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ORIGINAL RESEARCH

Fecal Microbiome Composition Does Not Predict Diet-Induced TMAO Production in Healthy Adults

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BACKGROUND: Trimethylamine-*N*-oxide (TMAO) is a small molecule derived from the metabolism of dietary nutrients by gut microbes and contributes to cardiovascular disease. Plasma TMAO increases following consumption of red meat. This metabolic change is thought to be partly because of the expansion of gut microbes able to use nutrients abundant in red meat.

METHODS AND RESULTS: We used data from a randomized crossover study to estimate the degree to which TMAO can be estimated from fecal microbial composition. Healthy participants received a series of 3 diets that differed in protein source (red meat, white meat, and non-meat), and fecal, plasma, and urine samples were collected following 4 weeks of exposure to each diet. TMAO was quantitated in plasma and urine, while shotgun metagenomic sequencing was performed on fecal DNA. While the cai gene cluster was weakly correlated with plasma TMAO (rho=0.17, *P*=0.0007), elastic net models of TMAO were not improved by abundances of bacterial genes known to contribute to TMAO synthesis. A global analysis of all taxonomic groups, genes, and gene families found no meaningful predictors of TMAO. We postulated that abundances of known genes related to TMAO production do not predict bacterial metabolism, and we measured choline- and carnitine-trimethylamine lyase activity during fecal culture. Trimethylamine lyase genes were only weakly correlated with the activity of the enzymes they encode.

CONCLUSIONS: Fecal microbiome composition does not predict systemic TMAO because, in this case, gene copy number does not predict bacterial metabolic activity.

REGISTRATION: URL: https://www.clinicaltrials.gov; Unique identifier: NCT01427855.

Key Words: fecal microbiome
metagenomics
rimethylamine lyase
rimethylamine N-oxide

Girculating trimethylamine *N*-oxide (TMAO) is a contributor to cardiovascular disease (CVD), which in turn is the leading cause of disability and mortality worldwide.¹ TMAO enhances atherosclerosis, thrombosis, and platelet aggregation in animal models and strongly predicts incident CVD in humans.^{2–7} Understanding the determinants of plasma TMAO may lead to the discovery of novel targets for the treatment and prevention of CVD via TMAO reduction. Although fish and other foods contain TMAO, the major source of circulating TMAO is gut microbial production of the metabolic precursor trimethylamine.^{2,8} A phylogenetically diverse group of bacterial strains convert compounds abundant in beef such as carnitine to trimethylamine, which is absorbed and converted to TMAO by host enzymes.^{9,10} Suppression of bacterial trimethylamine production with antibiotics also suppresses TMAO production in humans,^{9–11} and

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For Sources of Funding and Disclosures, see page 12.

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CLINICAL PERSPECTIVE

What Is New?

- Although the presence of gut microbes is required for trimethylamine-*N*-oxide synthesis, the fecal abundance of microbial genes known to produce trimethylamine only predicted a small amount of variance in plasma trimethylamine-*N*-oxide.
- The abundance of these genes only weakly predicted the activity of the enzymes they encode.
- The fecal abundance of the cutC gene was correlated with plasma trimethylamine-*N*-oxide only in participants who consumed a non-meat diet before being exposed to a red meat diet.

What Are the Clinical Implications?

• Fecal microbiome composition cannot be substituted for trimethylamine-*N*-oxide as a biomarker for cardiovascular risk.

Nonstandard Abbreviations and Acronyms

RMSE root-mean-square-error

RPKM reads per kilobase per million reads

TMAO trimethylamine-N-oxide

trimethylamine-lyase enzyme inhibitors suppress TMAO production in mice.^{12,13} Furthermore, self-described vegans without prior meat exposure produce far less TMAO from ingested carnitine than omnivores.^{9–11,14} A small diet intervention study found that 8 weeks of a meat-free diet prevented a meat-induced increase in TMAO, but no bacterial taxa were associated with TMAO.¹⁵ Stool 16S rRNA gene sequencing has identified bacterial genera more abundant in omnivores and high TMAO producers, suggesting that consumption of animal products, and specifically red meat, raises TMAO by providing a competitive advantage to gut microbes with the capacity to use carnitine and other substrates abundant in red meat while producing trimethylamine as a waste product.^{8,11,16–18}

If red meat increases TMAO via the growth of trimethylamine-producing gut microbes, then the abundance of their genes, especially trimethylaminelyases, should predict TMAO. Previous efforts have been made to model TMAO and CVD as a function of microbial gene abundance, but these experiments were insufficient to test the hypothesis that dietinduced changes in microbial composition predict changes in TMAO. A recent multiomic study of TMAO determinants modeled human plasma TMAO as a function of stool microbial taxa and basic clinical labs, but those factors could only account for 14% of the variability in plasma TMAO.¹⁹ Importantly, this multiomic study could not quantify specific bacterial strains or genes. One cross-sectional study of atherosclerotic CVD used metagenomics sequencing to quantify 2 trimethylamine-lyases, and these genes were not strongly associated with CVD.²⁰ Both studies were observational and may have been affected by variation in dietary habits that impact both TMAO and the gut microbiota.

We present our findings from a randomized dietary study of stool microbial gene composition and its effect on TMAO in 113 individuals that used shotgun metagenomics (mean 11 million reads per sample) of stool to quantify microbial taxa and genes. We used a gene cluster-based approach to quantify low-abundance trimethylamine-lyases and expanded our analysis to include millions of microbial genes. Finally, we measured fecal trimethylaminelyase enzyme activities to better understand the relationships between gene abundance, trimethylamine production in stool, and TMAO production in the host.

METHODS

To minimize the possibility of unintentionally sharing information that can be used to re-identify private information, a subset of the data generated for this study are available at the National Center for Biotechnology Information Sequence Read Archive and can be accessed at https://www.ncbi.nlm.nih.gov/bioproject/ PRJEB44883/.

Study Participants

Healthy subjects were enrolled in the APPROACH (Animal and Plant Protein and Cardiovascular Health) study, as previously described (NCT01427855^{18,21,22}). Briefly, APPROACH assigned healthy participants to consume 4 diets prepared in a metabolic kitchen. The first diet was designed to reflect a typical American diet while the other 3 differed in protein source: red meat, white meat, or non-meat. Participants were assigned to the latter 3 diets in random order, each lasting 4 weeks and separated by a 2- to 7-week washout period during which participants returned to their habitual diets. Following each 4-week period, stool, plasma, urine, and body measurements were collected in a single clinic visit. TMAO and creatinine were quantitated in plasma and urine as previously described.²² Fractional renal excretion of TMAO was calculated as $FER_{TMAO} = 100 \times \frac{U_{TMAO}P_{creatinine}}{P_{TMAO}U_{creatinine}}$. Fecal samples were homogenized and splite into 2 parts for metagenomics sequencing and determination of trimethylamine-lyase enzyme activities. This study has been approved by the local institutional review board, and all participants provided written informed consent.

Metagenomic Sequencing

Fecal samples were prepared for shotgun metagenomic sequencing using the previously reported BGI methods workflow.²³ DNA library construction was performed following the manufacturer's instruction (Illumina Hiseq2000). One paired-end library with insert size of 350 bp for each sample was built and sequenced with 100 bp read length from each end. SolexaQA²⁴ was used to trim bases with Phred score <20 and remove reads <45 bp. SOAPAligner2 was then used to mapping highquality reads to the human genome (build hg19, Genome Reference Consortium Human Build 37), and only unmapped, high-quality reads were used for metagenomic analysis (mean 11 million reads per sample).²⁵

Metagenomic data sets are publicly available through National Center for Biotechnology Information's Sequence Read Archive Database under BioProject PRJEB44883.

Bioinformatics

Microbial genes were quantified globally with MOCAT2²⁶ using the Integrated Genome Catalog²⁷ as a reference. Gene families were quantified with Humann2.28 To quantify low-abundance trimethylamine-related genes. gene cluster reference sequences were constructed based on Kyoto Encyclopedia of Genes and Genomes (KEGG) Sequence Similarity Database annotations.²⁹ For each gene, a representative gene was chosen to query the prokaryote Sequence Similarity Database with a similarity score threshold of 100. The application program interface KEGGREST was used to download genomic coordinates for gene and gene cluster homologs. These coordinates were mapped to sequence data from KEGG GENOMES. Because of distant genes occasionally included in KEGGREST records, gene clusters >40 kb were excluded from further analysis. The implementation of this gene cluster-based quantification is available at: https://github.com/maf167/Fecal -Microbiome-Composition-Does-Not-Predict-Diet-Induced-TMAO-Production-in-Healthy-Adults.

High-quality reads were aligned to the resulting reference sequences using Bowtie2 with "verysensitive-local" parameters. Gene cluster alignments allowed multimapping, and homologs were analyzed independently. Gene alignments did not allow multimapping, and read counts were pooled among all homologs. Gene and gene cluster abundance was expressed in reads per kilobase per million reads (RPKM) units, except where indicated. A second method of normalization based on universal single copy genes yielded nearly identical results. Similar to the Metagenomic Universal Single-Copy Correction method,³⁰ universal single copy genes (USCGs) were quantified in RPKM units and used to normalize gene abundances:

Gene copy per genome = $\frac{\text{gene RPKM}}{\text{sum (USCG RPKM)}}$ = $\frac{\text{gene counts*USCG mean length}}{\text{USCG counts*gene length}}$

Determination of Fecal Enzyme Activities

All fecal samples (n=430) were weighed by subtracting the mass of a 2-mL Eppendorf tube. After weighing, 20% fecal slurries were prepared with TT media (3% Trypticase Soy Broth [Becton Dickson]; 1% Trehalose) in a Coy anaerobic chamber. Raw fecal slurries were filtered by centrifugation at 200 g for 2 minutes using 100 μ m EASYstrainers. DMSO was then added to 5% of total volume, and the complete fecal slurries were then aliquoted and stored at -80 °C for later incubation with labeled trimethylamine precursors.

