

RESEARCH ARTICLE

Open Access

Diet and exercise orthogonally alter the gut microbiome and reveal independent associations with anxiety and cognition

Silvia S Kang¹, Patricio R Jeraldo², Aishe Kurti¹, Margret E Berg Miller^{3,4}, Marc D Cook^{3,4}, Keith Whitlock^{3,4}, Nigel Goldenfeld^{5,6}, Jeffrey A Woods^{3,4}, Bryan A White⁵, Nicholas Chia^{2*} and John D Fryer^{1,7,8*}

Abstract

Background: The ingestion of a high-fat diet (HFD) and the resulting obese state can exert a multitude of stressors on the individual including anxiety and cognitive dysfunction. Though many studies have shown that exercise can alleviate the negative consequences of a HFD using metabolic readouts such as insulin and glucose, a paucity of well-controlled rodent studies have been published on HFD and exercise interactions with regard to behavioral outcomes. This is a critical issue since some individuals assume that HFD-induced behavioral problems such as anxiety and cognitive dysfunction can simply be exercised away. To investigate this, we analyzed mice fed a normal diet (ND), ND with exercise, HFD diet, or HFD with exercise.

Results: We found that mice on a HFD had robust anxiety phenotypes but this was not rescued by exercise. Conversely, exercise increased cognitive abilities but this was not impacted by the HFD. Given the importance of the gut microbiome in shaping the host state, we used 16S rRNA hypervariable tag sequencing to profile our cohorts and found that HFD massively reshaped the gut microbial community in agreement with numerous published studies. However, exercise alone also caused massive shifts in the gut microbiome at nearly the same magnitude as diet but these changes were surprisingly orthogonal. Additionally, specific bacterial abundances were directly proportional to measures of anxiety or cognition.

Conclusions: Thus, behavioral domains and the gut microbiome are both impacted by diet and exercise but in unrelated ways. These data have important implications for obesity research aimed at modifications of the gut microbiome and suggest that specific gut microbes could be used as a biomarker for anxiety or cognition or perhaps even targeted for therapy.

Keywords: Neuroscience, Gut-brain axis, Microbiome, Anxiety, Cognition, Exercise, Diet

Background

Obesity is a rising public health concern, especially in the United States where over a third of the population is categorized as obese [1]. It is a pervasive co-morbidity and is associated with a variety of diseases such as hypertension, stroke, coronary artery disease, diabetes, cancer, age-associated dementias, depression, and anxiety [2,3]. Obesity in children is associated with poorer academic

achievement [4] and adult obesity is correlated with poorer performance on cognitive tasks [5-7]. Obese children and adults are more likely to have lower core self-evaluation and suffer from anxiety and depression [8]. Exercise is one proven approach to combat obesity in both humans and rodent models and likely operates via many potential mechanisms [9]. While many studies have demonstrated the positive effects of exercise on cognitive abilities or the anxiogenic effects of a high fat diet (HFD), virtually no studies have examined the intersection of these manipulations and whether exercise can normalize or prevent abnormalities of cognition or anxiety in a well-controlled animal model. This

* Correspondence: chia.nicholas@mayo.edu; fryer.john@mayo.edu

²Department of Surgical Research, Mayo Clinic, 200 First St SW, Rochester, MN 55905, USA

¹Department of Neuroscience, Mayo Clinic, 4500 San Pablo Rd, Jacksonville, FL 32224, USA

Full list of author information is available at the end of the article

is of critical importance since some individuals assume that exercise can reverse all negative effects of a HFD including behavioral problems such as anxiety or impaired cognition.

We now know that obesity involves a number of factors and one of much recent interest has been the role of the gut microbiota in both weight gain and severe weight loss [10-13]. HFD is known to induce changes in the commensal gut bacterial community and seminal studies by the Gordon lab demonstrated that the gut microbiota are not merely reflective of the dietary intake, but they are actually key mediators of the metabolic state [11]. For example, fecal transfers from mice fed a HFD into naïve germ-free recipient mice led to increased adiposity in the recipients even when fed a normal diet (ND) [10]. However, very little is known about whether or how exercise affects the gut microbiome and whether alterations in the gut microbiome relate to changes in behavior. Interestingly, germ-free mice are less anxious than conventionally raised mice and re-colonization with a normal gut microbiota early in life is anxiogenic [14,15]. In this study we sought to determine the interaction of HFD and exercise on behavioral outcomes as well as the gut microbial community structure and define their relationships in adult mice. We hypothesized that HFD would cause behavioral problems and alter the gut microbiome but that the effects on both of these would be mitigated by exercise. Much to our surprise, we found that both HFD and exercise substantially altered behavior and the gut microbial community but exercise did not normalize or “rescue” the effects of a HFD and, in fact, the relationships were orthogonal.

Results

High fat diet causes anxiety that is not attenuated by exercise

To test the relationship between diet and exercise, we randomly assigned 8-week-old male C57BL/6 J mice to four groups: ND, ND + exercise, HFD, and HFD + exercise ($n = 10/\text{group}$). We chose a forced exercise paradigm so that we could precisely control the amount or “dose” of exercise. Importantly, to control for environmental enrichment and handling, non-exercised mice were placed along side their exercising counterparts in running wheels that rotated at a speed that just prevented them from sleeping (~ 1 m/min). As expected, HFD alone caused a significant increase in body weight compared to ND as early as 4 weeks after beginning treatment (Figure 1A). Exercised mice of both groups had significantly lower body weight than their sedentary counterparts beginning at 5 weeks of treatment for ND groups and 8 weeks of treatment for the HFD groups (Figure 1A). This amount of exercise was not able to completely counteract the effects of the HFD,

but this was not surprising given the extremely high fat content of this diet (60% kcal from fat) and the limited amount of exercise from these one hour sessions.

We performed behavioral testing of these groups to determine the impact of HFD or exercise on anxiety and cognition. Open field analysis did not demonstrate any significant differences in total activity or rearing among the groups (Additional file 1: Figure S1). We also found no differences between the groups in sociability as measured in the three-chamber social interaction test (Additional file 1: Figure S1). Using the light/dark exploration apparatus as a measure of anxiety, we found that mice fed a HFD were significantly more anxious as measured by percentage of time spent in the light compartment and in distance traveled while occupying the light compartment (Figure 1B and C). However, exercise was not able to counteract these anxiogenic effects of the HFD (Figure 1B and C).

Exercise improves memory but is not impacted by high fat diet

We next measured learning and memory in the contextual fear conditioning paradigm where mice are placed in a chamber and a loud tone is paired with a negative stimulus (foot shock). Mice that remember the environment will spend more time freezing when placed again in the same apparatus (contextual memory). Exercised mice on a normal diet had significantly increased contextual memory compared to their sedentary counterparts (Figure 1D). Exercised mice on a HFD diet had a trend toward increased contextual memory compared to their sedentary HFD counterparts, but this effect did not quite achieve statistical significance (Figure 1D). Similar results were obtained with the cued portion of the task (Additional file 1: Figure S1). However, HFD alone did not substantially alter either contextual or cued memory nor did it significantly reduce the positive effects of exercise.

Effect of diet and exercise on the gut microbiome

We also assessed whether exercise alone or in the context of a HFD could alter the gut microbial community. We purified DNA from fecal samples from each of the mice and profiled their gut microbiome using next generation barcoded sequencing of amplicons from the variable regions V3-V5 of the bacterial 16S ribosomal DNA (a “fingerprint” of different bacterial taxa). A total of 35,197,991 reads were obtained from 40 samples (averaging $879,950 \pm 249,310$ reads; minimum 431,900 reads; maximum 1,416,925 reads) using a 300-cycle kit on a single Illumina MiSeq sequencing run. Paired reads were then analyzed using an extension of the TORNADO pipeline for taxa, operational taxonomic unit (OTU), and phylogeny [16]. QIIME [17,18] was used to calculate

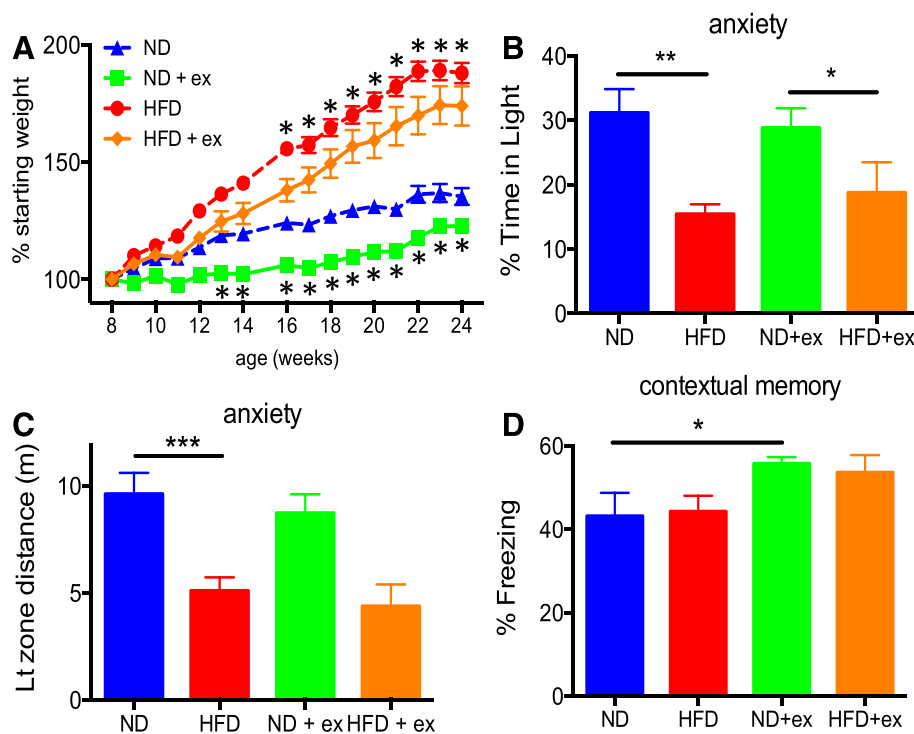


Figure 1 Effects of diet and exercise on body weight, anxiety, and cognitive behavior. (A) HFD-fed mice were significantly heavier starting at 4 weeks of treatment (ND vs HFD) while exercised mice weighed significantly less at 5 weeks of treatment for ND groups (ND vs ND + exercise, * $p < 0.05$) and at 8 weeks of treatment for HFD groups (HFD vs HFD + exercise, * $p < 0.05$). HFD-fed mice were more anxious as measured in the Light/Dark exploration assay for both percentage of time spent in the lit compartment (B) and distance traveled while in the lit compartment (C). Exercised mice had enhanced learning and memory measured in the contextual fear conditioning assay (D). Body weight analyzed by repeated measures ANOVA with post-hoc t-test and significance defined as $p < 0.05$. Behavioral data was analyzed by two-way ANOVA with post-hoc Fisher's LSD with significance indicated as * $p < 0.05$, ** $p < 0.01$, and *** $p < 0.001$. All data presented as mean \pm SEM from $n=10$ /group.

beta-diversity, which was then visualized using R with the vegan package for ordination plots.

