management and chronic disease risk. In the mediation analysis, we did not see a strong total model effect, despite seeing significant direct and mediation effects. This kind of result is expected when the total and mediation effects are of a similar magnitude, when there are opposing effect directions between treatments and mediators, or when there are other more complicated effects (e.g., non-linear associations).⁶⁶ Ultimately, future intervention trials should be done to assess the potential for managing BMF throughout the lifespan as a strategy to reduce chronic disease risk

STAR * METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j. xcrm.2024.101646.

ACKNOWLEDGMENTS

We thank Amy Willis for helpful advice on ordinal regression and members of the Gibbons, Hood-Price, and Hadlock labs for helpful discussions on this work. This research was funded by Washington Research Foundation Distinguished Investigator Award and by startup funds from the Institute for Systems Biology (to S.M.G.). Research reported in this publication was supported by the National Institute of Diabetes and Digestive and Kidney Diseases of the National Institutes of Health (NIH) under award no. R01DK133468 (to S.M.G.) and by a Global Grants for Gut Health Award from Nature Portfolio and Yakult (to S.M.G.). The content is solely the responsibility of the authors and does not necessarily represent the official views of the NIH. The funders had no role in designing, carrying out, or interpreting the work presented in the manuscript. Figure 1 was created with BioRender.com and the rest of the figures were created with R, Adobe Photoshop, and Python libraries.

AUTHOR CONTRIBUTIONS

J.P.J.-M. and S.M.G. conceived the study. J.P.J.-M. conducted the analyses, wrote the code, and wrote the first draft of the manuscript. S.M.G. provided supervision. C.D., T.W., A.E.L., and A.R. contributed code and expert input on the analyses. A.E.L., T.W., D.L.S., A.R., J.H., A.T.M., L.H., and N.R. contributed to interpretation of the results and editing the final manuscript.

DECLARATION OF INTERESTS

L.H. is a former shareholder of Arivale. A.T.M. was a former employee of Arivale. Arivale is no longer a commercially operating company as of April 2019.

DECLARATION OF GENERATIVE AI AND AI-ASSISTED TECHNOLOGIES IN THE WRITING PROCESS

During the preparation of this work, the first author used Google Gemini (formerly Bard) and Microsoft Copilot in order to query primary sources in the references, suggest minor revisions of written excerpts, and debug code. After using this tool/service, the authors reviewed and edited the content as needed. The authors take full responsibility for the content of this publication.

Received: April 11, 2023 Revised: February 22, 2024 Accepted: June 14, 2024 Published: July 16, 2024

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STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Software and Algorithms		
Custom Python and Jupyter Notebooks and R markdown notebooks	This study	https://github.com/Gibbons-Lab/Aberrant- BMF-Cell-Reports
R and RStudio	Posit.co ⁶⁷	https://posit.co/download/rstudio- desktop/
CORNCOB R Package	Martin et al. ³⁰	https://github.com/statdivlab/corncob
Bioconductor R Package	Bioconductor ^{30,68}	https://www.bioconductor.org/install/
tidyverse R Packages	tidyverse ⁶⁹	https://github.com/tidyverse/tidyverse/
LIMMA R Package	Smyth et al. ^{70,71}	https://kasperdanielhansen.github.io/ genbioconductor/html/limma.html
POLR/MASS R Package	Venables and Ripley et al. ²⁹	https://rdocumentation.org/packages/ MASS/versions/7.3-60.0.1

RESOURCE AVAILABILITY

Lead contact

Additional requests and information regarding resources, experimental materials, reagents, and assay vendors should be directed to and will be fulfilled by the lead contact, Sean Gibbons (sgibbons@isbscience.org).

Materials availability

This study did not generate reagents.

Data and code availability

- Code used to analyze 16S rRNA gene amplicon sequencing data can be found at https://github.com/gibbons-lab/mbtools.
- Code used to run the statistical analyses described in this paper is available at https://github.com/Gibbons-Lab/Aberrant-BMF-Cell-Reports .
- Qualified researchers can access the full Arivale deidentified dataset, including all raw data, supporting the findings in this study for research purposes through signing a Data Use Agreement (DUA). Inquiries to access the data can be made at data-access@ isbscience.org and will be responded to within 7 business days.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Institutional review board approval for the study

The procedures for this study were reviewed and approved by the Western Institutional Review Board, under the institutional review board study number 20170658 for the Institute for Systems Biology and 1178906 for Arivale, Inc.