Eight replicates of each fecal slurry were diluted 100× in M9 media under anaerobic conditions. Both d6-choline and d9-carnitine, which were chemically synthesized and purified as previously reported,²² were added for a final concentration of 200 μ mol/L. This culture was split in 3 parts. One part was used to measure baseline optical density, a second part was used to measure baseline cell viability with Thermo PrestoBlue, and the final part was incubated at 37 °C under anaerobic conditions for 36 hours.

At 18- and 36-hour time points, the incubated culture was vortexed and a 100 μ L volume was removed. This 100 μ L sample was quenched with excess formic acid and stored at -80 °C for later mass spectrometric analysis. All analytes were quantified with stable isotope dilution high-performance liquid chromatography with on-line electrospray ionization as previously described.²²

TMAO Modeling With Elastic Net and Random Forest

From 461 clinic visits, clinical data, gene cluster abundances (RPKM), fecal trimethylamine-lyase activities, TMAO fractional excretion rate, and plasma and urine TMAO were concatenated. For 100 participants who completed 4 weeks of the non-meat and red meat diets, the percent change from non-meat to red meat was calculated for each measurement as $\Delta x = 100 \times \frac{x_{red meat} - x_{non-meat}}{x_{non-meat}}$. Urine and plasma TMAO were modeled with clinic measurements, while Δ TMAO in urine and plasma were modeled with percent changes in measurements.

One hundred training/testing sets were selected (70/30, with replacement) using fractional stratification

(in-house *stratFrac* function) to ensure the distribution of the response variable was approximately the same in training and testing sets. Performance measures are reported across all 100 testing sets.³¹

Preprocessing was performed separately for training and testing sets. Continuous values used as predictors were scaled to represent the number of standard deviations from the mean. The R package caret was used to perform 10-fold cross validation of elastic net and obtain the trained linear model by minimizing rootmean-square error, as previously described.^{31,32} Model training is described in greater depth in Data S1. When a single predictor was used, linear regression was used to obtain the model. Root-mean-square error (RMSE) was defined as:

$$RMSE = \sqrt{\frac{(predicted - actual)^2}{n}}$$

For each elastic net run, we predicted either plasma TMAO (Figure 4), the change in plasma TMAO between non-meat and red meat diet (Δ TMAO, Figure 4), or choline consumption in a fecal culture (18-hour d6choline, Figure 6). The predictors included either the clinical measurements listed in Table 1, the homologs of a trimethylamine-related microbial gene cluster (cai, cut, tor, yea), fecal trimethylamine-lyase enzyme activities, or, as a sanity check, TMAO fractional excretion rate. Interactions and other non-linear terms were not included, as they would complicate interpretation.

To quantify non-linear associations between plasma TMAO and the predictors listed above, we implemented a random forest regressor with the R package random-Forest. The number of variables sampled at each node was tuned with caret, and each model consisted of 500 trees. TMAO estimates were obtained from 100 training/ testing sets, and variable importance was quantified as the mean cross validation residual sum of squares increase across all trees after permuting a given variable.

The implementation of this method in R is accessible at: https://github.com/maf167/Fecal-Microbiome -Composition-Does-Not-Predict-Diet-Induced-TMAO-Production-in-Healthy-Adults.

RESULTS

Diet-Induced Changes in TMAO Are Not Predicted by Trimethylamine-Lyase Abundance

We sought to measure changes in TMAO and the fecal microbiota in healthy individuals undergoing changes in diet (baseline characteristics shown in Table 1). Plasma and urine TMAO were increased significantly during red meat consumption compared with a meatfree period, and isotope tracer studies implicated

	Baseline	Non-meat	Red meat	White meat	P value
n	116	115	115	115	>.99
Sex (percent women)	60.3	61.7	60.9	60.9	>.99
Age, y	45 (33–56)	45 (33–56)	45 (33–56)	45 (33–56)	>.99
Triglycerides	71.8 (57.1–95.5)	69.2 (54.5–87.4)	67.8 (52.9–93.6)	66.2 (54.6-83.6)	0.33
TC	172 (152–196)	153 (141–177)	160 (144–184)	164 (144–183)	0.003
LDL-C	101 (83.9–122)	86.9 (73.7–106)	92.2 (80.2–111)	94.8 (77.3–114)	0.003
HDL-C	53 (45–63.5)	51 (42–59.5)	51.8 (43–61.5)	53.5 (44–60.8)	0.6
Apo A1	133 (120–145)	130 (114–140)	130 (117–144)	132 (119–140)	0.5
АроВ	73.8 (60.5–86.5)	65 (52.6–74.4)	69 (55.4–78.1)	69.5 (54.8-82)	0.022
BMI	25.4 (22.7–27.9)	25.5 (22.8–27.9)	25.6 (22.8–27.9)	25.6 (22.8–28.3)	0.99
SBP	108 (102–116)	107 (101–116)	107 (99.8–117)	107 (102–115)	0.95
DBP	69.5 (63–74)	69 (62.2–73)	68 (63–73)	68 (62–73)	0.78
Hip circumference, cm	101 (95.4–107)	101 (94–106)	100 (95.3–107)	101 (94.4–108)	0.95
Waist circumference, cm	82 (73.3–90)	81.2 (73–89.6)	81.6 (74.2-89.9)	81.8 (74.1–89.5)	0.98
WaistIC	88 (81.9–97.8)	87.2 (80.9–95.4)	87.8 (81.2–96.4)	88.6 (81.7–96.4)	0.98
Weight, Ibs	158 (140–181)	159 (139–181)	161 (140–182)	160 (139–182)	0.99
Height, cm	168 (161–177)	168 (161–177)	168 (161–177)	168 (161–177)	>.99
Diet compliance	5 (4.5–5)	5 (4.12–5)	5 (4–5)	5 (4.12–5)	>.99

 Table 1.
 Participant Characteristics at the Time of Sample Collection

Fecal and plasma samples were collected at baseline and following exposure to 3 prepared diets. Data are shown as median (interquartile range). Diet groups were compared with the Kruskal–Wallis test, except for sex (Chi-square test). BMI indicates body mass index; DBP, diastolic blood pressure; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol; SBP, systolic blood pressure; TC, Total cholesterol; and Waistlc, waist circumference measured at superior border of the iliac crest.

increased microbial production of trimethylamine from carnitine, possibly because of expansion of bacterial strains selectively using carnitine.²² We hypothesized that individual differences in TMAO production among the protein sources could be predicted by the copy number of genes related to TMAO synthesis in stool, and our analyses are summarized in Figure 1.

The Uniprot Knowledgebase lists 5 non-mammalian enzymes directly involved in trimethylamine production in the human gut (Figure 2): the choline trimethylamine-lyases cutC/D,^{12,33,34} the carnitine oxygenase cntA/B,^{16,35–37} the non-specific trimethylamine-lyase yeaW/X,³⁵ the betaine reductase component grdH,^{38–40} and the TMAO reductase torA.^{40,41} We also considered homologs of the crotonobetainyl-CoA reductase system caiA/B/C, as it catalyzes the first step in a 2-step synthesis of trimethylamine from carnitine.^{10,42–44} For 6 out of 10 of these genes, >30% of samples had no sequencing reads align to any reference sequence (Table S1). We hypothesized that these genes were not absent, but present at low abundance. To quantify low-abundance genes, we used a novel approach based on conserved gene clusters.

The 6 enzyme systems described above are coded by gene clusters conserved within phylogenetic clades. We estimated the abundance of each related gene cluster as a proxy for the gene of interest. After normalizing for reference length, homologs of all gene clusters except *cnt* and *cut* were highly correlated with gene-level quantification, and present in >70% of fecal samples (Figure S1). The *cntA* gene has been estimated to have <30% prevalence in humans,⁴⁵ while the *cut* gene cluster is phylogenetically diverse and not well conserved.³³ Gene cluster-based quantifications allowed us to test whether each homolog of each gene predicts diet-induced changes in TMAO.

Homologs of the *cai* gene cluster were weakly correlated with plasma TMAO (Table 2, Table S2). The *cai* gene cluster was also among the most important predictors of plasma TMAO in random forest models (Figure S4).

A recent diet intervention study observed that participants who consumed a meat-free diet first did not have increased plasma TMAO after switching to a red meat diet, while those assigned to the red meat diet first had increased TMAO that decreased after switching to a meat-free diet.¹⁵ This diet order effect could not be explained by 16S rRNA taxonomic profiling, so we examined correlations between trimethylamine-related genes and plasma TMAO in participants assigned to non-meat before red meat and vice versa. The fecal abundance of *cutC* was weakly correlated with plasma TMAO in those assigned to non-meat before red meat (rho=0.21, P=0.002, Figure 3). The same diet order effect was observed with an alternative normalization method based on universal single copy genes (Tables S3 through S5).