Our initial hypothesis was that exercise would rescue many of the shifts in the gut microbiome caused by a HFD. While we found a few examples where exercise modulated HFD-induced changes in the microbiome, including a remarkable “bloom” of a *Streptococcus* genus (OTU115, family Streptococcaceae, order Lactobacillales) that was returned to undetectable levels with exercise (Figure 2A), the majority of microbiome alterations exerted by diet and exercise followed an unexpected pattern. Not only did exercise alone cause large shifts in the two most abundant phyla, Firmicutes and Bacteroidetes, but the directionality and magnitude of the effect was similar to the changes caused by HFD (Figure 2B,C). Both diet and exercise also independently decreased the relative abundance of Tenericutes (Figure 2D).

While the changes in phyla induced by HFD or exercise appear similar, this broad grouping of bacteria does not provide the resolution for determining the effects of diet or exercise on the community structure. In order to refine the analysis, we analyzed these data at the OTU

level with each OTU representing bacteria grouped at 97% identical sequences from the 16S sequencing (see Methods). Metric Dimensional Analysis (MDA) is one such way to visualize similarities or dissimilarities in large data sets while preserving their distance relationships. MDA revealed that a high percentage of variation is explained by just the first and second principal components with each of the four groups of mice separating rather cleanly into four distinct clusters (Figure 3). This indicates that most diet- or exercise-induced shifts in microbial populations are completely unrelated (i.e. orthogonal). More detailed examination of what drives this separation in ordination space revealed multiple strong signatures separating ND from HFD groups and secondarily separating sedentary from exercised groups (Figure 4 with OTU probabilities shown in Additional file 2: Table S2). Additionally, we analyzed different levels of taxonomy and found significant main effects of diet on the abundance of many different families and an equal number of main effects of exercise (Table 1). Several main effects of diet or exercise were also seen at higher levels of taxonomy such as order, class, and phylum (Additional file 3: Table S1).

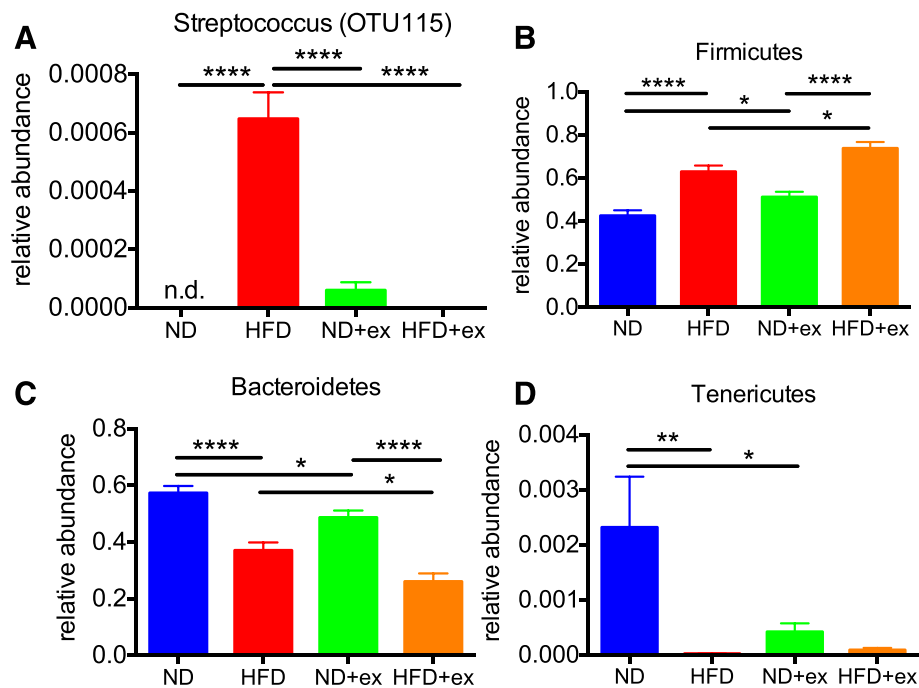


Figure 2 Changes in the relative abundances of gut bacteria by HFD or exercise. (A) HFD caused a bloom of OTU115 from the genus *Streptococcus* that returned to nearly undetectable levels in HFD + exercise mice. Bacteria of phyla Firmicutes (B), Bacteroidetes (C), and Tenericutes (D) of the gut microbiome were significantly altered by both HFD and exercise. Data was analyzed by two-way ANOVA with post-hoc Holm-Sidak t-tests with significance indicated as * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, and **** $p < 0.0001$. All data presented as mean \pm SEM from $n=10$ /group.

Associations between specific taxa with anxiety and cognition

We next wanted to test whether specific taxa would correlate with behavioral measures. We analyzed the top 100 OTU's and found that the relative abundance of OTU69, OTU90, and OTU97, all from the family Lachnospiraceae, positively correlated with % time in light, that is higher levels correlated with less anxiety (Figure 5). Other

specific bacteria such as OTU17 (family Lachnospiraceae), OTU30 (family Streptococcaceae, genus *Lactococcus*), OTU72 (family Ruminococcaceae), and OTU89 (family Ruminococcaceae, genus *Butyrivibrio*) negatively correlated with % time in light, that is lower levels correlated with less anxiety (Figure 5). In fact, anxiety relationships were found at all levels of taxonomy including family, order, class, and even phylum (Additional file 4: Figure S2), although with these sample sizes they did not reach statistical significance when corrected for false discovery rate. We also found weaker associations with cognitive performance between bacteria from the order Clostridiales such as OTU39, 82, 79, and 57 (Figure 6), although these also did not quite reach significance when corrected for false discovery rate. Nevertheless, it is interesting that they all fell within a single order (Clostridiales). In agreement with numerous other studies, several highly significant relationships with body weight were seen at all levels of taxonomy including the phyla Bacteroidetes, Firmicutes, and Proteobacteria (Additional file 5: Table S3).

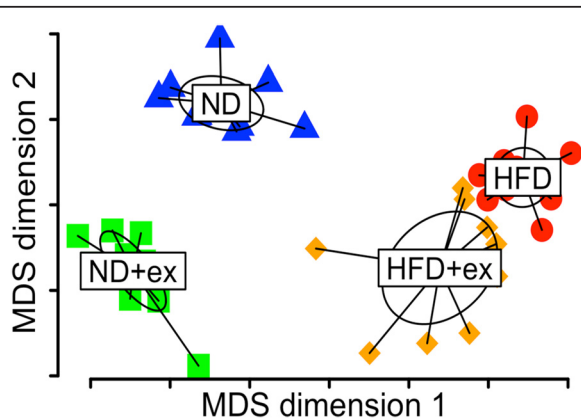
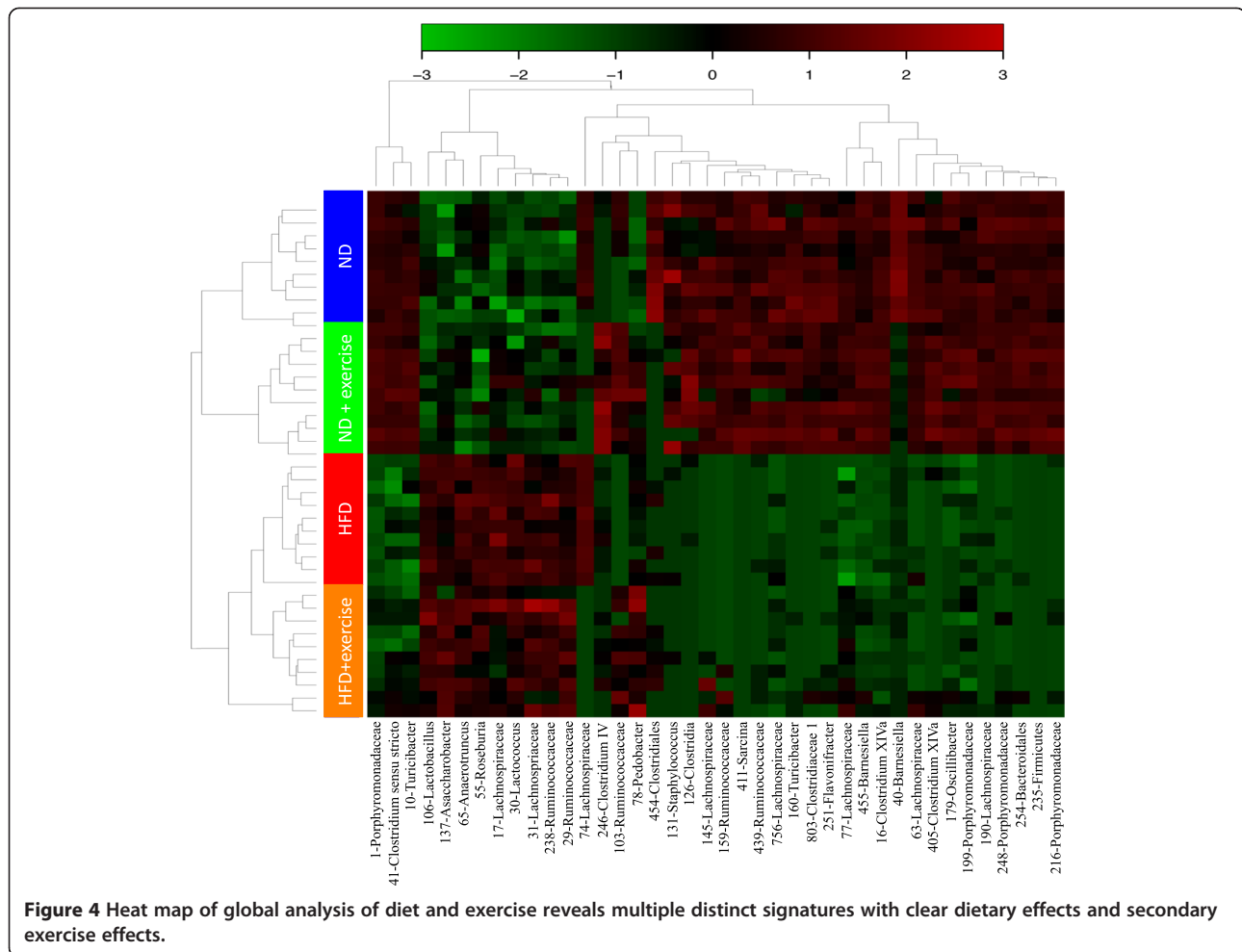


Figure 3 Multidimensional analysis of diet and exercise reveals orthogonal changes in the gut microbiome. This analysis in multidimensional space demonstrates clear segregation of each of the four groups of mice with no overlap between groups.