Generally healthy cohort

All study participants were subscribers in the Arivale Scientific Wellness program (2015–2019) and provided informed consent for the use of their anonymized, de-identified data for research purposes. Participants were community-dwelling, residents of Washington State and California (which are slightly leaner and healthier than other parts of the USA), over the age of 19, non-pregnant, but were not pre-screened for the presence or absence of any particular disease. Participants provided detailed questionnaire data that included self-reported information about medical conditions and medications, along with blood and stool samples that were used to generate blood plasma metabolomics, proteomics, clinical laboratory chemistries, and gut microbiome data (Figure 1; Table S1). 3,955 individuals with at least one BMF data point available from the Arivale program were initially considered for this analysis. Their data were merged in Python (Jupyter Notebooks) with those with eGFR data (N = 6,133) and those with available survey (N = 5,764), ASVs (N = 3,694), proteomics (N = 2,859), clinical laboratory chemistries (N = 4,881), or metabolomics (N = 2,043) data samples to yield dataframes representative of the cohorts at the intersections of these merges. The final dataframes for the regressions for survey (N = 1,420), ASVs (N = 1,062), proteomics (N = 523), clinical laboratory chemistries (N = 1,425), and metabolomics (N = 486) cohorts were used for downstream regression analyses (Figure S6).



Only baseline time point samples were used for each participant for the baseline 'omics analyses. A 30% prevalence filter was implemented across the gut microbiome, blood plasma metabolomics, proteomics, chemistries, and ordinal questionnaire data analyses. This meant that each feature in the data could contain no more than 70% missing data from the final cohort of samples in order to be retained for downstream analysis. For microbiome analyses, a filtered subcohort of 1,062 individuals with ASV-level taxa counts, BMF, gender, age, eGFR, BMI, LDL, CRP, A1C, and genetic ancestry data were selected. This filtering resulted in a total of 135 genera. For the metabolomics analysis, a cohort of 486 participants with BMF, gender, age, eGFR, BMI, CRP, LDL, A1C, PC1, PC2, and PC3, and blood metabolomics data were selected. 956 metabolites were retained for downstream analyses. 274 blood proteins that met the prevalence (\geq 50%) filter in the cohort of 823 individuals were retained for downstream analyses. A \geq 30% prevalence filter was applied to yield 1,425 samples with blood plasma clinical laboratory chemistries data, resulting in 55 features retained for downstream analyses. Similarly, for ordinal regression of the questionnaire data (e.g., diet, lifestyle, and stress/pain/ health factors) using the respective R package, polr,²⁹ we collated all the responses and filtered out questions that contained more than 10% "NAs" (≥90% prevalence; and for binary variables in downstream depression/anxiety analyses: ≥10% affirmative or "True" responses). We also excluded binary response variables for the general survey questionnaire analysis (separate from the anxiety/depression analysis, which leveraged binary response features), which are incompatible with ordinal regression, resulting in 138 variables across 1,420 participants, in addition to having paired data on age, gender, eGFR, BMI, BMF, CRP, LDL, A1C, PC1, PC2, and PC3. The final features considered needed to retain at least 2 non-missing factors (or categories) and contain at least 10 responses per category, which resulted in 99 features. BMF data was captured from responses to a survey question on how many bowel movements an individual has per week, on average. The available responses to this question were: (1) Twice per week or less; (2) 3-6 times per week; (3) 1-3 times daily; or (4) 4 or more times daily. While the normal range of BMF encompasses both the second and third responses to this question (i.e., between three times a week and three times a day),⁷² we chose to define 1-3 times per day (high-normal) as the reference group for the purposes of regression.

Finally, we imposed disease-related exclusion criteria, in order to generate a "generally healthy" sub-cohort. These include any participants who reported affirmative or "true" to a history of taking cholesterol, laxative, or blood pressure medication, as well as those who reported a self- or family- history presence of the following diseases: bladder or kidney disease, inflammatory bowel disease (IBD), celiac disease, diverticulosis, gastroesophageal reflux disease (GERD), irritable bowel syndrome (IBS), or peptic ulcers (see Table S1 in the supplemental information). 988 (25%) out of the initial 3,955 Arivale individuals with BMF data were excluded by these filters.

METHOD DETAILS

Gut microbiome data

Fecal samples from Arivale participants were collected (described in Diener et al.¹³ and detailed here) from proprietary at-home kits developed by two microbiome vendors (DNA Genotek and Second Genome). Using the KingFisher Flex instrument, the MoBio PowerMag Soil DNA isolation kit⁷³ (QIAGEN) enabled the isolation of stool DNA from 250 mL of homogenized human feces, after performing an additional glass bead-beating step. Qubit measurement and spectrophotometry were also performed using an A260/A280 absorbance ratio.