To assess whether any genes directly related to trimethylamine metabolism predict TMAO production, we trained an elastic net linear model with 70% of available samples using 10-fold cross validation to minimize RMSE. We then used the resulting linear model to predict TMAO concentrations in the held-out samples and used Pearson correlation coefficient to estimate the portion of variance in TMAO predicted by the model. Finally, we repeated training and testing on 100 randomly selected held-out groups. Using clinical data summarized in Table 1, the model accounted for 41% of the variance in plasma TMAO (Figure 4). We used TMAO fractional excretion rate as a sanity check to confirm that a valid predictor of plasma TMAO has the expected effects on model performance metrics (increased adjusted R^2 and decreased RMSE). We also implemented a random forest to predict TMAO with non-linear associations. The percent change in the fractional excretion rate of TMAO from a non-meat to a red meat diet improved prediction of the percent change in urine TMAO by 12%, as predicted by the findings of Wang, et al.²² However, trimethylaminelyase gene abundances did not predict plasma or urine TMAO nor did they improve the linear models or nonlinear random forest models based on clinical data (Figure 4 and Figure S3).

TMAO is Not Predicted by Functional Metagenomic Analysis

While the abundance of candidate, trimethylaminerelated genes did not predict TMAO, there are many undescribed microbial genes that may. Diet-related changes in microbial taxa have been reported in this cohort using 16S rRNA gene sequencing,¹⁸ and we used metagenomic data to quantify taxa, genes, and gene families. No single taxonomic unit or gene was directly correlated with plasma or urine TMAO. Three Uniref gene families were weakly correlated with plasma TMAO with borderline significance (Table 3). These correlations were also significant after multiple testing adjustment with the method of Benjamini and Hochberg.⁴⁶ We next sought to understand why TMAO is not more strongly predicted by fecal microbiota composition.

Trimethylamine-Lyase Gene Abundance Does Not Predict Direct Measurements of Enzyme Activity in Fecal Samples

TMAO is a product of both commensal and host metabolism, while trimethylamine is produced only by gut microbes.² Microbial gene abundance may not be an effective predictor of microbial trimethylamine production, and trimethylamine production may not be a strong predictor of TMAO. We first compared trimethylamine and TMAO production in 13 individuals.

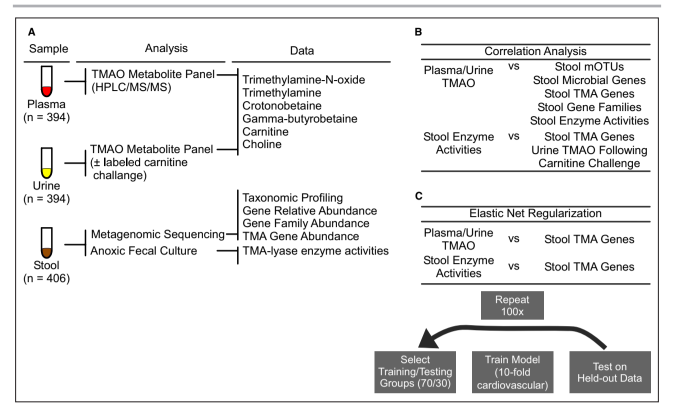


Figure 1. Overview of analyses.

A, Healthy adults were provided non-meat, white meat, or red meat diets for 4-week periods and donated plasma, urine, and stool samples during clinic visits. Trimethylamine *N*-oxide (TMAO) and related metabolites were measured in plasma and urine samples. A subset of urine samples (n=38) was taken after oral challenge with isotope labeled choline and carnitine. Isotope-labeled TMAO was measured in these urine samples. Stool was used for both anoxic cultures to determine the activities of enzymes related to trimethylamine production and metagenomics sequencing. Metagenomic sequence data were used to compute metagenomic operational taxonomic units, as well as global gene and gene family abundances. Low abundance trimethylamine-related genes were quantified using a gene cluster-base d approach. **B**, Plasma and urine TMAO were not correlated with any stool data set, and stool trimethylamine -related enzyme activities were not correlated genes or with TMAO production during the oral challenge. **C**, Elastic net models trained with trimethylamine-related gene abundances did not predict TMAO or stool enzyme activities. CV indicates cardiovascular; FMO, flavin mono-oxygenases; GBB, gamma-butyrobetaine; HPLC/MS/MS, high performance liquid chromatography with tandem mass spectrometry; MI, myocardial infarction; mOTUs, metagenomic operational taxonomic units; and TMAO, trimethylamine-*N*-oxide.

Isotope tracer studies involving a subset of the cohort (n=38) reported increased conversion of oral carnitine to urine trimethylamine and TMAO during a red meat diet versus white meat and non-meat, consistent with unlabeled TMAO.²² We anoxically cultured fecal samples from the same clinic visits in the presence of isotopically labeled carnitine and choline and measured labeled trimethylamine and synthetic intermediates after 18- and 36-hour incubations. These measures allowed us compare trimethylamine production by gut microbes to TMAO production in the microbe-host system.

Although there were no strong correlations between trimethylamine production in feces and TMAO production in urine, choline metabolite concentrations were moderately correlated in urine and feces (rho=-0.46, P=0.003, Figure 5). This correlation was also significant

after multiple testing adjustment (adjusted P=0.04) with the method of Benjamini and Hochberg.⁴⁶ In the prediction of d6-choline consumption, elastic net linear models trained with *cutC* abundances outperformed those trained with clinical data (Table 1), however no model of fecal enzyme activity predicted >6 percent of the variation in labeled metabolites (Figure 6).

Production of trimethylamine from carnitine in feces was not correlated with TMAO production from carnitine in the host (Spearman correlation, P=0.5), and carnitine trimethylamine-lyase gene abundances did not predict carnitine metabolism in feces (Figure S2). In short, abundances of genes known to encode enzymes that produce trimethylamine in feces fail to predict systemic TMAO because they cannot predict total fecal enzyme activity, at least under the culture conditions described in Methods.

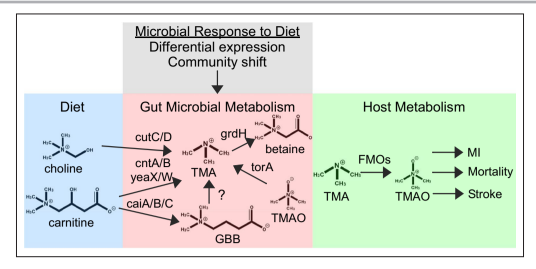


Figure 2. Synthesis of trimethylamine *N*-oxide requires dietary substrate as well as host and microbial enzymes.

Nutrients including choline and carnitine, which are abundant in red meat, are converted to trimethylamine via multiple synthetic pathways. The copy number of related genes is altered by diet and may contribute to trimethylamine *N*-oxide production by host enzymes. caiA/B/C, crotonobetainyl-CoA dehydrogenase; cntA/B, carnitine oxygenase/reductase; cutC/D, choline utilization gene cluster; FMOs, flavin monooxygenases; GBB, gamma-butyrobetaine; grdH, betaine reductase complex component B subunit beta; MI, myocardial infarction; TMA, trimethylamine; TMAO, trimethylamine-*N*-oxide; torA, trimethylamine-N-oxide reductase (cytochrome c); and yeaX/W, carnitine monooxygenase subunit YeaX.

Table 2. Abundance of Gene Cluster cai is Correlated With Plasma TMAO

Gene Cluster	Organism	ρ	P value*
cai	Escherichia coli O18:K1:H7	0.17	0.040
cai	Salmonella enterica subsp. enterica serovar Bareilly	0.16	0.069
cai	Citrobacter werkmanii	0.16	0.070
cai	Salmonella enterica subsp. enterica serovar Enteritidis EC20090332	0.16	0.073
cai	Salmonella enterica subsp. enterica serovar Enteritidis EC20090135	0.16	0.11

Four candidate genes were quantified using Kyoto Encyclopedia of Genes and Genomes Sequence Similarity Database gene clusters. Of 274 gene cluster homologs, 60 were detected in >70% of samples. Spearman correlations were calculated between these 60 homolog abundances and plasma TMAO. The top 5 *cai* homologs are shown, and all data are shown in Table S2. TMAO indicates trimethylamine-*N*-oxide.

*Adjusted according to the method of Bonferroni.

DISCUSSION

The gut microbial metabolite TMAO is a major contributor to cardiovascular disease, and intervention may be complicated by the complex interactions that result in elevated circulating TMAO.¹⁹ Plasma TMAO can be raised and lowered with dietary change or gut microbial suppression.^{12,22} In addition, TMAO response to these interventions is modulated by pre-existing vegan diet and renal function. A vegan diet, or more specifically a non-meat diet, has been observed to cause changes in both stool microbiome composition and TMAO metabolism.^{9,14} The abundance of carnitine in red meat presumably gives an advantage to gut microbial strains that consume carnitine, thus increasing production of trimethylamine and TMAO. However, TMA-related gene abundance did not predict systemic TMAO, possibly because of limitations in metagenomics technology or a large impact of host factors.

As interventions to lower TMAO are considered, it will be important to understand the relationships among diet, gut microbiota composition, and TMAO. We hypothesized that if the growth of specific strains plays a role in TMAO metabolism, then abundance of the relevant taxa or genes would predict TMAO levels. We used paired fecal, plasma, and urine samples from healthy participants assigned to diets with and without meat in a random crossover design to test predictive models of TMAO based on the abundance of genes related to trimethylamine production. We used shotgun metagenomic sequencing to quantify gene abundance in feces and measured TMAO in plasma and urine.