Discussion

Very few studies have examined how exercise alone impacts the gut microbiome. Voluntary exercise in rats



caused global changes in the gut microbiome but the technique used in those studies (PCR banding patterns) is only useful in determining whether differences exist and cannot be used to ascertain which bacteria are altered [19,20]. One study found that voluntary exercise in mice caused large alterations in the gut microbiome and with clear separations in ordination space [21], but the techniques used (PhyloChip arrays) make it difficult or impossible to compare to our data given the differences in accurately assigning bacteria to different taxa. Interestingly, one study that employed deep sequencing techniques failed to find significant changes in the gut microbiome [22], but they used voluntary wheel running and their first sampling of the microbiome was after 1 year of exercise compared to our changes found at 16 weeks. A recent study of similar design and duration to ours also found that diet and exercise exerted orthogonal effects on the gut microbiota [23]. However, they reported that exercise increased Bacteroidetes and decreased Firmicutes (opposite of our results) perhaps due to their voluntary wheel running paradigm [23].

Due to the nature of studying the interaction of exercise and diet, which requires subjects to be fed a high fat diet as one of the conditions, it is unlikely that similar studies will ever be conducted in humans. Therefore, it is important to understand how these factors interact in a well-controlled animal model system in order to determine how these parameters may impact human health. Although numerous studies have shown the benefits of exercise on cognitive function [7,24,25] as we have demonstrated here, few studies have actually examined the interaction between HFD and exercise, specifically in regard to behavioral outcomes. Rat studies have suggested that exercise can partially rescue the cognitive decline associated with HFD; however, an important parameter of behavior, anxiety, was not measured in these studies [26,27]. Our current study clearly indicates that exercise alone is unable to counteract all of the effects of a HFD. This is a critical finding given that some humans likely operate under the assumption that exercise can completely reverse the negative impact of HFD and change how we view weight loss that is conducted through exercise alone.

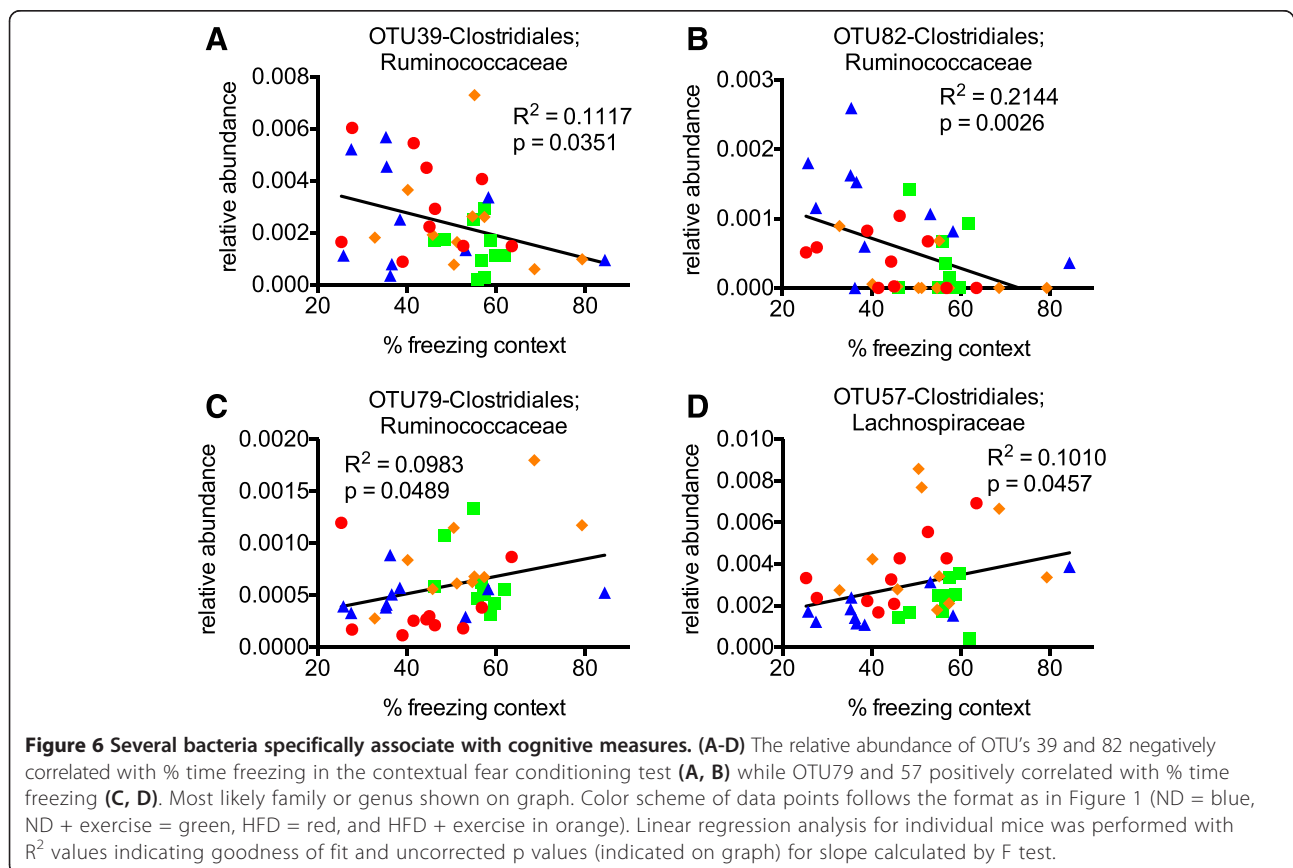
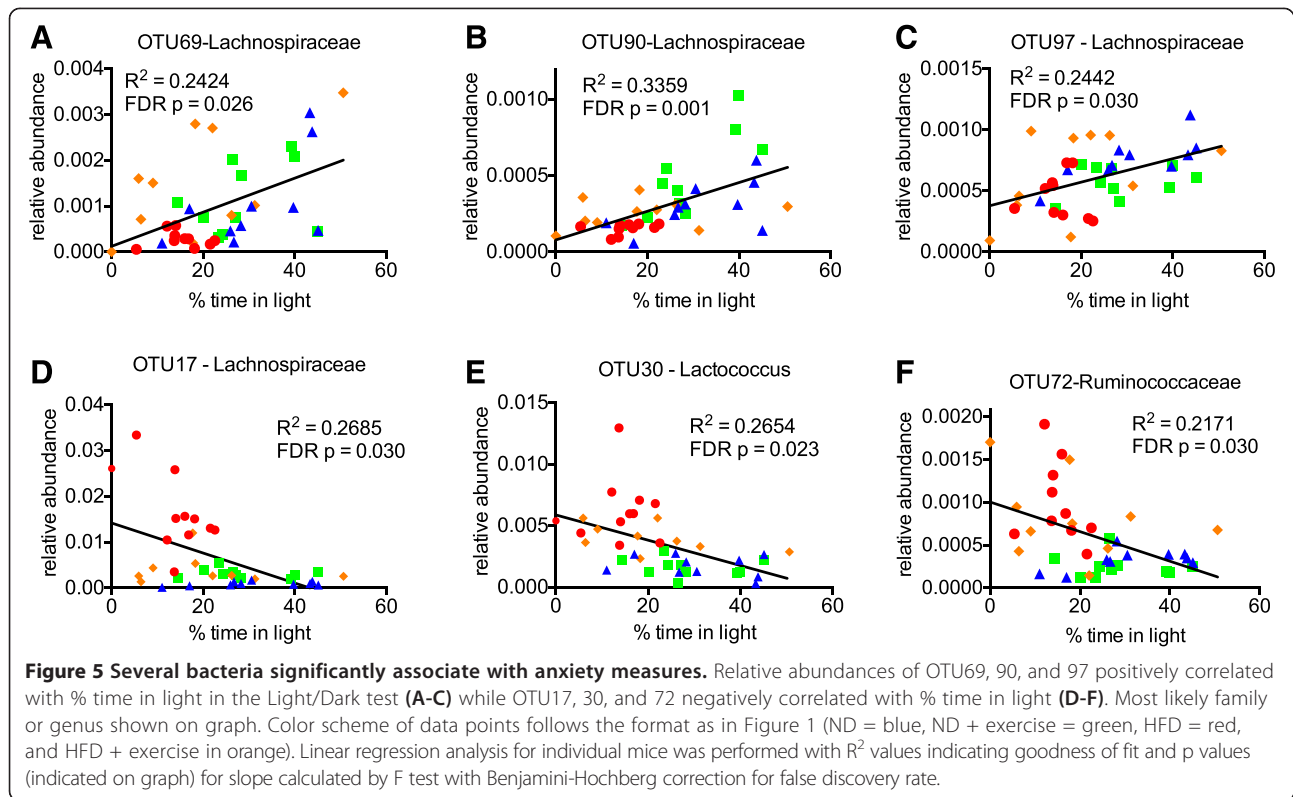
Table 1 Significant effects of diet and exercise at the level of family

| Family | Main effects 2 way ANOVA | | | Post-hoc testing | | | |
|-----------------------|--------------------------|----------|-------------|------------------|---------------------|---------------|-----------------|
| | High fat diet | Exercise | Interaction | ND vs HFD | ND + ex vs HFD + ex | ND vs ND + ex | HFD vs HFD + ex |
| Porphyromonadaceae | ↓ **** | ↓ *** | | **** | **** | * | * |
| Streptococcaceae | ↑ **** | ↓ ** | * | **** | ** | | ** |
| Lachnospiraceae | ↑ **** | ↑ * | | **** | **** | | |
| Erysipelotrichaceae | ↓ **** | | | ** | *** | | |
| Clostridiaceae_1 | ↑ **** | | | ** | **** | * | |
| Ruminococcaceae | ↑ **** | | | | *** | | |
| Peptostreptococcaceae | ↑ *** | ↑ *** | ** | | *** | | *** |
| Eubacteriaceae | ↑ *** | | * | | ** | | |
| Peptococcaceae_1 | ↓ ** | ↓ ** | ** | ** | | ** | |
| Anaeroplasmataceae | ↓ ** | | * | ** | | * | |
| Staphylococcaceae | ↓ ** | | | | | | |
| Cryomorpaceae | ↑ * | ↑ ** | * | | ** | ** | |
| Phyllobacteriaceae | ↑ * | ↑ * | | | * | | * |
| Alcaligenaceae | ↑ * | ↑ * | | | | | |
| Pseudomonadaceae | ↑ * | ↑ * | | | * | | * |
| Bacteroidaceae | ↑ * | | | * | | | |
| Veillonellaceae | ↑ * | | | | | | |
| Enterobacteriaceae | ↑ * | | | | | | |
| Lactobacillaceae | ↑ * | | | | | | |
| Gracilibacteraceae | ↑ * | | | | | | |
| Peptococcaceae_2 | | ↓ *** | | | | * | |
| Rhizobiaceae | | ↑ ** | | | | | |
| Incertae_Sedis_IV | | ↑ ** | | | | * | |
| Microbacteriaceae | | ↑ * | | | | | |
| Nocardiaceae | | ↑ * | | | | | |
| Coriobacteriaceae | | ↑ * | | | | * | |
| Flavobacteriaceae | | ↑ * | | | | | * |
| Sphingobacteriaceae | | ↑ * | | | * | | * |
| Bradyrhizobiaceae | | ↑ * | | | | | |
| Caulobacteraceae | | ↑ * | | | | | * |
| Burkholderiaceae | | ↑ * | | | | | * |
| Comamonadaceae | | ↑ * | | | | | |

*P<0.05, **P<0.01, ***P<0.001, ****P<0.0001.