16S amplicon sequencing was run on a MiSeq⁷⁴ (Illumina, USA) with either paired-end 300-bp protocol (DNA Genotek) or pairedend 250-bp protocol (SecondGenome). The FASTQ files were provided by the Illumina Basespace platform after the phiX reads were removed with basecalling. Length cutoffs of 250-bp for the forward reads and 230-bp for the reverse reads were employed. Any reads with more than 2 expected errors or ambiguous base calls under the Illumina error model were eliminated. Over 97% of the reads passed these filters, resulting in approximately 200,000 reads per sample.

Final truncated and filtered reads were then used to infer amplicon sequence variants (ASVs) with DADA2.^{75–77} Each sequencing run separately resulted in its own error profiles. The final ASVs and counts were then joined, with chimeras removed using DADA2's "consensus" strategy. After this step, ~16% of reads were removed. Taxonomic assignment of ASVs was then achieved using the naive Bayes classifier in DADA2 with the SILVA database (version 128).⁷⁸

Nearly 90% of the ASVs were classified down to the genus level, which was the taxonomic level chosen for this analysis. 3,694 samples across 609 taxa were available from these methods, which were then filtered down to 135 taxa after using a 30% prevalence filter. Samples were rarefied to an even depth of 13,703 reads prior to calculating alpha-diversity metrics (using the "rarefy_even_depth()" function in the phyloseq R package⁷⁹; rng seed = 111). ASV richness (Observed ASVs), Shannon Diversity, and Pielou's evenness were calculated. Merging with covariate data resulted in 1,062 samples with 135 taxa for downstream analyses.

Olink proteomics

Blood plasma proteomic data were generated by Olink Biosciences using the ProSeek Cardiovascular II, Cardiovascular III, and Inflammation arrays. The proteins were filtered down to 274 proteins and 823 samples, retaining proteins with \geq 50% prevalence across samples and samples with the full set of covariate data. Post-filtering, NAN values were assumed to be below detection and imputed to be the median across samples for that particular protein. The values used for the proteomics analysis were from protein readings previously batch-corrected and normalized based on the overlapping reference samples within the batch plates (i.e., a set of Arivale plasma samples that are run with each batch). The corrected values were also scale-shifted to the reference sample and



the original delivered data (using the seventh run as a baseline). Olink's Proximity Extension Assay (PEA), a 2-antibody-barcode technology,^{75,76,80} is used to tag protein biomarkers with a proximity probe (which binds specifically to the target protein biomarker) and an extension probe (which carries a DNA barcode sequence) as described by Illumina in conjunction with Olink.^{81–83} Once both probes bind to each other due to a protein-protein interaction or by proximity, they trigger the activation of the extension probe, beginning the hybridization of the probe with a detection bead's complementary DNA sequence. Each bead contains an individual identifier, which allows target proteins to be decoded according to a barcode. These methods are also described further in Zubair et al.⁷⁵

Metabolon metabolomics

Metabolon obtained metabolomics data on the previously mentioned plasma samples using preparation, quality control, and collection methods described in previous studies.⁷⁶ During sample processing, the plasma samples were thawed and proteins were removed using methanol extraction. Samples were then divided into 5 fractions including a backup fraction. Organic solvents were removed using TurboVap and measurements were then performed using high-performance liquid chromatography (HPLC) and high-resolution mass spectrometry (MS).⁸⁴ Four separate measurements were performed using different fractions combinations: positive-ion and negative-ion modes optimized for both hydrophobic and hydrophilic compounds. Batch correction was performed using quality control samples (i.e., a set of Arivale plasma samples that were run with each batch) and abundance data were normalized to these quality control samples. Metabolites were annotated according to 3 standards: Tier 1, matching to an internal standard; Tier 2, matching to a published MS spectrum; or Tier 3, matching to a known chemical formula. Unknown metabolites were unannotated and labeled with an "X - "label followed by an identifier.⁸⁵ 956 total metabolites showed at least 70% prevalence across 486 samples. In this analysis, missing values were imputed to be the median of the non-missing samples for each metabolite, and final downstream metabolites were log-transformed and merged with the full set of covariates.