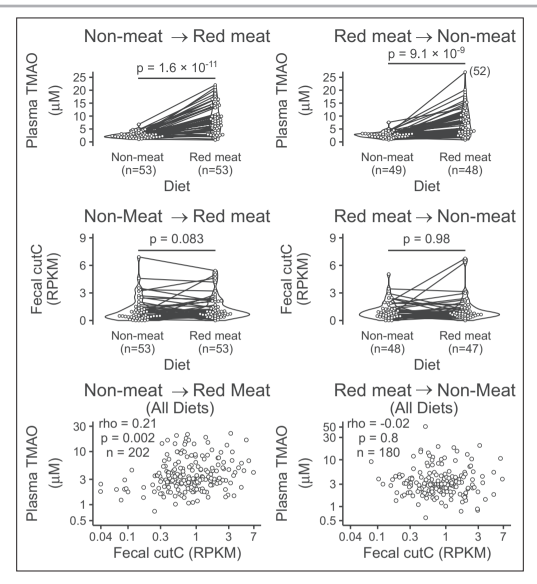


Figure 3. *cutC* predicts trimethylamine *N*-oxide in those assigned to non-meat before red meat. Plasma trimethylamine *N*-oxide is higher during red meat consumption vs a meat-free diet regardless of diet order. Although *cutC* is not differentially abundant in red meat vs meat-free, *cutC* abundance is correlated with plasma trimethylamine *N*-oxide among those assigned to non-meat before red meat. *P* values represent Wilcox tests. RPKM indicates reads per kilobase per million reads; TMAO, trimethylamine-*N*-oxide.

We first tested whether the abundance of 5 genes known to be involved in trimethylamine production could predict diet-induced changes in TMAO. Although the *cai* gene cluster and *cutC* gene were correlated with TMAO, (Table 2, Figure 3) none improved machine learning models of plasma TMAO over clinical data alone (Figure 4). The *cutC* gene was correlated with plasma TMAO only in those assigned to the non-meat diet before red meat (Figure 3), but the observed effect of diet order on cutC might be diminished by the study design. The APPROACH study was designed to measure independent effects of multiple diet interventions in the same individuals. Participants were instructed to return to their habitual diets for at least 2 weeks between experimental diets to "wash out" any transient effects of the previous diet. Minimizing the effect of diet order was intended to be an advantage, but given the diet order effects on *cutC*, the washout periods may be a limitation by diminishing the strength of relationships that require a non-meat diet before a red meat diet or vice versa.

Expanding our analysis to the entire metagenome did not reveal any associations with plasma or urine TMAO. No gene, taxonomic group, or gene family met the Bonferroni significance threshold for correlation with TMAO. Three gene families met a less stringent

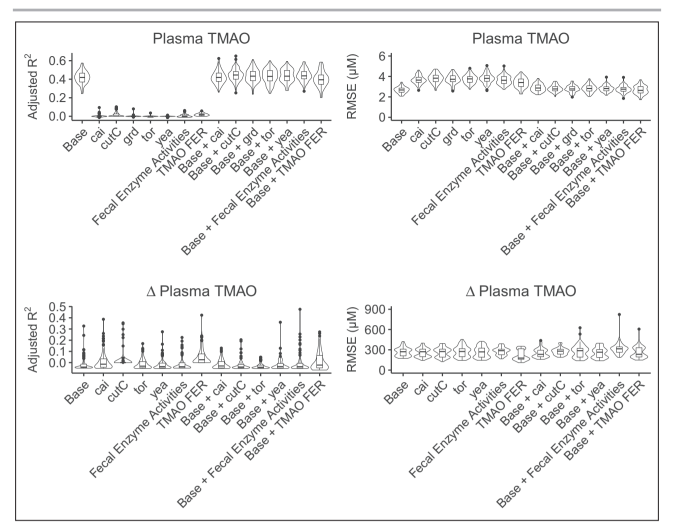


Figure 4. Abundance of genes related to trimethylamine synthesis do not improve models of trimethylamine *N*-oxide (TMAO).

Elastic net was used to train and test linear models of plasma TMAO, as well as percent changes in TMAO from non-meat to red meat diets. Models were tested with 100 randomly selected training/testing groups, using adjusted R^2 and root-mean-squared error as performance metrics. The base model, using data summarized in Table 1, predicted 41% of the variance in TMAO, and the addition of trimethylamine-related gene abundances did not improve model performance. TMAO fractional excretion rate outperformed the base model in predicting changes in plasma TMAO, in terms of root-mean-squared error (P<0.001, Wilcox test). FER, fractional excretion rate; RMSE indicates root-mean-squared error; and TMAO, trimethylamine-N-oxide.

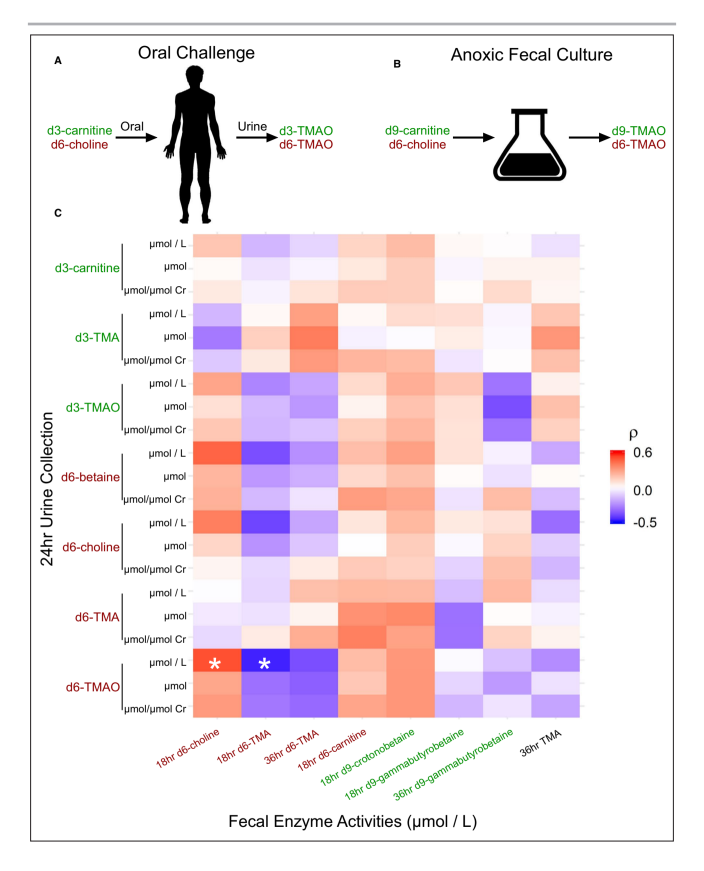
significance threshold for correlation (Tables 2 and 3). The metagenome of human feces did not predict systemic TMAO levels (Figure 4) or in vitro trimethylamine production (Figure S2), and this lack of association may be because of the limitations of metagenomic functional profiling.

Table 3.	Spearman	Correlation	Between	Plasma	TMAO	and	Gene	Families
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UniRef90 ID	Name	ρ	P value*
A6BD41	Uncharacterized protein	0.22	0.047
F0R5N1	SF3 helicase domain-containing protein	-0.22	0.047
F7JHG5	Uncharacterized protein	0.22	0.047
C0BEB7	Uncharacterized protein	0.22	0.050
G2SXI9	Phosphomethylpyrimidine synthase	0.22	0.050

Gene abundances were aggregated into Uniref90 gene family abundances using Humann2. Of 28 441 gene families detected in >75% of samples, 5 were weakly correlated with plasma trimethylamine-*N*-oxide. TMAO indicates trimethylamine-*N*-oxide.

*Adjusted according to the method of Benjamani and Hochberg.46



Functional metagenomic analysis aims to quantify the metabolic potential of a microbial community given genomic information.⁴⁷ The presence of genes coding for an enzyme is one of many requirements for enzyme activity. Enzymatic reactions require transcription and translation of the gene, sufficient substrate, and a

Figure 5. Fecal trimethylamine-lyase activities are not correlated with trimethylamine *N*-oxide (TMAO) production in host. In a single visit, 13 participants randomly selected from the APPROACH cohort donated a fecal sample and participated in a heavyisotope-labeled choline and carnitine challenge (38 visits total). A, Participants were dosed orally with 250mg d6-choline and d3carnitine. Labeled TMAO and synthetic intermediates were quantitated in 24-hour urine collection. B, Paired fecal samples were cultured with 200 µmol/L d6-choline and d9-carnitine and labeled TMAO and synthetic intermediates were quantitated after 18- and 36-hour incubations. Analytes not detected in >30% of samples were excluded. C, The heatmap shows Spearman correlations between fecal enzyme activities and the indicated isotopologue in 24-hour urine collection. TMAO produced from carnitine in the host was not highly correlated with carnitine-derived trimethylamine in fecal cultures. Urine choline-derived TMAO was negatively correlated with choline-trimethylamine-lyase activity in feces (ρ =-0.46, *P*=0.003). Cr, creatinine; RMSE indicates root-mean-squared error; and TMAO, trimethylamine-*N*-oxide. * Adjusted *P*<0.05. *P* values adjusted with the method of Benjamani and Hochberg.⁴⁶

microenvironment conducive to maintaining the structure of the enzyme and the health of the cell.⁴⁸

Multiomic analyses, in which the metagenome, metabolome, etc are surveyed in the same samples, elucidate the extent to which the metagenome predicts metabolic activity and should create an expectation that TMAO is not predicted by the metagenome. Comparisons of fecal gene and transcript abundances have estimated the metagenome accounts for less than half the variation in the meta-transcriptome.⁴⁹ Strong associations between gene abundances and metabolites should not be expected when the intermediate processes (ie, transcription) are so loosely connected to gene copy number.