Somewhat to our surprise, our data suggest that HFD and exercise independently impact the different behavioral domains of anxiety and cognition, much like the orthogonal effects on the gut microbiome. Though the cognition associations with specific OTUs were weaker compared to anxiety, we nonetheless found significant relationships that all fell within Clostridiales, a diverse taxonomic order within the phyla Firmicutes that has not been previously associated with cognition. Clearly further studies with larger samples sizes are required to confirm these associations.

Numerous studies have described how the gut microbiome is altered by a HFD and obesity including changes in Firmicutes and Bacteroidetes [28-31]. Though only a very minor phylum in terms of representation, we also found that HFD reduced the relative abundance of Tenericutes, driven almost entirely by a single OTU (#67) from the genus *Anaeroplasm* (>99% probability as shown in Additional file 2: Table S2) that has not been previously implicated in dietary manipulations. HFD induced large changes at all levels of taxonomy,



including in the two most abundant classes/orders (Bacteroidia/Bacteroidales and Clostridia/Clostridiales). We also found HFD caused many significant changes in taxa of more minor abundance. For example, highly significant alterations were seen for several orders (e.g. Lactobacillales, Erysipelotrichales, and Anaeroplasmatales) and classes (e.g. Erysipelotrichia, Mollicutes, and Bacilli). Subsequent studies will determine which of these are the most important players, but probiotic studies support the idea that even taxa of minor abundance can still elicit large effects on the host. Current methods of bacterial profiling as we have used here do not provide sufficient coverage of the entire 16S region to reliably identify changes at the species level, but such studies would allow for more direct translational potential of findings (e.g. pre- or probiotics).

Many studies have documented the negative consequences of dietary fat intake and obesity on behavior and brain function in rodents [32-36]. Diet-induced obesity has been shown to promote depressive-like behavior due to altered brain reward circuitry in mice [32]. Interestingly, caloric restriction has been shown to reduce anxiety and depression related behaviors in mice [37]. One study demonstrated HFD-induced anxiety behavior and changes in the gut microbiome at the level of phylum, but the techniques used did not allow for high resolution analysis [38]. In addition to low anxiety phenotypes in germ-free mice [14,15], some studies have shown that modulation of the gut microbiota either by diet or probiotics can impact anxiety [39,40]. The majority of anxiety associations from our study were a subset of the same taxa that associated with body weight. Interestingly, alterations of Ruminococcaceae and Lachnospiraceae were found in mice exposed to a grid floor stress paradigm [41]. We found a few associations with anxiety that were independent of body weight alterations, including OTU37 (Ruminococcaceae; *Clostridium_IV*), OTU40 (*Barnesiella*), and OTU48 (*Eubacterium*), and these as well as OTU30 (*Lactococcus*) have no reported associations with anxiety. It is important to point out that we cannot establish causality from these studies. Indeed, others have found that manipulations that cause anxiety in rodents can also result in changes to the gut microbial community [42,43]. These “top down” vs “bottom up” dynamics between the CNS and the gut microbiota require further study to fully delineate the causality, but it is highly likely that both mechanisms are at play with some feedback at both levels (i.e. a true system). Likewise, further studies on high fat diet and exercise are required to determine whether these specific bacteria are biomarkers of anxiety or actually drivers of the anxiety phenotypes or both.

Conclusions

In this study we found that a HFD was able to cause significant anxiety with no rescue by exercise while exercise,

but not HFD, was able to enhance cognition. We also found that exercise alone robustly altered the gut microbial community and did not rescue the changes induced by a HFD but, in fact, the changes caused by exercise were completely orthogonal to those induced by a HFD. Additionally, we found numerous, independent associations of specific OTUs and taxa with body weight, anxiety, and cognition that will need to be empirically tested to determine their importance. These data have important public health implications not only in terms of obesity but also determining how the gut microbial community relates to behavioral domains and how it may be used as a biomarker or reshaped to effect changes in anxiety and cognition.

Methods

Mice and treatments

Adult male C57BL/6 J mice were purchased from Jackson Laboratories (Bar Harbor, MN) and were randomly assigned to $n = 10/\text{group}$. Beginning at 8 weeks of age, mice were randomized to one of four groups: ND, ND + exercise, HFD, and HFD + exercise. Diet consisted of normal diet (10% of calories from fat; Research Diets, New Brunswick, NJ #D12450B) or high fat diet (60% of calories from fat; Research Diets #D12492). After an initial 2 week training period where the speed was gradually increased from 3 to 7 m/min and duration from 6 to 60 minutes per day, exercised groups were placed in running wheels (Lafayette Instruments, Lafayette, IN) for one hour at 7 m/min every morning for 5 days/week for the remainder of the study with the exception of the behavioral testing days. Control “sedentary” groups were placed in adjacent running wheels that rotated at a speed that just prevented them from sleeping (~1 m/min) to control for environmental enrichment and handling. All procedures were in accordance with guidelines from the Institutional Animal Care and Use Committee under an approved protocol.

Behavioral analysis

For all behavioral tests, mice were acclimated in the room for an hour prior to the onset of testing. Data was recorded and monitored using overhead cameras and tracked with Anymaze software (Stoelting Co., Wood Dale, IL). Three days following the last exercise session, mice were tested on consecutive days in the open field assay, light/dark box, three-chamber social apparatus, and contextual fear conditioning (over two days) always in the first half of the light phase.

Open field assay (OFA)

Mice were placed in a 40 × 40 × 30 cm (W × L × H) opaque Perspex box and activity was tracked during the entire 15 minute observation period.

Light/Dark exploration (LDE) assay

Mice were placed in a square 40 × 40 × 30 cm Perspex box equally divided into two compartments with a small open door joining the light and dark compartments. The dark compartment was completely covered and mice were tested by placing them at the far end of the lit chamber facing away from the dark chamber and their activity was tracked for 10 minutes.

Three chamber social interaction (3SI)

Social behavior was tested in a three-chamber Perspex 40 × 40 cm box consisting of two 17x40 cm regions separated by 2 dividers forming a smaller 5 × 40 cm center region. Mice were able to move freely through a small 8 × 5cm opening that was aligned in both dividers. Each of the larger chambers contained an inverted wire mesh cylinder in opposing corners. Mice were initially acclimated to the box and empty cylinders for 4 min and then placed in temporary holding cages. A male probe mouse of the same strain was placed in one of the inverted mesh cylinders and allowed to acclimate for 3 minutes prior to placing the test mouse back in the box. The open mesh enabled visual, olfactory, and auditory interactions between probe and test mice. Test mice were subsequently monitored for 10 min in the presence of the probe mouse and interaction score was calculated with the following formula: (Time Mouse cup – Time empty cup) / (Time Mouse cup + Time empty cup).

Contextual fear conditioning assay (CFC)

This test was conducted in a sound attenuated chamber with a grid floor capable of delivering an electric shock and freezing was measured with an overhead camera and FreezeFrame software (Actimetrics, Wilmette, IL). Mice were initially placed into the chamber and undisturbed for 2 minutes, during which time baseline freezing behavior was recorded. An 80-dB white noise served as the conditioned stimulus (CS) and was presented for 30 sec. During the final 2 sec of this noise, mice received a mild foot shock (0.5 mA), which served as the unconditioned stimulus (US). After 1 minute, another CS-US pair was presented. The mouse was removed 30 sec after the second CS-US pair and returned to its home cage. Twenty-four hours later, each mouse was returned to the test chamber and freezing behavior was recorded for 5 minutes (context test). Mice were returned to their home cage and placed in a different room than previously tested in reduced lighting conditions for a period of no less than one hour. For the auditory CS test, environmental and contextual cues were changed by: wiping testing boxes with 30% isopropyl alcohol instead of 30% ethanol; replacing white house lights with red house lights; placing a colored plastic triangular insert in the chamber to alter its shape and spatial cues; covering the

wire grid floor with opaque plastic; and altering the smell in the chamber with vanilla extract. The animals were placed in the apparatus for 3 min and then the auditory CS was presented and freezing was recorded for another 3 min (cued test). Baseline freezing behavior obtained during training was subtracted from the context or cued tests to control for variability in each animal.

Fecal DNA isolation and 16S amplicon preparation and sequencing

Fecal samples left behind in the open field apparatus were collected and immediately frozen on dry ice until processing. Fecal DNA was isolated using a PowerSoil kit (MoBio, Carlsbad, CA) according to directions.

Amplification targeted V3 and V5 regions of the bacterial 16S subunit [44] using primers 357 F (AATGATACGGCGACCACCGAGATCTACACTATGGTAATTGTCTACGGGAGGCAGCAG) and 926R (CAAGCAGAAGACGGCATAACGAGAT-NNNNNNNNNNNN-AGTCAGTCAAGCCCGTCAATTCMTTTRAGT) with barcodes 1–40 from Caporaso et al [45]. PCR was run through 34 cycles of 98°C for 15 seconds, 70°C for 20 seconds, 72°C for 15 seconds with Kapa Hotstart Hi-Fi (Kapa Biosystems, Boston, MA). Electrophoresis of a small sample was used to verify amplicon specificity and purification was carried out using magnetic beads. DNA for each amplicon was then diluted to 10nM and pooled for sequencing on a MiSeq (Illumina, San Diego, CA) using a 300 cycle kit and custom read1 (TATGGTAATTGTCTACGGGAGGCAGCAG), read2 (AGTCAGTCAAGCCCGTCAATTCMTTTRAGT), and index (ACTYAAAKGAATTGACGGGGCTGACTGACT) sequencing primers.