For the multi-linear regression and causal mediation analysis, those with paired eGFR, BMF-associated metabolomics results, and BMF were filtered using the "generally healthy" exclusionary criteria and the previously mentioned prevalence filtering for metabolomics. The remaining individuals (Figures 7B and S7; N = 572) were processed in a multi-linear regression (OLS) with eGFR ~ BMF (encoded as a value between 1 and 4 with 4 being diarrhea or the highest BMF) + the obtained metabolomics values for the 9 BMF-associated metabolites (Figure S6). The other multi-omics covariates (gender, age, BMI, CRP, LDL, A1C, and PC1-PC3) were not considered for the subsequent mediation analysis (Figure 7B; N = 562), which was performed using a mediation model with the mediate() function from the mediation package in R.³¹ Using this modeling function, the outcome model was specified as eGFR ~ 3-IS + BMF (where BMF was encoded as a binary categorical variable, with "Low" including those with low-normal BMF and constipation, and "High" containing those with high-normal BMF and diarrhea. "Low" was the control value for BMF and "High" was the treatment value) and the mediation model was assumed to be 3-IS ~ BMF. ACME and ADE values were obtained from the model and reported using the diagram in Figure 7B. A GLM was also performed between eGFR ~ BMF, 3-IS ~ BMF, and eGFR ~ 3-IS to obtain the β -coefficients and p-values for the relationships between the mediated variables (Figure 7B).

Blood plasma chemistries

LabCorp and Quest⁸⁶ phlebotomists collected blood from Arivale participants using methods described previously by Wilmanski et al. and others.¹³ Individuals were asked to abstain from alcohol, vigorous exercise, monosodium glutamate and aspartame at least 24 h prior to drawing of the blood, as well as fasting at least 12 h beforehand. Blood samples were collected for clinical chemistries, metabolomics and proteomics at the same time, and within 21 days of stool sampling. BMI was calculated from weight and height using the following formula $BMI = \frac{weight (kg)}{(height (m))^2}$. 4,881 samples and 68 laboratory values were filtered down using the same prevalence filtering as the metabolomics data. 1,425 samples and 55 chemistries were retained. The final 55 features were log-transformed, with missing samples imputed to be the median value of the non-missing samples. These features were merged with the full set of covariates. eGFR was calculated based on the CKD Epidemiology Collaboration (CKD-EPI) creatinine Equation, as recommended by the current guidelines of the National Kidney Foundation⁸⁷: eGFR_{cr} = 142 × min(Scr/κ, 1)^α × max(Scr/κ, 1)^{-1.200} × 0.9938^{Age} × 1.012 [if female], where Scr = standardized serum creatinine in mg/dL, $\kappa = 0.7$ (female) or 0.9 (male), and $\alpha = -0.241$ (female) or -0.302 (male).

Questionnaire data

3,482 self-reported questionnaire features were retrieved across 5,764 Arivale participants. After health and prevalence filtration, 138 features remained, which were subsequently filtered down again to 99 features by removing factored features with fewer than 10 responses per level and keeping features with at least 2 non-missing levels to the factor. Category responses were organized and numbered to be ordinally ascending in magnitude or intensity, with relatively even-spaced differences in magnitude between categories wherever possible (i.e., for a factored feature with levels from 1, ...,n, the level labeled "1" represented responses such as "Strongly Disagree", "Never", "None", or the lowest frequency/intensity, and the level labeled "n" represented responses such as "Strongly Agree", "Always", or the greatest frequency or intensity). These features were merged with the full set of covariate data.



Depression and anxiety health history data

We used logistic regression to scrutinize associations between 23 (anxiety) and 35 (depression) binary ("true" or "false") self-reported questions based on "self-current", "self-past", and "family" histories of depression or anxiety, with depression or anxiety encoded as a binary dependent variable, and BMF encoded as a categorical independent variable, and with the standard set of covariates.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical analyses