Although the presence or absence of 50% of metabolites detected by liquid chromatography mass spectrometry can be predicted with metagenomic data, <10% of the variation in metabolite levels is accounted for by DNA-level data.^{49,50} Metabolite production is predicted to be low when required genes are not detected, but we have demonstrated that aligning

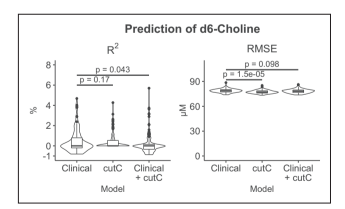


Figure 6. *cutC* abundance marginally outperforms clinical data in predicting fecal choline-trimethylamine lyase activity.

Fecal samples were cultured with 200 μ mol/L d6-choline and labeled trimethylamine and synthetic intermediates were quantitated after 18-hour incubation. Elastic net models were trained to predict 18-hour d6-choline with 70% of samples and tested on the remaining 30%, and testing was iterated over 100 different training/testing sets. Although the *cutC* models had less error than models trained with clinical data (Table 1), *cutC* models did not result in higher correlations between predicted and actual d6-choline concentration. RMSE indicates rootmean-squared error. reads to reference gene sequences often fails to detect low-abundance genes whose conserved gene clusters are present (Table S1). Quantitative polymerase chain reaction has been used to measure fecal *cutC* and *grdH* as low as 0.05% of bacterial genomes.⁴⁵

The trimethylamine/TMAO synthetic pathway has many of the features that complicate prediction of activity levels with meta-genomic data. Even with simulated sequence data, analysis of low-abundance strains is challenging, and bacterial strains at <0.4% abundance in the mouse gut can have an outsized influence on trimethylamine and TMAO production.34,51 The regulation of transcription can also limit the prediction of metabolites from gene abundance data, and in fact, the transcription of at least 1 trimethylaminelyase, *cutC/D*, is regulated by substrate availability.³³ Regulation via translation rate, post-translational modifications, and cofactor availability may also impact enzyme activity independent of gene copy number. Trimethylamine-producing microbes active in the gut may be inactive in stool or vice versa. Finally, circulating trimethylamine and TMAO concentrations are impacted by host liver enzymes and renal filtration as well as microbial activity.6,22,52

Our isotope tracer studies in humans and fecal cultures challenged with labeled carnitine revealed discrepancies in the metabolism of microbes and the host-microbe system (Figure 4). This metabolic discrepancy is consistent with at least one report of transcriptionally inactive strains in human stool.53 A lack of transcription would also account for the observed inability of trimethylamine-lyase gene abundance to predict enzyme activity in stool (Figure 4). Finally, tools like MelonPann that use metagenomic data to predict the presence or absence of metabolites would have no translational application to TMAO, because low, detectable plasma or urine TMAO does not impact cardiovascular risk.50,54,55 Metabolomic analysis of metagenomes using flux balance analysis and optimization predicts many microbial metabolites well, but it assumes a relationship between gene abundance and metabolite concentration and requires high-quality annotated genomes of relevant microbes.⁵⁶ While the quantitative prediction of nearly any metabolite with metagenomics data are challenging, TMAO is an especially difficult case because of the impact of host metabolism and the limitations of metagenomic analysis.

While microbial trimethylamine production is required for TMAO synthesis, red-meat-induced TMAO production is not strongly predicted by fecal microbial composition or the abundance of genes related to trimethylamine production.

ARTICLE INFORMATION

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Supplementary Material

Data S1 Tables S1–S5 Figures S1–S4

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SUPPLEMENTAL MATERIAL

Data S1.

Supplemental Methods

Induced TMAO Production in Healthy Adults

Elastic Net Modeling

The goals of the present analyses were first, to test whether plasma TMAO is associated with the abundance of specific strains of bacteria in the fecal microbiome and second, identify microbial enzymes or strains whose associations with TMAO are comparable to more easily measured predictors such as sex and diet. We were especially focused on identifying simple linear relationships amenable to future experiments in model systems. Finally, we expected a high degree of collinearity when comparing closely related strains and gene homologs.

Elastic net accomplishes both goals by "selecting" a relatively small number of predictors that minimize model error and reporting an effect size (β coefficient) for each predictor that can be directly compared.

For each elastic net run, we predicted either plasma TMAO (Figure 4), the change in plasma TMAO between non-meat and red meat diet (Δ TMAO, Figure 4), or choline consumption in a fecal culture (18-hour d6-choline, Figure 6). The predictors included either the clinical measurements listed in Table 1, the homologs of a TMA-related microbial gene cluster (cai, cut, tor, yea), fecal TMA-lyase enzyme activities, or, as a sanity check, TMAO fractional excretion rate (see Response #2). Interactions and other nonlinear terms were not included, as they would make interpretation more difficult.

Each outcome-predictor set pair was analyzed as follows. The pseudocode below summarizes the elastic net analysis.

main

- 1 Define a predictor set P = [...] and outcome \overrightarrow{O}
- 2 Define training sets $T_{train} = \{T_1, T_2, \dots, T_{100}\}$ and T_{test} where $T_i = \{p_i \subset P, \vec{o_i} \subset \vec{O}\}$
- 3 for each training set T do
- 4 Train linear model
- 5 Predict testing set T_{test_i}
- 6 Compute adjusted R² and RMSE
- 7 **end**

Train

- 1 Define a grid of alpha and lambda values
- 2 for each $\{\alpha, \lambda\}$:
- 3 Randomly divide training data into ten folds
- 4 *for each* fold **do**
- 5 | Hold out the nth fold
- 6 Preprocess the remaining nine folds
- 7 Fit linear model with preprocessed data
- 8 Predict data in the nth fold
- 9 **end**
- 10 Compute average RMSE across 10 folds
- 11 end
- 12 Select the $\{\alpha, \lambda\}$ set with smallest RMSE
- 13 Fit final model using all training data and optimal $\{\alpha, \lambda\}$

From the set of 461 clinical visits, those visits with complete data were randomly divided into training (70%) and testing (30%) sets. Due to the wide variation in model performance, we repeated training/testing assignment with replacement to produce a total of 100 training/testing sets. Within each training/testing set, model error and variance explained were quantified as follows.

Elastic net models were trained and tested in R 3.5.0 by invoking the glmnet function via the train function of the caret package. Briefly, model parameters are tuned by minimizing a cost function for each of combination of 9 out of 10 folds within the training set, and the root-mean-square error (RMSE) is determined for the prediction of the tenth, held-out fold. The optimal parameters producing the lowest RMSE are used to produce a model using the entire training set, and this model is then used to predict the held-out testing set. The error and variance explained by the model in the testing set is then computed by comparing predicted to measured values.

A grid of λ and α values based on preliminary testing was defined for $\lambda = \{0, 0.1, 0.2 \dots 12\}$ and $\alpha = \{0, 0.1, 0.2 \dots 1\}$. These two parameters were optimized, or tuned, by quantifying prediction error for all 1200 combinations of λ and α values using 10-fold cross-validation.

For cross-validation, the training set was divided into ten folds, and each fold was iteratively held out. The remaining nine folds are preprocessed independently of the held out fold: continuous predictors are scaled and centered, categorical predictors are recoded as "dummy variables", and predictors with missing or zero data in >30% samples are excluded.

Within each cross-validation, $\hat{\beta}$ coefficients were computed to minimize the cost function *C* using the given values of λ and α .

$$C(\hat{\beta},\lambda,\alpha) = \frac{1}{2n} \sum_{i} (\hat{y}_i - y_i)^2 + \lambda \left(\frac{1-\alpha}{2} \sum \hat{\beta}^2 + \alpha \sum |\hat{\beta}|\right)$$

Each 10-fold cross validation produced 10 models tested on held out folds to compute error as root-mean-square error (RMSE), a value used when predicting data with a skewed distribution, such as TMAO.

$$RMSE = \sqrt{\frac{1}{n} \sum_{i} (\hat{y}_i - y_i)^2}$$

Nearly all the measured values in our manuscript appear to follow a right-skewed distribution. We wanted to give more weight to larger absolute errors to construct a more clinically useful model. Larger absolute errors will hamper the clinical goal of risk stratification more than smaller absolute errors.

The values of λ and α with the least average RMSE were deemed optimal, and the final model was trained as above with all training data using the optimal values of λ and α .

The resulting linear model was then used to predict the data in the held-out testing data. These predicted values are compared to actual values using RMSE and adjusted R-squared. Predictor sets that produce higher median adjusted R-squared and/or lower median RMSE across 100 testing sets are said to improve the model. The Wilcox test is used to assess a difference in medians.

Table S1. Proportion of samples in which gene or gene cluster is detected. Metagenomic sequencing reads from 405 fecal samples were aligned to genes related to TMA production as well as the associated gene cluster. In almost every case, conserved gene clusters are more frequently detected than the genes of interest. P values represent Bonferroni-adjusted Chi-square tests comparing each gene to the related gene cluster.