Pipeline for processing of 16S data

Pre-processed sequence files are then subject to quality filtering using Trimmomatic [46] version 0.22, with a hard cutoff of PHRED score Q3 for 5' and 3' ends of the reads (parameters LEADING:3 and TRAILING:3), trimming of the 3' end with a moving average score of Q15, with a window size of 4 bases (parameter SLIDINGWINDOW:4:15), and removing any remaining reads shorter than 75% of the original read length (parameter MINLEN:112 for reads of 150 bp long). Finally reads with any ambiguous base calls or with homopolymers longer than 10 bases long are discarded using mothur [47]. Only the read pairs that survived the quality filter were processed further. Any surviving reads that were unpaired (that is, they lost their matching pair due to low quality) were discarded. Surviving read pairs were then grouped into two files, one each for "read 1" and "read 2" sequences. Reads were also dereplicated, consolidating identical reads to avoid redundant processing.

Each of the two read libraries was then checked for chimeras using UCHIME in de novo mode.

Taxonomy assignment

To prepare the reads for this step, we took reads from the previous step just before the stitching procedure, remove the gaps and then stitched them with a pad sequence of “N” bases. The specific scripts used for these file manipulations are publically available as part of the IM-TORNADO 16S rRNA analysis pipeline (<http://sourceforge.net/projects/intornado/>). Since by default most Bayesian classifiers use 8-mers to perform the classification, we used “NNNNNNNN” as the padding. The stitched reads were then classified using the Greengenes taxonomy (Greengenes99 database version 12.10) as the reference [48] using the following mothur “classify.seqs” command with “iter = 1000”.

Clustering, representative sequences, and chimera removal

Paired-end reads were concatenated directly with no padding and dereplicated. OTU representatives were selected and used to generate a reference set for clustering using USEARCH command “usearch7.0.1090_i86linux32” [49]. Clustering width was set at 97% (-otu_radius_pct 3.0).

Ordination plot

Distance between communities was measured using unweighted Unifrac as implemented in QIIME version 1.8 [16]. Ordination plot was generated using multidimensional scaling analysis as implemented in R and displayed using the vegan package for ecological analyses (<http://CRAN.R-project.org/package=vegan>).

Heatmap

METAGENassist [50] heatmap was produced using the 40 most significant OTU features. Data was log normalized in METAGENassist. Clustering of the groups was carried out using complete linkage clustering and the Spearman distance metric.

Correlation analysis

Relative abundances of bacteria from different levels of taxonomy were analyzed by linear regression with R^2 values indicating goodness of fit and p values for slope calculated by F test with Benjamini-Hochberg correction for false discovery rate (FDR) where indicated.

Additional files

Additional file 1: Figure S1. Impact of diet and exercise on other mouse behaviors. Neither HFD nor exercise altered total locomotor activity (A) or rearing (B) in the open field assay or in sociability (C) in the three chamber social test. As with the contextual memory, exercised

mice had a trend toward increased cued memory (D, $p = 0.051$) in the cued portion of the contextual fear conditioning assay.

Additional file 2: Table S2. OTU and probabilities of correct taxonomic classification (in parentheses).

Additional file 3: Table S1. Significant effects of diet and exercise at different levels of taxonomy. * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$, **** $P < 0.0001$.

Additional file 4: Figure S2. Anxiety levels were associated with several bacteria at the level of phyla, class, order, and family. Color scheme of data points follows the format as in Figure 1 (ND = blue, ND + exercise = green, HFD = red, and HFD + exercise in orange). Linear regression analysis for individual mice was performed with R^2 values indicating goodness of fit and uncorrected p values for slope calculated by F test.

Additional file 5: Table S3. Associations of bacterial abundances with body weight. * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$, **** $P < 0.0001$.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

SSK, NC, and JDF designed and performed this study, analyzed data, and wrote the manuscript. PRJ, MEBM, MDC, KW, NG, JAW, and BAW assisted in purification, sequencing, and analysis of fecal DNA. AK assisted in behavioral analysis. All authors discussed the results and provided comments on the manuscript. All authors read and approved the final manuscript.

Acknowledgements

This work was funded by the Mayo Clinic Center for Individualized Medicine (JDF, NC), Mayo Clinic/University of Illinois Strategic Alliance for Technology-Based Healthcare (PRJ, BAW), Gerstner Family Foundation Career Development Award (JDF), Division of Nutritional Science Vision 20/20 Program at the University of Illinois Urbana-Champaign Hatch project ILLU-971-335 (JAW), and the National Institutes of Health/National Cancer Institute (NIH/NCI) #CA179243 (NC).

Author details

¹Department of Neuroscience, Mayo Clinic, 4500 San Pablo Rd, Jacksonville, FL 32224, USA. ²Department of Surgical Research, Mayo Clinic, 200 First St SW, Rochester, MN 55905, USA. ³Departments of Kinesiology and Community Health, University of Illinois at Urbana-Champaign, Urbana, IL, USA. ⁴Integrative Immunology and Behavior Group, University of Illinois at Urbana-Champaign, Urbana, IL, USA. ⁵Institute for Genomic Biology, University of Illinois at Urbana-Champaign, Urbana, IL, USA. ⁶Loomis Laboratory of Physics, University of Illinois at Urbana-Champaign, Urbana, IL, USA. ⁷Center for Immunology and Immune Therapies, Mayo Clinic, 4500 San Pablo Rd, Jacksonville, FL 32224, USA. ⁸Neurobiology of Disease Graduate Program, Mayo Clinic College of Medicine, 4500 San Pablo Rd., Jacksonville, FL 32224, USA.

Received: 26 June 2014 Accepted: 9 September 2014

Published: 13 September 2014

References

1. Flegal KM, Carroll MD, Kit BK, Ogden CL: Prevalence of obesity and trends in the distribution of body mass index among us adults, 1999–2010. *JAMA* 2012, **307**:491–497.
2. Dixon JB: The effect of obesity on health outcomes. *Mol Cell Endocrinol* 2010, **316**:104–108.
3. Guh DP, Zhang W, Bansback N, Amarsi Z, Birmingham CL, Anis AH: The incidence of co-morbidities related to obesity and overweight: A systematic review and meta-analysis. *BMC Public Health* 2009, **9**:88.
4. Taras H, Potts-Datema W: Obesity and student performance at school. *J Sch Health* 2005, **75**:291–295.
5. Gunstad J, Lhotsky A, Wendell CR, Ferrucci L, Zonderman AB: Longitudinal examination of obesity and cognitive function: results from the Baltimore longitudinal study of aging. *Neuroepidemiology* 2010, **34**:222–229.
6. Elias MF, Elias PK, Sullivan LM, Wolf PA, D'Agostino RB: Lower cognitive function in the presence of obesity and hypertension: the Framingham heart study. *Int J Obes* 2003, **27**:260–268.