Using Python (Jupyter Notebooks)⁸⁸ and R (RStudio)⁶⁷ linear regressions were performed. The response variables were either: centered log ratio-transformed bacterial genus data, log-transformed plasma metabolomics data, batch-corrected plasma proteomics data, log-transformed plasma chemistries data, or ordinal response variables from questionnaire data, depending on the analysis. For the blood proteomics, plasma chemistries, and metabolite associations, generalized linear regression models were run using the LIMMA package in R.⁷⁰ BMF was encoded as a categorical variable (or in the case of analyzing alpha-diversity, it was also computed as an ordinal variable with a linear model coefficient) with categories: 1 = constipation (1-2 bowel movements per week), 2 = low-normal (3-6 bowel movements per week), 3 = high-normal (1-3 bowel movements per day), and 4 = diarrhea (4 or more bowel movements per day). To begin characterizing the main variables in the cohorts: BMF and eGFR, a POLR regression²⁹ (N = 1,425) was performed on BMF (encoded as an ordinal variable with categories "Constipation", "Low Normal", "High Normal", and "Diarrhea" BMF in ascending order of magnitude) \sim eGFR + other covariates (gender, age, BMI, CRP, LDL, A1C, PC1, PC2, and PC3). Similarly, a GLM (N = 1.425) was computed for eGFR \sim BMF (also encoded ordinally) + other covariates (gender, age, BMI, CRP, LDL, A1C, PC1, PC2, and PC3). These were used to determine the significant covariates affecting each subsequent analysis (Figure 2). Next, in each baseline regression, the following covariates were all included: gender, age, BMI, eGFR, CRP, LDL, A1C, PC1, PC2, and PC3. Gut bacterial genus-level counts were modeled with a beta-binomial distribution using the CORNCOB package in R.³⁰ For the questionnaire data (ordinal response categories across diet, exercise, stress, pain, and other lifestyle factors), polr in R was used for the ordinal regression analysis (POLR).²⁹ For the anxiety and depression data, which were binary in response ("True" or "False"; Nonresponders to each feature were not considered and features were filtered to have at least 5 non-missing responses for each binary outcome), logistic regression was performed using the "glm(family = "binomial")" function in R. All questionnaire and anxiety/depression response modeling results were FDR-corrected for significance. Finally, for the Arivale cohort, the initial time point or baseline value for eGFR was obtained alongside the initial or earliest time point sample for the BMF-related metabolites. eGFR was regressed against the BMF-associated metabolites in an OLS-based linear regression to determine visible effects of these metabolites on our available samples. Finally, a mediation analysis was run using the mediate() function in the mediation library available for R^{31} on the individuals who met the "generally healthy" exclusion criteria with paired eGFR, BMF, and 3-IS data. BMF was the treatment variable, 3-IS was the mediator, and eGFR was the response variable. ACME, ADE, total effect and proportion mediated were determined with nonparametric bootstrap confidence intervals.

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Supplemental information

Aberrant bowel movement frequencies coincide

with increased microbe-derived blood metabolites

associated with reduced organ function

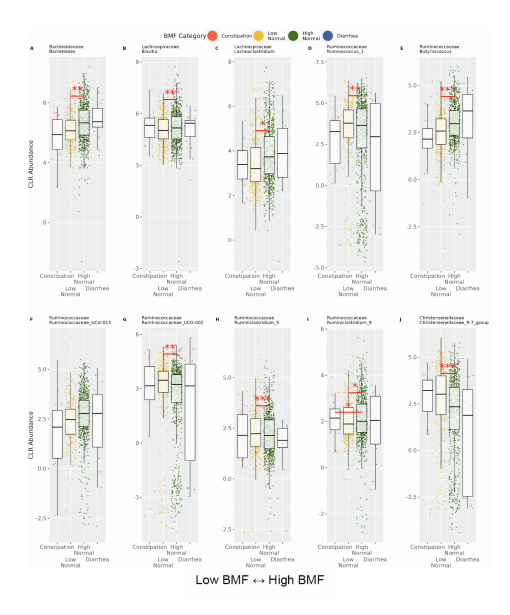
Johannes P. Johnson-Martínez, Christian Diener, Anne E. Levine, Tomasz Wilmanski, David L. Suskind, Alexandra Ralevski, Jennifer Hadlock, Andrew T. Magis, Leroy Hood, Noa Rappaport, and Sean M. Gibbons