Gene Cluster	Gene	Detection Rate (%)	P (Chi-Square)
	Gene		i (cili-oquale)
cai		99.8 (404/405)	
	caiA	76.8 (311/405)	< 0.0001
	caiB	94.8 (384/405)	0.0004
	caiC	93.6 (379/405)	< 0.0001
cnt		13.1 (53/405)	
	cntAB	12.8 (52/405)	>0.99
cut		97.3 (394/405)	
	cutC	94.8 (384/405)	>0.99
	cutD	38.8 (157/405)	< 0.0001
grd		97.8 (396/405)	
	grdH	94.3 (382/405)	0.19
tor		95.3 (386/405)	
	torA	64.9 (263/405)	< 0.0001
yea		95.1 (385/405)	
	yeaW	65.2 (264/405)	< 0.0001
	yeaX 4	5.2 (183/405)	< 0.0001

Table S2. Universal single copy genes used for alternative normalization. Manor and Banorstein (1) selected prokaryotic orthologous groups with exactly one copy in almost every prokaryotic genome listed by KEGG. We used only the top ten universal single copy genes to quantify relative bacterial genome count in fecal samples as an alternative form of normalization.

KEGG Orthology	COG	Annotation	Prokaryotic Genomes With Gene (%)	Prokaryotic Genomes With >1 Copy (%)
K02988	COG0098	small subunit ribosomal protein S5	6112 (99)	5 (0.08)
K02994	COG0096	small subunit ribosomal protein S8	6111 (99)	5 (0.08)
K02933	COG0097	large subunit ribosomal protein L6	6108 (99)	8 (0.13)
K02931	COG0094	large subunit ribosomal protein L5	6097 (99)	13 (0.21)
K02863	COG0081	large subunit ribosomal protein L1	6096 (99)	5 (0.08)
K02876	COG0200	large subunit ribosomal protein L15	6094 (99)	8 (0.13)
K02967	COG0052	small subunit ribosomal protein S2	6093 (98)	7 (0.11)
K02864	COG0244	large subunit ribosomal protein L10	6081 (98)	8 (0.13)
K02881	COG0256	large subunit ribosomal protein L18	6076 (98)	6 (0.10)
K03110	COG0552	fused signal recognition particle receptor	5961 (96)	8 (0.13)

Table S3. Spearman correlations with plasma TMAO among all samples. Four candidate genes were quantified using KEGG SSDB gene clusters. Of 274 gene cluster homologs, 60 were detected in >70% of samples. Spearman correlations were calculated between these 60 homolog abundances and plasma TMAO. Gene clusters are named with KEGG org identifier and genomic coordinates. Universal single-copy gene normalization (USCG) utilized ten USCGs.

	RPKM				USCG			
Gene Cluster	Rho	Р	P-Bonferroni	Rho	Р	P-Bonferroni		
cai-eih-27336-52938	0.17	0.00065	0.04	0.17 0.0)0086	0.053		
cai-senv-57098-100887	0.16 0.	.0012	0.073	0.16	0.0016	0.099		
cai-sena-57098-100887	0.16 0.	.0018	0.11	0.16	0.002	0.12		
cai-seeb-3884824-3928761	0.16 0.	.0011	0.069	0.17 0.0)0094	0.057		
cai-cwe-3656832-3683530	0.16 0.	.0011	0.07	0.17 0.0)0083	0.051		
cai-ecv-27317-52919	0.15 0.	.0026	0.16	0.15	0.0032	0.19		
cai-sek-57350-101254	0.15 0.	.0028	0.17	0.15	0.0035	0.21		
cai-senq-57098-100887	0.15 0.	.0033	0.2	0.15	0.0036	0.22		
cai-sel-57340-101063	0.15 0.	.0029	0.18	0.15	0.0029	0.18		
cai-senc-1519486-1563447	0.15 0.	.0029	0.18	0.15	0.0034	0.21		
cai-ses-2844264-2873996 0.1	14 0.0043	3	0.26	0.14	0.004	0.24		
cai-ebt-3409854-3439396 0.1	14	0.005	0.31	0.14	0.0041	0.25		
cai-edh-3821075-3845551 0.	13	0.013	0.77	0.12	0.016	0.98		
cai-ecz-27336-52937	0.13 0	.0074	0.45	0.14	0.0055	0.34		
cai-setc-3466245-3510218	0.12	0.02	1	0.11	0.023	1		
cai-cko-3102867-3134413 0.1	12	0.018	1	0.12	0.018	1		
cai-ecl-3947888-3972396	0.11	0.022	1	0.12	0.02	1		
cai-eci-27336-52938	0.11	0.034	1	0.11	0.036	1		
cai-sea-57341-101275	0.11	0.029	1	0.11	0.025	1		
cai-senj-2127653-2171775	0.11	0.023	1	0.12	0.019	1		
cutC-Pooled	0.11	0.029	1	0.11	0.023	1		
cai-ecoi-27459-53061	0.1	0.042	1	0.1	0.041	1		
cai-seep-57183-100834	0.1	0.039	1	0.1	0.038	1		
cai-senl-57098-100886	0.1	0.042	1	0.1	0.044	1		
cai-seno-57098-100887	0.097 0	0.055	1	0.095	0.058	1		
cai-sens-57342-100598	0.097 0	0.054	1	0.097	0.054	1		
grd-cace-803252-808152 0.0	96 0.056		1	0.1	0.048	1		
grd-shi-2784756-2816008	0.091 0	0.071	1	0.09	0.072	1		
tor-eck-1158853-1178787	0.089 0).078	1	0.091	0.069	1		

cai-elu-4788091-4813693	0.088 0.0	082	1	0.087	0.083	1
cai-sene-57344-101308	0.087 0.	085	1	0.088	0.079	1
cai-eal-1458476-1483446	0.079	0.12	1	0.083	0.1	1
grd-cfm-2173856-2178773	0.077	0.12	1	0.081	0.11	1
cai-sega-57339-101062	0.073	0.15	1	0.069	0.17	1
cai-cama-23975-56364	0.069	0.17	1	0.07	0.16	1
cai-cyo-4142522-4169176	0.066	0.19	1	0.071	0.16	1
cai-ema-2472536-2497872	0.064	0.2	1	0.065	0.19	1
cai-eco-25826-50302	0.063	0.21	1	0.065	0.2	1
cai-sec-51598-95536	0.062	0.22	1	0.069	0.17	1
cai-eko-4058406-4082885	0.055	0.28	1	0.056	0.27	1
cai-seec-1052456-1096396	0.053	0.29	1	0.053	0.29	1
cai-ena-14772-39563	0.051	0.31	1	0.053	0.29	1
cai-sew-57348-101465	0.047	0.36	1	0.048	0.34	1
cai-ebc-294299-320969	0.045	0.37	1	0.05	0.32	1
cai-spq-51689-95628	0.039	0.44	1	0.04	0.43	1
cai-cif-1056571-1081433 0.	037	0.46	1	0.041	0.42	1
cai-sed-57342-96817	0.033	0.51	1	0.034	0.5	1
cai-ebf-3985021-4010024	0.033	0.51	1	0.039	0.44	1
cai-sbg-52520-83258	0.028	0.58	1	0.031	0.53	1
yea-eco-1878874-1894731	0.027	0.59	1	0.03	0.55	1
cai-cro-32133-66456	0.021	0.68	1	0.024	0.64	1
cai-sbv-49201-79934	0.019	0.71	1	0.021	0.67	1
cai-cfar-891731-923473 0.0	0064	0.9	1	0.007	0.89	1
cai-sbz-49701-81116	0.0061	0.9	1	0.011	0.83	1
cai-cbra-2695495-2727410	0.0022 0	.96	1	0.0038	0.94	1
cai-seo-57344-102030	-0.0039 0	.94	1	-0.003	0.95	1
cai-efe-21905-59475	-0.014	0.78	1	-0.011	0.83	1
cai-kin-3809521-3836197 -0	0.03	0.55	1	-0.028	0.58	1
cai-cir-287785-319507	-0.056	0.27	1	-0.054	0.28	1
cai-caf-4639362-4680627	-0.062	0.22	1	-0.056	0.27	1
grd-cdrk-2091059-2111627	-0.066	0.19	1	-0.071	0.16	1

Table S4. Spearman correlations with plasma TMAO among those asigned to nonmeat before red meat. Four candidate genes were quantified using KEGG SSDB gene clusters. Of 274 gene cluster homologs, 60 were detected in >7% of samples. Spearman correlations were calculated between these 60 homolog abundances and plasma TMAO. Gene clusters are named with KEGG org identifier and genomic co ordinates. Universal single-copy gene normalization (USCG) utilized ten USCGs.