7. Hillman CH, Erickson KI, Kramer AF: **Be smart, exercise your heart: exercise effects on brain and cognition.** *Nat Rev Neurosci* 2008, **9**:58–65.
8. Hilbert A, Braehler E, Haeuser W, Zenger M: **Weight bias internalization, core self-evaluation, and health in overweight and obese persons.** *Obes Silver Spring Md* 2013, doi: 10.1002/oby.20561.
9. Mattson MP: **Energy intake and exercise as determinants of brain health and vulnerability to injury and disease.** *Cell Metab* 2012, **16**:706–722.
10. Turnbaugh PJ, Ley RE, Mahowald MA, Magrini V, Mardis ER, Gordon JI: **An obesity-associated gut microbiome with increased capacity for energy harvest.** *Nature* 2006, **444**:1027–1031.
11. Turnbaugh PJ, Gordon JI: **The core gut microbiome, energy balance and obesity.** *J Physiol* 2009, **587**:4153–4158.
12. Jumpertz R, Le DS, Turnbaugh PJ, Trinidad C, Bogardus C, Gordon JI, Krakoff J: **Energy-balance studies reveal associations between gut microbes, caloric load, and nutrient absorption in humans.** *Am J Clin Nutr* 2011, **94**:58–65.
13. Smith MI, Yatsunenko T, Manary MJ, Trehan I, Mkakosya R, Cheng J, Kau AL, Rich SS, Concannon P, Mychaleckyj JC, Liu J, Houpt E, Li JV, Holmes E, Nicholson J, Knights D, Ursell LK, Knight R, Gordon JI: **Gut microbiomes of Malawian twin pairs discordant for kwashiorkor.** *Science* 2013, **339**:548–554.
14. Clarke G, Grenham S, Scully P, Fitzgerald P, Moloney RD, Shanahan F, Dinan TG, Cryan JF: **The microbiome-gut-brain axis during early life regulates the hippocampal serotonergic system in a sex-dependent manner.** *Mol Psychiatry* 2012, doi: 10.1038/mp.2012.77.
15. Diaz Heijtz R, Wang S, Anuar F, Qian Y, Björkholm B, Samuelsson A, Hibberd ML, Forsberg H, Pettersson S: **Normal gut microbiota modulates brain development and behavior.** *Proc Natl Acad Sci* 2011, **20110529**. doi:10.1073/pnas.1010529108.
16. Sipos M, Jeraldo P, Chia N, Qu A, Dhillon AS, Konkel ME, Nelson KE, White BA, Goldenfeld N: **Robust computational analysis of rRNA hypervariable tag datasets.** *PLoS One* 2010, **5**:e15220.
17. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, Fierer N, Peña AG, Goodrich JK, Gordon JI, Huttley GA, Kelley ST, Knights D, Koenig JE, Ley RE, Lozupone CA, McDonald D, Muegge BD, Pirrung N, Reeder J, Sevinsky JR, Turnbaugh PJ, Walters WA, Widmann J, Yatsunenko T, Zaneveld J, Knight R: **QIIME allows analysis of high-throughput community sequencing data.** *Nat Methods* 2010, **7**:335–336.
18. Kuczynski J, Stombaugh J, Walters WA, González A, Caporaso JG, Knight R: **Using QIIME to analyze 16S rRNA gene sequences from microbial communities.** *Curr Protoc Microbiol* 2012, **1**:Unit 1E.5.
19. Matsumoto M, Inoue R, Tsukahara T, Ushida K, Chiji H, Matsubara N, Hara H: **Voluntary running exercise alters microbiota composition and increases n-butyrate concentration in the rat cecum.** *Biosci Biotechnol Biochem* 2008, **72**:572–576.
20. Queipo-Ortuño MI, Seoane LM, Murri M, Pardo M, Gomez-Zumaquero JM, Cardona F, Casanueva F, Tinahones FJ: **Gut microbiota composition in male rat models under different nutritional status and physical activity and its association with serum leptin and ghrelin levels.** *PLoS One* 2013, **8**:e65465.
21. Choi JJ, Eum SY, Rampersaud E, Daunert S, Abreu MT, Toborek M: **Exercise attenuates PCB-induced changes in the mouse gut microbiome.** *Environ Health Perspect* 2013, **121**:725–730.
22. Zhang C, Li S, Yang L, Huang P, Li W, Wang S, Zhao G, Zhang M, Pang X, Yan Z, Liu Y, Zhao L: **Structural modulation of gut microbiota in life-long calorie-restricted mice.** *Nat Commun* 2013, **4**:2163.
23. Evans CC, LePard KJ, Kwak JW, Stancukas MC, Laskowski S, Dougherty J, Moulton L, Glawe A, Wang Y, Leone V, Antonopoulos DA, Smith D, Chang EB, Ciancio MJ: **Exercise prevents weight gain and alters the gut microbiota in a mouse model of high fat diet-induced obesity.** *PLoS One* 2014, **9**:e92193.
24. Kramer AF, Erickson KI, Colcombe SJ: **Exercise, cognition, and the aging brain.** *J Appl Physiol* 2006, **101**:1237–1242.
25. Pang TYC, Hannan AJ: **Enhancement of cognitive function in models of brain disease through environmental enrichment and physical activity.** *Neuropharmacology* 2013, **64**:515–528.
26. Woo J, Shin KO, Park SY, Jang KS, Kang S: **Effects of exercise and diet change on cognition function and synaptic plasticity in high fat diet induced obese rats.** *Lipids Health Dis* 2013, **12**:144.
27. Molteni R, Wu A, Vaynman S, Ying Z, Barnard RJ, Gómez-Pinilla F: **Exercise reverses the harmful effects of consumption of a high-fat diet on synaptic and behavioral plasticity associated to the action of brain-derived neurotrophic factor.** *Neuroscience* 2004, **123**:429–440.
28. de Wit N, Derrien M, Derrien M, Bosch-Vermeulen H, Oosterink E, Keshtkar S, Duval C, de Vogel-van den Bosch J, Kleerebezem M, Müller M, Van der Meer R: **Saturated fat stimulates obesity and hepatic steatosis and affects gut microbiota composition by an enhanced overflow of dietary fat to the distal intestine.** *Am J Physiol Gastrointest Liver Physiol* 2012, **303**:G589–G599.
29. Murphy EF, Cotter PD, Hogan A, O'Sullivan O, Joyce A, Fouhy F, Clarke SF, Marques TM, O'Toole PW, Stanton C, Quigley EM, Daly C, Ross PR, O'Doherty RM, Shanahan F: **Divergent metabolic outcomes arising from targeted manipulation of the gut microbiota in diet-induced obesity.** *Gut* 2013, **62**:220–226.
30. Ley RE, Turnbaugh PJ, Klein S, Gordon JI: **Microbial ecology: human gut microbes associated with obesity.** *Nature* 2006, **444**:1022–1023.
31. David LA, Maurice CF, Carmody RN, Gootenberg DB, Button JE, Wolfe BE, Ling AV, Devlin AS, Varma Y, Fischbach MA, Biddinger SB, Dutton RJ, Turnbaugh PJ: **Diet rapidly and reproducibly alters the human gut microbiome.** *Nature* 2014, **505**:559–563.
32. Sharma S, Fulton S: **Diet-induced obesity promotes depressive-like behaviour that is associated with neural adaptations in brain reward circuitry.** *Int J Obes* 2013, **37**:382–389.
33. Davis JF, Tracy AL, Schurdak JD, Tschöp MH, Lipton JW, Clegg DJ, Benoit SC: **Exposure to elevated levels of dietary fat attenuates psychostimulant reward and mesolimbic dopamine turnover in the rat.** *Behav Neurosci* 2008, **122**:1257–1263.
34. Yamada N, Katsura G, Ochi Y, Ebihara K, Kusakabe T, Hosoda K, Nakao K: **Impaired CNS leptin action is implicated in depression associated with obesity.** *Endocrinology* 2011, **152**:2634–2643.
35. Kohsaka A, Laposky AD, Ramsey KM, Estrada C, Joshua C, Kobayashi Y, Turek FW, Bass J: **High-fat diet disrupts behavioral and molecular circadian rhythms in mice.** *Cell Metab* 2007, **6**:414–421.
36. Hilakivi-Clarke L, Cho E, Onojafe I: **High-fat diet induces aggressive behavior in male mice and rats.** *Life Sci* 1996, **58**:1653–1660.
37. Yamamoto Y, Tanahashi T, Kawai T, Chikahisa S, Katsura S, Nishida K, Teshima-Kondo S, Sei H, Rokutan K: **Changes in behavior and gene expression induced by caloric restriction in C57BL/6 mice.** *Physiol Genomics* 2009, **39**:227–235.
38. Ohland CL, Kish L, Bell H, Thiesen A, Hotte N, Pankiv E, Madsen KL: **Effects of *Lactobacillus helveticus* on murine behavior are dependent on diet and genotype and correlate with alterations in the gut microbiome.** *Psychoneuroendocrinology* 2013, **38**:1738–1747.
39. Bravo JA, Forsythe P, Chew MV, Escaravage E, Savignac HM, Dinan TG, Bienenstock J, Cryan JF: **Ingestion of *Lactobacillus* strain regulates emotional behavior and central GABA receptor expression in a mouse via the vagus nerve.** *Proc Natl Acad Sci U S A* 2011, **108**:16050–16055.
40. Li W, Dowd SE, Scurlock B, Acosta-Martinez V, Lyte M: **Memory and learning behavior in mice is temporally associated with diet-induced alterations in gut bacteria.** *Physiol Behav* 2009, **96**:557–567.
41. Bangsgaard Bendtsen KM, Krych L, Sørensen DB, Pang W, Nielsen DS, Josefsen K, Hansen LH, Sørensen SJ, Hansen AK: **Gut microbiota composition is correlated to grid food induced stress and behavior in the BALB/c mouse.** *PLoS One* 2012, **7**:e46231.
42. Bailey MT, Dowd SE, Galley JD, Hufnagle AR, Allen RG, Lyte M: **Exposure to a social stressor alters the structure of the intestinal microbiota: implications for stressor-induced immunomodulation.** *Brain Behav Immun* 2011, **25**:397–407.
43. Galley JD, Bailey MT: **Impact of stressor exposure on the interplay between commensal microbiota and host inflammation.** *Gut Microbes* 2014, **5**:390–396.
44. Yu Z, Morrison M: **Comparisons of different hypervariable regions of rrs genes for use in fingerprinting of microbial communities by PCR-denaturing gradient gel electrophoresis.** *Appl Environ Microbiol* 2004, **70**:4800–4806.
45. Caporaso JG, Lauber CL, Walters WA, Berg-Lyons D, Huntley J, Fierer N, Owens SM, Betley J, Fraser L, Bauer M, Gormley N, Gilbert JA, Smith G, Knight R: **Ultra-high-throughput microbial community analysis on the Illumina HiSeq and MiSeq platforms.** *ISME J* 2012, **6**:1621–1624.
46. Lohse M, Bolger AM, Nagel A, Fernie AR, Lunn JE, Stitt M, Usadel B: **RobiNA: a user-friendly, integrated software solution for RNA-Seq-based transcriptomics.** *Nucleic Acids Res* 2012, **40**:W622–W627.
47. Schloss PD, Westcott SL, Ryabin T, Hall JR, Hartmann M, Hollister EB, Lesniewski RA, Oakley BB, Parks DH, Robinson CJ, Sahl JW, Stres B, Thallinger GG, Van Horn DJ, Weber CF: **Introducing mothur: open-source,**

platform-independent, community-supported software for describing and comparing microbial communities. *Appl Environ Microbiol* 2009, **75**:7537–7541.

48. DeSantis TZ, Hugenholtz P, Larsen N, Rojas M, Brodie EL, Keller K, Huber T, Dalevi D, Hu P, Andersen GL: Greengenes, a chimera-checked 16S rRNA gene database and workbench compatible with ARB. *Appl Environ Microbiol* 2006, **72**:5069–5072.
49. Edgar RC: UPARSE: highly accurate OTU sequences from microbial amplicon reads. *Nat Methods* 2013, **10**:996–998.
50. Arndt D, Xia J, Liu Y, Zhou Y, Guo AC, Cruz JA, Sinelnikov I, Budwill K, Nesbø CL, Wishart DS: METAGENassist: a comprehensive web server for comparative metagenomics. *Nucleic Acids Res* 2012, **gks497**. doi:10.1093/nar/gks497.

doi:10.1186/1750-1326-9-36

Cite this article as: Kang *et al.*: Diet and exercise orthogonally alter the gut microbiome and reveal independent associations with anxiety and cognition. *Molecular Neurodegeneration* 2014 **9**:36.