1 SUPPLEMENTAL FIGURES AND TABLES

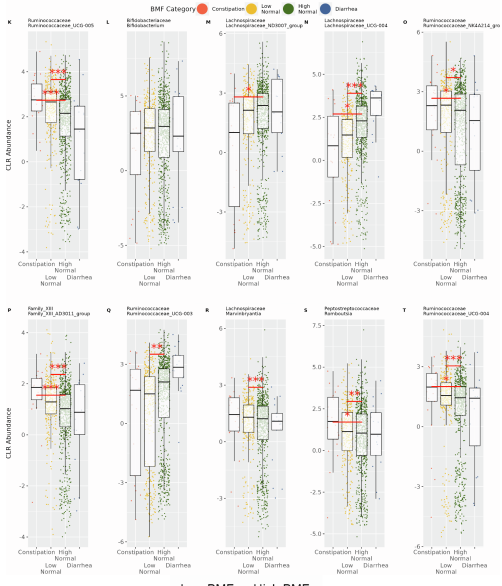
Covariates:	Mean ± standard deviation, or % across Arivale:			
Gender	65.1% Female			
BMI	27.2 ± 5.89			
Age	46.36 ± 12.96			
eGFR	89.07 ± 20.20			
CRP	2.40 ± 4.76			
LDL	114.17 ± 33.77			
A1C	5.49 ± 0.57			
Highlighted exclusionary criteria:				
Percent with self-reported kidney disease:	3.00% (119 out of 3,955 participants with BMF data available withheld from cohort)			
Percent IBS or IBD:	3.23% (128 out of 3,955 participants with BMF data available withheld from cohort)			
Self - current history - bladder infection Self - current history - kidney disease				
Self - current history - kidney infection Self - current history - kidney stones				
Self - current history - kidney stones Self - current history - bladder/kidney - other				
Self - current history - polycystic kidney disease (PKD)				
Self - current history - urinary incontinence				
Self - current history - kidney cancer				
Self - current history - celiac disease				
Self - current history - colonic Crohn's disease				
Self - current history - diverticulosis				
Self - current history - gastroesophageal reflux o	disease (GERD)			
Self - current history - ileal Crohn's disease				
Self - current history - irritable bowel syndrome	(IBS)			
Self - current history - inflammatory bowel disease (IBD)				
Self - current history - ulcerative colitis				
Self - current history - peptic ulcer				
Self - laxatives usage				
Self - anticoagulation or cholesterol drugs usage	e			
Self - blood pressure drugs usage				

2

3 Table S1. The modeling covariates and exclusionary criteria, related to results in Figures 2-7. Out of the 3,955 total Arivale participants that had BMF data, 3.00% self-reported kidney 4 5 disease (the kidney-related questions in the exclusionary features) and 3.23% self-reported IBS 6 or IBD. An initial baseline cohort of 3,132 participants that had health history survey questionnaire 7 data was available. The participants that answered affirmatively to the exclusionary features were 8 removed from the analysis, resulting in 25% of the initial cohort with BMF data being filtered down 9 to N = 1,561, and subsequently, a final baseline cohort of 1,425 individuals after merging for 10 covariates.



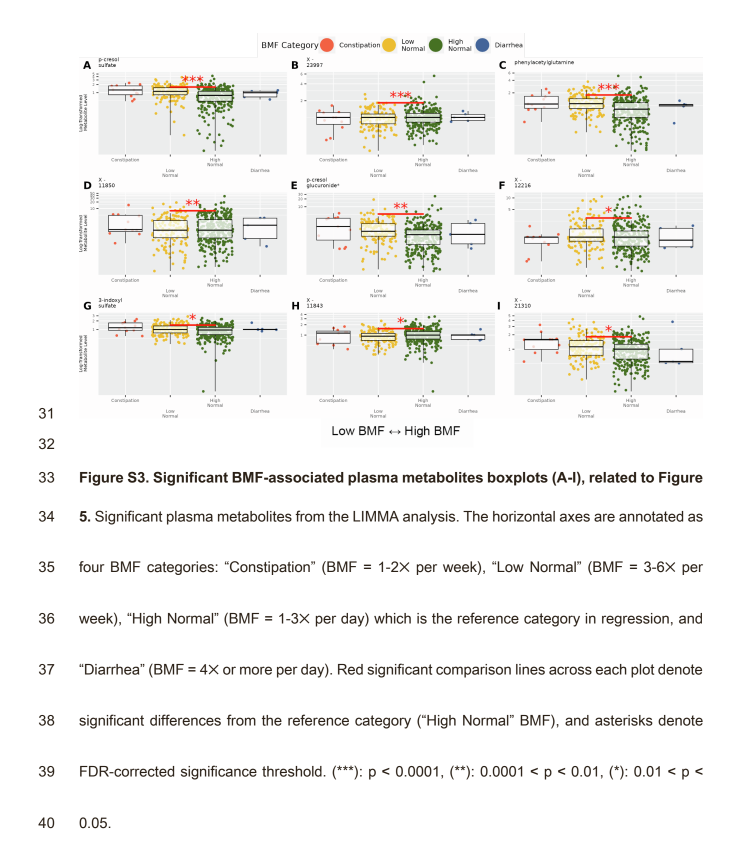
13	Figure S1. The top 10 most abundant genera significantly associated with BMF (A-J),
14	related to Figure 4. Significant genera from the CORNCOB analysis in order of decreasing CLR-
15	transformed abundance. The line in each plot denotes significant differences from the reference
16	category ("High Normal" BMF), and asterisks denote FDR-corrected significance threshold. (***):
17	p < 0.0001, (**): 0.0001 < p < 0.01, (*): 0.01 < p < 0.05. The horizontal axes are annotated as four
18	BMF categories: "Constipation" (BMF = $1-2\times$ per week), "Low Normal" (BMF = $3-6\times$ per week),
19	"High Normal" (BMF = $1-3 \times$ per day) which is the reference category in regression, and "Diarrhea"
20	(BMF = $4 \times$ or more per day).