	RPKM			USCG			
Gene Cluster	Rho	Р	P-Bonferroni	Rho	Р	P-Bonferroni	
cai-ebt-3409854-3439396	0.23	0.00092	0.056	0.220.	0017	0.1	
cutC-Pooled	0.22	0.0016	0.1	0.210.	0019	0.12	
cai-eih-27336-52938	0.21	0.0019	0.11	0.2	0.0037	0.23	
cai-cwe-3656832-3683530	0.21	0.0029	0.18	0.2	0.0035	0.22	
cai-ecl-3947888-3972396	0.2	0.0036	0.22	0.190.	0057	0.35	
cai-sel-57340-101063	0.19	0.0059	0.36	0.190.	0071	0.43	
cai-ecz-27336-52937	0.18	0.0098	0.6	0.180.	0088	0.54	
cai-sena-57098-100887	0.17	0.012	0.73	0.17	0.015	0.91	
cai-ecv-27317-52919	0.16	0.019	1	0.15	0.03	1	
cai-edh-3821075-3845551	0.15	0.026	1	0.14	0.04	1	
cai-seeb-3884824-3928761	0.15	0.028	1	0.14	0.037	1	
cai-senc-1519486-1563447	0.15	0.034	1	0.14	0.046	1	
cai-senv-57098-100887	0.14	0.045	1	0.13	0.063	1	
cai-ses-2844264-2873996	0.14	0.047	1	0.13	0.059	1	
cai-eci-27336-52938	0.13	0.058	1	0.13	0.069	1	
cai-senj-2127653-2171775	0.13	0.064	1	0.12	0.083	1	
cai-sbv-49201-79934	0.13	0.068	1	0.12	0.077	1	
cai-ebc-294299-320969	0.13	0.063	1	0.12	0.079	1	
cai-setc-3466245-3510218	0.12	0.077	1	0.11	0.11	1	
cai-elu-4788091-4813693	0.11	0.11	1	0.1	0.15	1	
cai-senq-57098-100887	0.11	0.1	1	0.11	0.12	1	
cai-sens-57342-100598	0.1	0.15	1	0.094 ().18	1	
grd-shi-2784756-2816008	0.1	0.13	1	0.099 (0.16	1	
cai-cama-23975-56364	0.098	0.16	1	0.093 ().18	1	
cai-sek-57350-101254	0.093	0.18	1	0.086 ().22	1	
cai-eko-4058406-4082885	0.092	0.18	1	0.085 ().22	1	
cai-sea-57341-101275	0.085	0.22	1	0.079 ().26	1	
cai-sbz-49701-81116	0.082	0.24	1	0.079 ().26	1	
cai-sene-57344-101308	0.08	0.25	1	0.072	0.3	1	

g	rd-cfm-2173856-2178773 0	.078	0.26	1	0.072	0.3	1
	cai-sbg-52520-83258	0.077	0.27	1	0.07	0.32	1
	cai-seo-57344-102030	0.073	0.29	1	0.063 0.	37	1
С	ai-cyo-4142522-4169176	0.073	0.29	1	0.061 0.3	38	1
ç	rd-cace-803252-808152	0.067	0.33	1	0.057 0.4	41	1
	cai-ecoi-27459-53061	0.065	0.35	1	0.061 0.3	38	1
Ca	ai-seec-1052456-1096396	0.059	0.39	1	0.059	0.4	1
С	ai-eal-1458476-1483446	0.057	0.41	1	0.044 0.	53	1
	cai-seno-57098-100887	0.057	0.41	1	0.055 0.4	43	1
	cai-seep-57183-100834	0.053	0.44	1	0.05	0.47	1
	cai-senl-57098-100886	0.052	0.46	1	0.044 0.	52	1
C	cai-cif-1056571-1081433	0.044	0.53	1	0.038 0.	59	1
	cai-ena-14772-39563	0.032	0.65	1	0.03	0.67	1
	cai-sega-57339-101062	0.029	0.68	1	0.016 0.	82	1
С	ai-cko-3102867-3134413	0.029	0.68	1	0.023 0.	74	1
to	or-eck-1158853-1178787	0.024	0.73	1	0.024 0.	73	1
	cai-cfar-891731-923473	0.022	0.75	1	0.015 0.	83	1
	cai-sec-51598-95536	0.017	0.8	1	0.017	0.8	1
	cai-spq-51689-95628	0.013	0.85	1	0.0057	0.94	1
ye	ea-eco-1878874-1894731	0.0076	0.91	1	0.007 0.	92	1
Ca	ai-ema-2472536-2497872	0.00064	0.99	1	-0.0014	0.98	1
	cai-sew-57348-101465	-0.0013	0.99	1	-0.0066	0.92	1
	cai-cir-287785-319507	-0.0036	0.96	1	-0.011	0.88	1
	cai-eco-25826-50302	-0.0045	0.95	1	-0.0075	0.91	1
	cai-cro-32133-66456	-0.011	0.88	1	-0.02	0.78	1
	cai-efe-21905-59475	-0.018	0.79	1	-0.022	0.76	1
Ca	ai-cbra-2695495-2727410 -(0.02	0.78	1	-0.032	0.64	1
С	ai-ebf-3985021-4010024 -0	.034	0.62	1	-0.038	0.59	1
С	ai-caf-4639362-4680627 -0	.036	0.61	1	-0.037	0.6	1
	cai-sed-57342-96817	-0.045	0.52	1	-0.056	0.42	1
gr	d-cdrk-2091059-2111627	-0.076	0.27	1	-0.097	0.16	1
C	ai-kin-3809521-3836197	-0.08	0.25	1	-0.09	0.2	1

Table S5. Spearman correlations with plasma TMAO among those assigned to red meat before non-meat. Four candidate genes were quantified using KEGG SSDB gene clusters. Of 274 gene cluster homologs, 60 were detected in >7% of samples. Spearman correlations were calculated between these 60 homolog abundances and plasma TMAO. Gene clusters are named with KEGG org identifier and genomic co ordinates. Universal single-copy gene normalization (USCG) utilized ten USCGs.

	RPKM			USCG			
Gene Cluster	Rho	Р	P-Bonferroni	Rho	Р	P-Bonferroni	
cai-cko-3102867-3134413	0.24	0.0011	0.064	0.24	0.00089	0.055	
cai-sek-57350-101254	0.23	0.0016	0.1	0.23	0.0018	0.11	
cai-senq-57098-100887	0.19	0.0089	0.54	0.2	0.0064	0.39	
cai-senv-57098-100887	0.18	0.012	0.72	0.18	0.012	0.72	
cai-seeb-3884824-3928761	0.18	0.011	0.7	0.2	0.0058	0.36	
cai-seep-57183-100834	0.17	0.021	1	0.18	0.014	0.83	
tor-eck-1158853-1178787	0.16	0.027	1	0.17	0.02	1	
cai-ecoi-27459-53061	0.15	0.039	1	0.16	0.033	1	
cai-senl-57098-100886	0.15	0.038	1	0.16	0.025	1	
cai-ses-2844264-2873996	0.15	0.046	1	0.15	0.036	1	
cai-eco-25826-50302	0.14	0.053	1	0.15	0.042	1	
cai-sea-57341-101275	0.14	0.06	1	0.15	0.042	1	
cai-senc-1519486-1563447	0.14	0.049	1	0.15	0.037	1	
cai-eih-27336-52938	0.13	0.087	1	0.13	0.067	1	
cai-ecv-27317-52919	0.13	0.067	1	0.14	0.053	1	
cai-ema-2472536-2497872	0.13	0.085	1	0.13	0.071	1	
cai-seno-57098-100887	0.13	0.076	1	0.13	0.069	1	
cai-sena-57098-100887	0.13	0.083	1	0.13	0.067	1	
cai-sed-57342-96817	0.13	0.082	1	0.14	0.054	1	
cai-sega-57339-101062	0.12	0.11	1	0.12	0.096	1	
cai-sec-51598-95536	0.12	0.12	1	0.13	0.075	1	
cai-ebf-3985021-4010024	0.12	0.1	1	0.13	0.067	1	
grd-cace-803252-808152 0.	12	0.11	1	0.14	0.06	1	
cai-eal-1458476-1483446	0.11	0.14	1	0.13	0.071	1	
cai-setc-3466245-3510218	0.11	0.13	1	0.12	0.094	1	
cai-cwe-3656832-3683530	0.11	0.14	1	0.13	0.082	1	
cai-edh-3821075-3845551 0	.1	0.17	1	0.11	0.14	1	
cai-sens-57342-100598	0.1	0.16	1	0.11	0.15	1	
cai-senj-2127653-2171775	0.099	0.18	1	0.11	0.12	1	

cai-sel-57340-101063	0.098	0.18	1	0.1	0.16	1
cai-sene-57344-101308 0.0	098	0.18	1	0.12	0.11	1
cai-sew-57348-101465	0.095	0.19	1	0.1	0.15	1
cai-ecz-27336-52937	0.088	0.23	1	0.098	0.18	1
cai-ena-14772-39563	0.077	0.3	1	0.085	0.25	1
grd-cfm-2173856-2178773	0.076	0.3	1	0.091	0.22	1
cai-eci-27336-52938	0.074	0.32	1	0.08	0.28	1
grd-shi-2784756-2816008	0.074	0.31	1	0.082	0.26	1
cai-spq-51689-95628	0.067	0.36	1	0.076	0.3	1
cai-elu-4788091-4813693	0.064	0.38	1	0.072	0.33	1
cai-cro-32133-66456	0.061	0.41	1	0.079	0.29	1
cai-cyo-4142522-4169176	0.057	0.44	1	0.083	0.26	1
yea-eco-1878874-1894731	0.057	0.44	1	0.067	0.36	1
cai-seec-1052456-1096396	0.042	0.57	1	0.053	0.47	1
cai-kin-3809521-3836197	0.04	0.59	1	0.051	0.49	1
cai-cama-23975-56364	0.038	0.61	1	0.046	0.53	1
cai-ebt-3409854-3439396	0.038	0.61	1	0.054	0.46	1
cai-cbra-2695495-2727410	0.034	0.64	1	0.047	0.53	1
cai-efe-21905-59475	0.022	0.77	1	0.034	0.65	1
cai-ecl-3947888-3972396	0.02	0.79	1	0.033	0.65	1
cai-cif-1056571-1081433 0.	02	0.78	1	0.038	0.61	1
cai-eko-4058406-4082885	0.011	0.88	1	0.02	0.79	1
cai-cfar-891731-923473	-0.015	0.83	1	-0.0066	0.93	1
cutC-Pooled	-0.018	0.81	1	-0.0023	0.97	1
cai-sbg-52520-83258	-0.03 0.	68	1	-0.013	0.86	1
grd-cdrk-2091059-2111627 -	0.041	0.58	1	-0.032	0.66	1
cai-ebc-294299-320969	-0.051	0.49	1	-0.036	0.63	1
cai-sbz-49701-81116	-0.084	0.25	1	-0.065	0.37	1
cai-seo-57344-102030	-0.09 0.	22	1	-0.075	0.31	1
cai-caf-4639362-4680627	-0.091	0.22	1	-0.073	0.32	1
cai-sbv-49201-79934	-0.12	0.1	1	-0.11	0.15	1
cai-cir-287785-319507	-0.12 0.	11	1	-0.1	0.16	1