**Submit your next manuscript to BioMed Central
and take full advantage of:**

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

Submit your manuscript at
www.biomedcentral.com/submit



Supplemental Figure 1

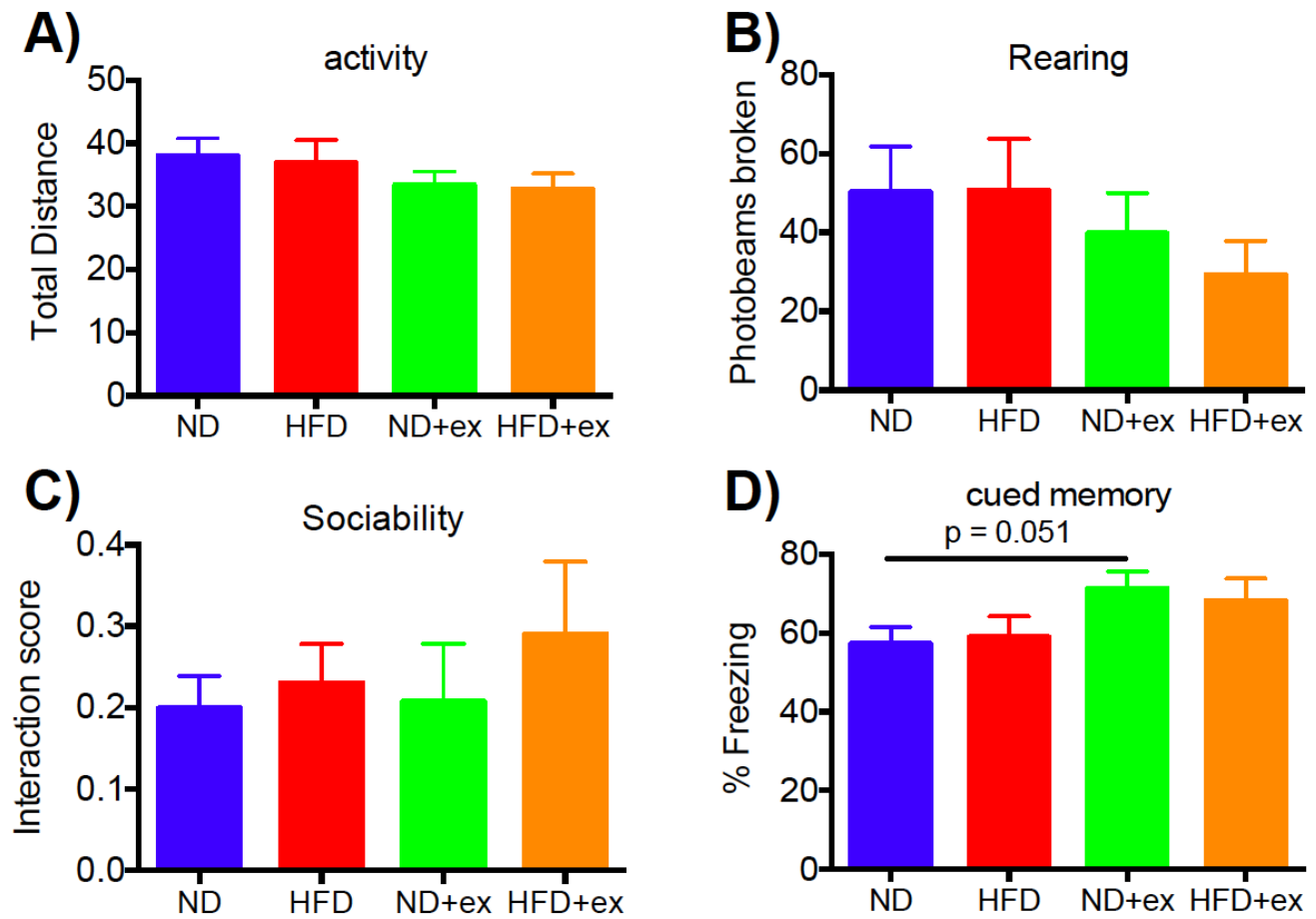


Figure S1. Impact of diet and exercise on other mouse behaviors. Neither HFD nor exercise altered total locomotor activity (A) or rearing (B) in the open field assay or in sociability (C) in the three chamber social test. As with the contextual memory, exercised mice had a trend toward increased cued memory (D, $p = 0.051$) in the cued portion of the contextual fear conditioning assay.

Supplemental Table S2. OTU and probabilities of correct taxonomic classification (in parentheses)

| OTU | Taxonomy probabilities |
|-----|--|
| 1 | Bacteria(100);"Bacteroidetes"(99.9);"Bacteroidia"(98.9);"Bacteroidales"(98.9);"Porphyromonadaceae"(94);Barnesiella(51.7); |
| 2 | Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(98);"Bacteroidales"(98);"Porphyromonadaceae"(96.6);Barnesiella(87.6); |
| 3 | Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(99.6);"Bacteroidales"(99.6);"Porphyromonadaceae"(97.7);Barnesiella(74.7); |
| 4 | Bacteria(100);Firmicutes(99);Clostridia(98.9);Clostridiales(98.9);Lachnospiraceae(90.5);Syntrophococcus(22.5); |
| 5 | Bacteria(100);Firmicutes(99.6);Clostridia(99.6);Clostridiales(99.6);Lachnospiraceae(95.5);Clostridium_XIVa(32.9); |
| 6 | Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(99.2);"Bacteroidales"(99.2);"Porphyromonadaceae"(96.9);Barnesiella(52.9); |
| 7 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(94.4); |
| 8 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(90.5); |
| 9 | Bacteria(99.9);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Lachnospiraceae(98.4);Marvinbryantia(26.6); |
| 10 | Bacteria(100);Firmicutes(100);Erysipelotrichia(100);Erysipelotrichales(100);Erysipelotrichaceae(100);Turicibacter(100); |
| 11 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Anaerotruncus(43.9); |
| 12 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Peptostreptococcaceae(100);Clostridium_XI(100); |
| 13 | Bacteria(100);Firmicutes(99.8);Clostridia(99.7);Clostridiales(99.7);Eubacteriaceae(59.5);Eubacterium(59.5); |
| 14 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Pseudoflavonifractor(87.6); |
| 15 | Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(98.5);"Bacteroidales"(98.5);"Porphyromonadaceae"(97.9);Barnesiella(68.2); |
| 16 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Clostridium_XIVa(79.9); |
| 17 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(97.1);Clostridium_XIVa(45.2); |
| 18 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Roseburia(54.7); |
| 19 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Marvinbryantia(33.5); |
| 20 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(100); |
| 21 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.5);Lachnospiraceae_incertae_sedis(85.3); |
| 22 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(97.8);Clostridium_XIVa(82.5); |
| 23 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Clostridium_XIVb(100); |
| 24 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Roseburia(73.6); |
| 25 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Flavonifractor(72.5); |
| 26 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(98.9); |
| 27 | Bacteria(100);Firmicutes(71.2);Clostridia(69.4);Clostridiales(68.1);Lachnospiraceae(36.8);Cellulosilyticum(15.1); |
| 28 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.4);Coproccoccus(28.7); |
| 29 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(39.7); |
| 30 | Bacteria(100);Firmicutes(100);Bacilli(100);Lactobacillales(100);Streptococcaceae(100);Lactococcus(100); |
| 31 | Bacteria(100);Firmicutes(99.7);Clostridia(99.7);Clostridiales(99.7);Lachnospiraceae(97.9);Syntrophococcus(38.7); |
| 32 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.2);Clostridium_XIVa(47.8); |
| 33 | Bacteria(100);Firmicutes(93.2);Clostridia(90.7);Clostridiales(89.5);Gracilibacteraceae(22);Gracilibacter(21.9); |
| 34 | Bacteria(100);Firmicutes(98.2);Clostridia(96.4);Clostridiales(96.4);Ruminococcaceae(81.7);Clostridium_IV(43.8); |
| 35 | Bacteria(100);Firmicutes(55.4);Clostridia(48.9);Clostridiales(47.7);Ruminococcaceae(31.3);Butyricoccus(20.4); |
| 36 | Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(98.1);"Bacteroidales"(98.1);"Porphyromonadaceae"(91.5);Barnesiella(51.7); |
| 37 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(71.7); |
| 38 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Hespellia(36.7); |
| 39 | Bacteria(100);Firmicutes(78.8);Clostridia(77.5);Clostridiales(77.3);Ruminococcaceae(60.4);Butyricoccus(31.3); |
| 40 | Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(97.2);"Bacteroidales"(97.2);"Porphyromonadaceae"(80.2);Barnesiella(75.7); |
| 41 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Clostridiaceae_1(100);Clostridium_sensu_stricto(99.4); |
| 42 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Moryella(39.1); |
| 43 | Bacteria(99.9);"Bacteroidetes"(99.4);"Bacteroidia"(96.7);"Bacteroidales"(96.7);"Porphyromonadaceae"(88.4);Barnesiella(59.3); |
| 44 | Bacteria(100);Firmicutes(98.8);Clostridia(98.7);Clostridiales(98.7);Lachnospiraceae(90.2);Marvinbryantia(38.4); |
| 45 | Bacteria(100);Firmicutes(99.5);Clostridia(99.2);Clostridiales(99.2);Lachnospiraceae(91.1);Clostridium_XIVb(48.7); |
| 46 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(98.2); |
| 47 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Acetitomaaculum(19.4); |
| 48 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(99.9);Eubacteriaceae(79.3);Eubacterium(79.3); |
| 49 | Bacteria(100);"Bacteroidetes"(100);Flavobacteria(100);"Flavobacteriales"(100);Flavobacteriaceae(100);Flavobacterium(100); |
| 50 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(98.4);Clostridium_XIVa(40.6); |
| 51 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Clostridium_XIVb(99.9); |
| 52 | Bacteria(100);Firmicutes(80.3);Clostridia(78.4);Clostridiales(78.2);Ruminococcaceae(55.3);Butyricoccus(33.1); |
| 53 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Syntrophococcus(16); |
| 54 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Lachnospiraceae_incertae_sedis(78.3); |
| 55 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Roseburia(100); |
| 56 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(99.4); |
| 57 | Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Lachnospiraceae(99.8);Roseburia(52.9); |
| 58 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(97.7);Coproccoccus(15.9); |
| 59 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Oribacterium(41.7); |
| 60 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(87.2); |
| 61 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Acetanaerobacterium(60.9); |
| 62 | Bacteria(100);Firmicutes(100);Erysipelotrichia(100);Erysipelotrichales(100);Erysipelotrichaceae(100);Clostridium_XVIII(100); |
| 63 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(95.1);Butyrivibrio(16.1); |
| 64 | Bacteria(100);Firmicutes(95.7);Clostridia(94.4);Clostridiales(94.2);Ruminococcaceae(89.2);Flavonifractor(16.3); |
| 65 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Anaerotruncus(85.4); |
| 66 | Bacteria(100);Firmicutes(81.1);Clostridia(76.5);Clostridiales(75);Ruminococcaceae(34.7);Butyricoccus(17.2); |
| 67 | Bacteria(100);"Tenericutes"(100);Mollicutes(100);Anaeroplasmatales(100);Anaeroplasmataceae(100);Anaeroplasmata(100); |
| 68 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(98.7); |
| 69 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(89.4);Coproccoccus(14.3); |
| 70 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(97.5);Moryella(32.5); |
| 71 | Bacteria(100);Firmicutes(46.2);Clostridia(45.2);Clostridiales(41.7);Incertae_Sedis_IV(6.7);Caldicoproccoccus(6.7); |
| 72 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Anaerotruncus(48.4); |
| 73 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Lachnospiraceae_incertae_sedis(39.1); |
| 74 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(93.3);Syntrophococcus(28); |
| 75 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(99.7);Pseudoflavonifractor(65.3); |
| 76 | Bacteria(100);Firmicutes(100);Erysipelotrichia(100);Erysipelotrichales(100);Erysipelotrichaceae(100);Coproccoccus(61.6); |
| 77 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(98.4);Lachnospiraceae_incertae_sedis(22); |
| 78 | Bacteria(100);"Bacteroidetes"(100);"Sphingobacteria"(100);"Sphingobacteriales"(100);Sphingobacteriaceae(100);Pedobacter(100); |
| 79 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(36.2); |
| 80 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(98.4);Clostridium_XIVa(40); |
| 81 | Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Clostridium_XIVa(50.3); |
| 82 | Bacteria(100);Firmicutes(75.4);Clostridia(69.9);Clostridiales(67.9);Ruminococcaceae(35.2);Acetivibrio(18); |
| 83 | Bacteria(100);Firmicutes(100);Erysipelotrichia(100);Erysipelotrichales(100);Erysipelotrichaceae(100);Erysipelotrichaceae_incertae_sedis(98.4); |