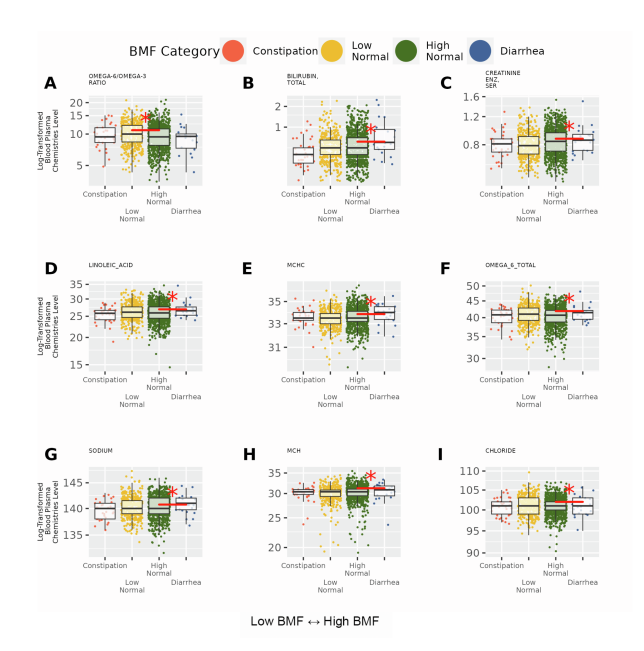


Low BMF \leftrightarrow High BMF

Figure S2. The top 11-20 most abundant genera associated with BMF (K-T), related to Figure 4. Significant genera from the CORNCOB analysis in order of decreasing CLRtransformed abundance. The line in each plot denotes significant differences from the reference category ("High Normal" BMF), and asterisks denote FDR-corrected significance threshold. (***): p < 0.0001, (**): 0.0001 , (*): <math>0.01 . The horizontal axes are annotated as fourBMF categories: "Constipation" (BMF = 1-2× per week), "Low Normal" (BMF = 3-6× per week),"High Normal" (BMF = 1-3× per day) which is the reference category in regression, and "Diarrhea"

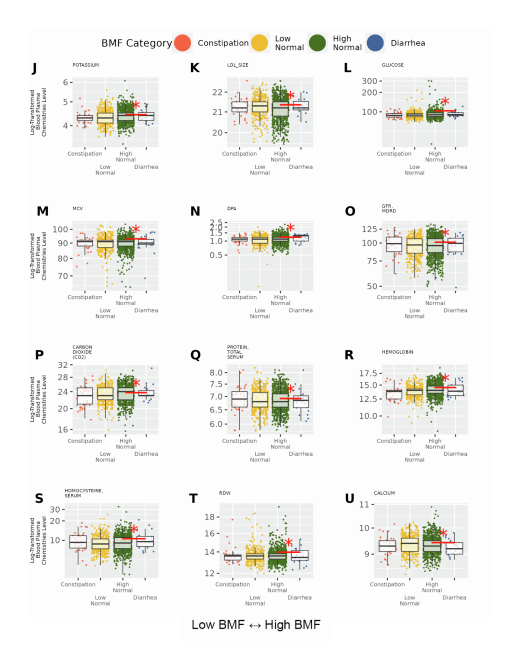
30 (BMF = $4 \times$ or more per day).