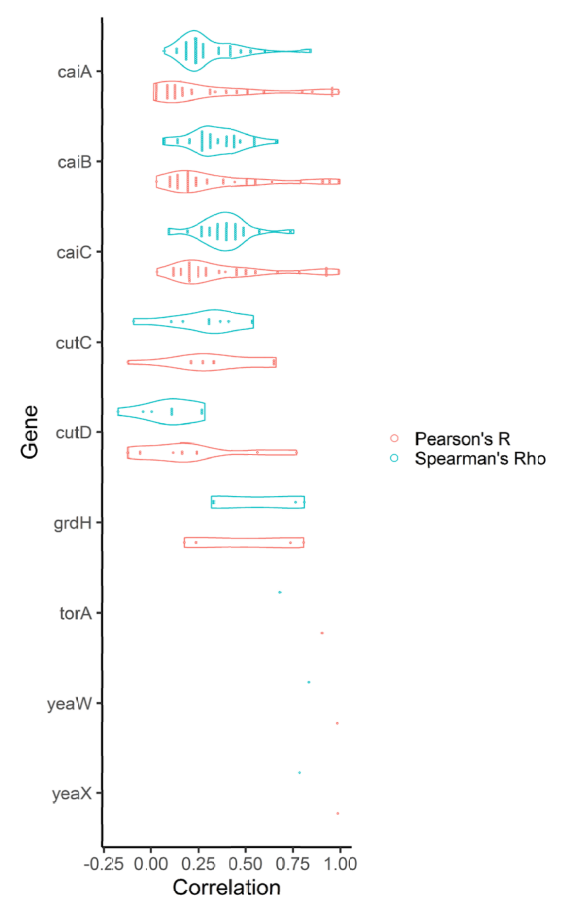
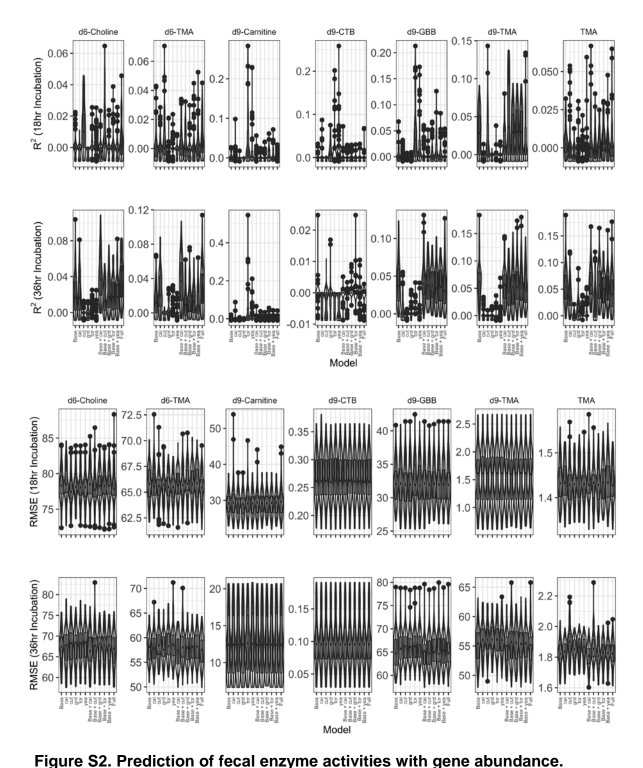
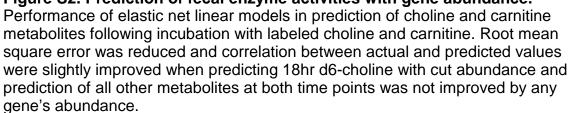


Figure S1. Gene cluster abundances are correlated with gene abundances. Genes known to be related to TMAO synthesis were quantified as RPKM according to the count of reads aligning to any gene homolog. Gene cluster homologs present in >70% of samples were quantified individually, and multiple homologs of each gene cluster were highly correlated with and detectable in more fecal samples than gene-level quantification.





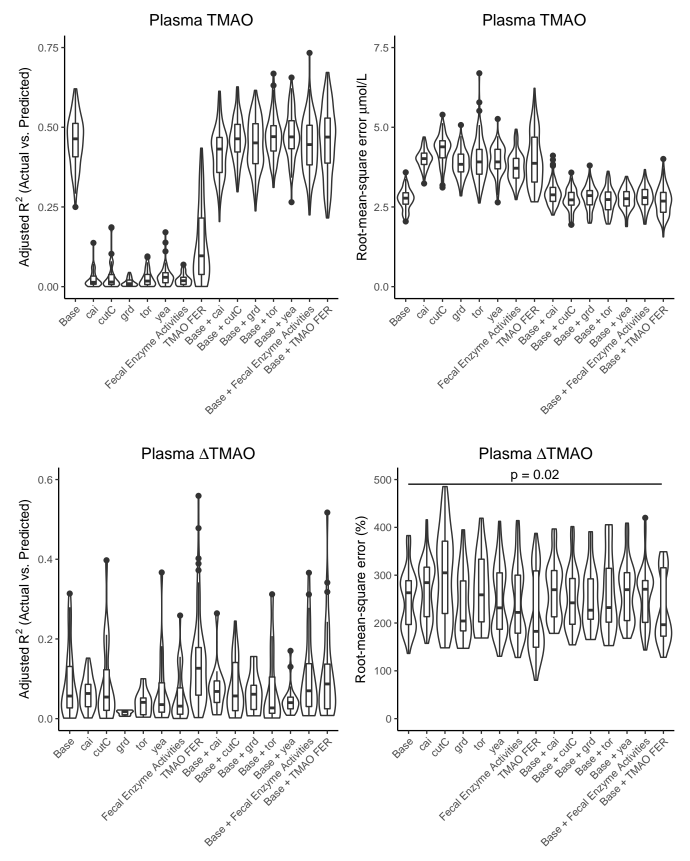
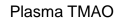


Figure S3. Abundance of genes related to TMA synthesis do not improve random forest models of TMAO. Random forest was used to train and test models of plasma TMAO, as well as percent changes in TMAO from non-meat to red meat diets (Δ TMAO). Models were tested with 100 randomly selected training/testing groups, using adjusted R² and root-mean-squared error (RMSE) as performance metrics. The base model, using data summarized in Table 1, predicted 46% of the variance in TMAO, and the addition of TMA-related gene abundances did not improve model performance. As a sanity check, TMAO fractional excretion rate (FER) outperformed the base model in predicting changes in plasma Δ TMAO, in terms of RMSE (p = 0.02, Wilcox test).



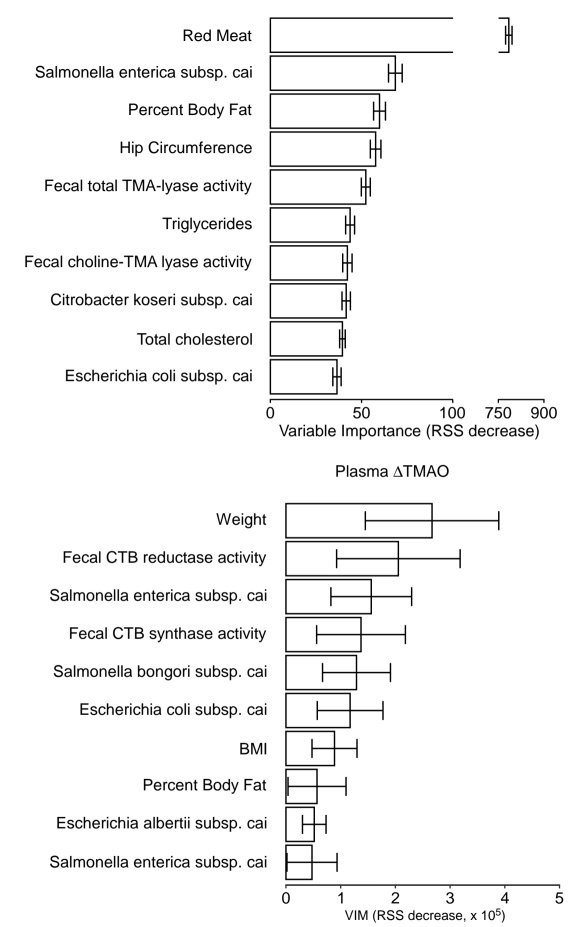


Figure S4. cai gene cluster ranks among the most important predictors of plasma TMAO. Random forest was used to train and test models of plasma TMAO, as well as percent changes in TMAO from non-meat to red meat diets (Δ TMAO). Models were tested with 100 randomly selected training/testing groups. During training, the cross-validation error (residual sum of squares, RSS) was computed after permuting each predictor. Variable importance is reported as the increase in RSS after permutation. Bars represent mean and standard error.