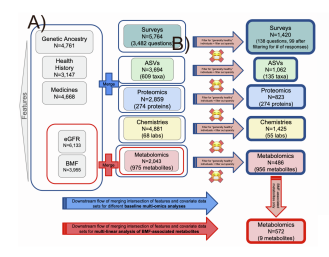


42	Figure S4. Significant BMF-associated clinical chemistries boxplots (A-I), related to Figure
43	6. Significant clinical chemistries from the LIMMA analysis. The horizontal axes are annotated as
44	four BMF categories: "Constipation" (BMF = $1-2\times$ per week), "Low Normal" (BMF = $3-6\times$ per
45	week), "High Normal" (BMF = $1-3 \times$ per day) which is the reference category in regression, and
46	"Diarrhea" (BMF = $4 \times$ or more per day). Red significant comparison lines across each plot denote
47	significant differences from the reference category ("High Normal" BMF), and asterisks denote
48	FDR-corrected significance threshold. (***): p < 0.0001, (**): 0.0001 < p < 0.01, (*): 0.01 < p <

49 0.05.



51	Figure S5. The remaining significant BMF-associated clinical chemistries boxplots (J-U),
52	related to Figure 6. The remaining significant clinical chemistries from the LIMMA analysis. The
53	horizontal axes are annotated as four BMF categories: "Constipation" (BMF = $1-2\times$ per week),
54	"Low Normal" (BMF = $3-6\times$ per week), "High Normal" (BMF = $1-3\times$ per day) which is the reference
55	category in regression, and "Diarrhea" (BMF = $4 \times$ or more per day). Red significant comparison
56	lines across each plot denote significant differences from the reference category ("High Normal"
57	BMF), and asterisks denote FDR-corrected significance threshold. (***): p < 0.0001, (**): 0.0001
58	< p < 0.01, (*): 0.01 < p < 0.05.



OLS Regression Results							
Dep. Variable: eGFR R-squared: 0.082							
Model:				R-squared:		0.067	
Method:	Least Squa					5,547	
Date:				(F-statistic):		2,42e-07	
Time:				Likelihood:		-2465.4	
No. Observations:			AIC:			4951.	
Df Residuals:		62	BIC:			4994.	
Df Model:		9					
Covariance Type:	nonrob	st					
	coef	sto	d err	t	P> t	[0.025	0.975]
const	115.0755		.841		0.000		
bowel	-3.9902	1	1.496	-2.667		-6.929	
p-cresol sulfate				-1.088		-7.548	
X – 23997				1.200		-1.087	
phenylacetylglutamine				0.982		-2.207	
X - 11850				-1.232		-1.147	
p-cresol glucuronide				0.750		-0.595	
X - 12216				-2.520		-3.528	
3-indoxyl sulfate						-14.104	
X - 11843	0.0527	6	.554	0.095	0.924	-1.036	1.141
Omnibus:				in-Watson:		1.841	
Prob(Omnibus):				ue-Bera (JB):		22.585	
Skew:		90	Prob			1.25e-05 45.4	
Kurtosis:	2	:99	cond	. No.		45.4	

60	Figure S6. A) Flow Chart for Cohort Selection of Baseline Population, related to the
61	Generally-healthy cohort section of the STAR Methods. B) OLS regression resulting from
62	eGFR ~ BMF-associated metabolites + BMF, related to Figure 7B. A) Individuals with the full
63	complement of covariate data (gender, age, BMI, and CRP, LDL, A1C, and PCs 1-3) were further
64	filtered for having available baseline data for each of the following: surveys, microbiome profiles,
65	proteomics, clinical chemistries (e.g. complete blood count, or CBC; and comprehensive
66	metabolic panel, or CMP) and metabolomics. The "generally-healthy" exclusion criteria were then
67	imposed (38.5% excluded; see Method Details), along with sparsity or non-missingness
68	minimums for the features in the 'omics data (\geq 30% prevalence for gut microbiome data,
69	metabolomics and clinical chemistries; \geq 50% prevalence for proteomics; and \geq 90% prevalence
70	and \geq 10% affirmative for binary responses in the survey questions). These filters resulted in the
71	final sub-cohort numbers shown on the right side of the figure in blue outlines. Additionally, the
72	eGFR and BMF data frames were merged with the metabolomics data frame and filtered by the
73	"generally-healthy" exclusionary criteria for the 9 BMF-associated metabolites eGFR regression
74	and mediation analysis. B) The p-value for the overall generalized-linear model (eGFR \sim BMF-
75	related metabolites) was significant (N = 572, p = 2.42E-7, R^2 = 0.082) and so were the p-values
76	of the individual β -coefficients for 3-IS (β_{3-IS} = -9.69, p = 1.96E-5), BMF (denoted "bowel"; β_{BMF} =
77	-3.99, p = 7.88E-3), and X - 12216 ($\beta_{X-12216}$ = -1.98, p = 1.20E-2).