84 Bacteria(100);Firmicutes(98.9);Negativicutes(66.4);Selenomonadales(66.4);Veillonellaceae(65.4);Zymophilus(23.4);
85 Bacteria(100);Firmicutes(99.8);Clostridia(99.8);Clostridiales(99.8);Ruminococcaceae(99.1);Clostridium_IV(79.5);
86 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Lachnospiraceae_incertae_sedis(88.1);
87 Bacteria(100);Firmicutes(100);Clostridia(99.9);Clostridiales(99.9);Lachnospiraceae(97.2);Marvinbryantia(54.1);
88 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Lachnospiraceae(96.8);Clostridium_XIVa(29.3);
89 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Butyricoccus(98.4);
90 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Clostridium_XIVa(63.8);
91 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Blautia(43.4);
92 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(99);"Bacteroidales"(99);"Porphyromonadaceae"(94.9);Barnesiella(65.8);
93 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Clostridium_XIVa(54.1);
94 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Clostridium_XIVa(62.3);
95 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Clostridium_XIVa(52.1);
96 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Clostridium_XIVa(73.2);
97 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(95.9);Lachnospiraceae_incertae_sedis(40.3);
98 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(99.9);"Bacteroidales"(99.9);"Porphyromonadaceae"(99.6);Barnesiella(89.8);
99 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(98.6);Clostridium_XIVa(44.4);
100 Bacteria(100);"Actinobacteria"(99.8);Actinobacteria(99.8);Coriobacteriales(99.8);Coriobacteriaceae(99.8);Paraeggerthella(48.2);
101 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Coriobacteriales(100);Coriobacteriaceae(100);Olsenella(80.5);
102 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(96.5);Acetivibrio(53.4);
103 Bacteria(100);Firmicutes(94.6);Clostridia(91.4);Clostridiales(91.4);Ruminococcaceae(85.9);Pseudoflavonifractor(26.4);
104 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.2);Clostridium_XIVa(60.3);
105 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.8);Clostridium_XIVa(32.9);
106 Bacteria(100);Firmicutes(100);Bacilli(100);Lactobacillales(100);Lactobacillaceae(100);Lactobacillus(100);
107 Bacteria(100);Firmicutes(99.8);Clostridia(99.8);Clostridiales(99.8);Lachnospiraceae(99);Marvinbryantia(36.1);
108 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.1);Blautia(31.5);
109 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Dorea(30.6);
110 Bacteria(100);Firmicutes(96.9);Clostridia(59.4);Clostridiales(53.2);Peptococcaceae_2(19.9);Cryptanaerobacter(19.2);
111 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Clostridium_XIVa(38.3);
112 Bacteria(100);Firmicutes(100);Erysipelotrichia(100);Erysipelotrichales(100);Erysipelotrichaceae(100);Erysipelotrichaceae_incertae_sedis(100);
113 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(96.6);Coproccoccus(43.5);
114 Bacteria(100);Firmicutes(100);Clostridia(99.2);Clostridiales(99.2);Ruminococcaceae(92.3);Acetivibrio(77.9);
115 Bacteria(100);Firmicutes(100);Bacilli(100);Lactobacillales(100);Streptococcaceae(100);Streptococcus(100);
116 Bacteria(100);"Proteobacteria"(100);Gammaproteobacteria(100);Pseudomonadales(100);Pseudomonadaceae(100);Pseudomonas(100);
117 Bacteria(100);Firmicutes(100);Bacilli(100);Lactobacillales(100);Enterococcaceae(100);Enterococcus(100);
118 Bacteria(99.9);Firmicutes(87);Erysipelotrichia(53.6);Erysipelotrichales(53.6);Erysipelotrichaceae(53.6);Coproccoccus(13.1);
119 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(96.8);Clostridium_XIVa(66.2);
120 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Clostridium_XIVa(46.4);
121 Bacteria(100);Firmicutes(89.2);Clostridia(82.5);Clostridiales(78.6);Clostridiales_Incertae_Sedis_XIII(60.1);Anaerovorax(60.1);
122 Bacteria(100);"Proteobacteria"(100);Alphaproteobacteria(100);Caulobacteriales(100);Caulobacteraceae(100);Brevundimonas(100);
123 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Anaerotruncus(99.6);
124 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(94.4);
125 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(97.4);Clostridium_XIVa(39.4);
126 Bacteria(100);Firmicutes(84.9);Clostridia(72.1);Clostridiales(69.6);Ruminococcaceae(20.5);Acetivibrio(12.5);
127 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Coriobacteriales(100);Coriobacteriaceae(100);Enterorhabdus(99.2);
128 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Lachnospiraceae_incertae_sedis(36.2);
129 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Coproccoccus(47.7);
130 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Clostridium_XIVb(99.7);
131 Bacteria(100);Firmicutes(100);Bacilli(100);Bacillales(100);Staphylococcaceae(100);Staphylococcus(100);
132 Bacteria(100);Firmicutes(80.5);Clostridia(77.1);Clostridiales(75.6);Ruminococcaceae(48.3);Acetivibrio(22.6);
133 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(43.8);
134 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Flavonifractor(82.7);
135 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(97.1);Blautia(29.5);
136 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(99.9);
137 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Coriobacteriales(100);Coriobacteriaceae(100);Asaccharobacter(81.6);
138 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Clostridium_XIVa(75.8);
139 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Lachnospiraceae_incertae_sedis(94.3);
140 Bacteria(100);Firmicutes(95);Clostridia(93.9);Clostridiales(93.5);Lachnospiraceae(64.7);Cellulosilyticum(22.2);
141 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(95.7);Clostridium_XIVa(31.1);
142 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Eubacteriaceae(52.3);Eubacterium(52.3);
143 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(97.4);Clostridium_XIVa(25.4);
144 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(97);
145 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Butyrivibrio(23.3);
146 Bacteria(100);Firmicutes(99.4);Clostridia(99.4);Clostridiales(99.4);Ruminococcaceae(99);Hydrogenoanaerobacterium(42.5);
147 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(94.3);
148 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Eubacteriaceae(63.7);Eubacterium(63.7);
149 Bacteria(100);Firmicutes(99.7);Clostridia(99.7);Clostridiales(99.7);Ruminococcaceae(99.7);Clostridium_IV(53.2);
150 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(99.1);"Bacteroidales"(99.1);"Porphyromonadaceae"(98.9);Barnesiella(95.5);
151 Bacteria(100);Firmicutes(92.9);Erysipelotrichia(74.9);Erysipelotrichales(74.9);Erysipelotrichaceae(74.9);Coproccoccus(32.5);
152 Bacteria(100);Firmicutes(84.3);Clostridia(83.2);Clostridiales(81.8);Ruminococcaceae(24.2);Butyricoccus(15.1);
153 Bacteria(100);Firmicutes(99.5);Clostridia(89.2);Clostridiales(88.4);Peptococcaceae_1(81);Peptococcus(80.2);
154 Bacteria(100);Firmicutes(83.3);Clostridia(81.6);Clostridiales(81.3);Incertae_Sedis_IV(11.4);Caldicoprobacter(11.4);
155 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Flavonifractor(65);
156 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.7);Lachnospiraceae_incertae_sedis(50.9);
157 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Flavonifractor(95.1);
158 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(99.9);Clostridium_IV(70.7);
159 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(99.9);Clostridium_IV(43.7);
160 Bacteria(100);Firmicutes(99.8);Erysipelotrichia(95.4);Erysipelotrichales(95.4);Erysipelotrichaceae(95.4);Turicibacter(95.4);
161 Bacteria(100);"Bacteroidetes"(66.4);"Bacteroidia"(54);"Bacteroidales"(54);"Porphyromonadaceae"(43.4);Tannerella(11.6);
162 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(96.3);Johnsonella(24.5);
163 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Lachnospiraceae_incertae_sedis(93);
164 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(84.4);
165 Bacteria(100);"Proteobacteria"(100);Gammaproteobacteria(100);Xanthomonadales(100);Xanthomonadaceae(100);Stenotrophomonas(99.7);
166 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Clostridium_XIVa(66.7);
167 Bacteria(100);Firmicutes(75.6);Erysipelotrichia(75.6);Erysipelotrichales(75.6);Erysipelotrichaceae(75.6);Holdemania(55.6);
168 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.1);Oribacterium(58.6);
169 Bacteria(100);Firmicutes(100);Bacilli(100);Bacillales(100);Staphylococcaceae(100);Staphylococcus(100);

170 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Roseburia(77.7);
171 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Clostridium_XIVa(75.2);
172 Bacteria(100);Firmicutes(100);Bacilli(100);Bacillales(100);Staphylococcaceae(100);Jeotgallcoccus(100);
173 Bacteria(100);Firmicutes(99.9);Clostridia(99.3);Clostridiales(99.3);Eubacteriaceae(63.4);Eubacterium(63.4);
174 Bacteria(99.9);"Bacteroidetes"(54.8);"Bacteroidia"(39.8);"Bacteroidales"(39.8);"Porphyromonadaceae"(36.5);Tannerella(22.9);
175 Bacteria(100);Firmicutes(94.4);Clostridia(91.3);Clostridiales(90.4);Clostridiales_Incertae_Sedis_XII(39.3);Acidaminobacter(30.5);
176 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Lachnospiraceae_Incertae_Sedis(55.8);
177 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(97);Clostridium_XIVa(66.8);
178 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.7);Clostridium_XIVa(44);
179 Bacteria(100);Firmicutes(98.4);Clostridia(98.3);Clostridiales(98.3);Ruminococcaceae(98.3);Oscillibacter(82.9);
180 Bacteria(100);"Proteobacteria"(100);Gammaproteobacteria(100);"Enterobacteriales"(100);Enterobacteriaceae(100);Cronobacter(43);
181 Bacteria(100);"Proteobacteria"(100);Alphaproteobacteria(100);Rhizobiales(100);Phyllobacteriaceae(100);Phyllobacterium(100);
182 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Bifidobacteriales(100);Bifidobacteriaceae(100);Bifidobacterium(100);
183 Bacteria(100);Firmicutes(80.7);Clostridia(70.8);Clostridiales(70.4);Gracilibacteraceae(31.1);Gracilibacter(29.6);
184 Bacteria(100);Firmicutes(100);Bacilli(100);Bacillales(99.9);Bacillaceae_1(44.5);Bacillus(44.5);
185 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Lachnospiraceae_Incertae_Sedis(80.8);
186 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Anaerotruncus(78.9);
187 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Roseburia(60.2);
188 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.7);Coproccoccus(35.1);
189 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Coriobacteriales(100);Coriobacteriaceae(100);Paraeggerthella(29.4);
190 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(96.2);Lachnospiraceae_Incertae_Sedis(27);
191 Bacteria(100);Firmicutes(92.9);Clostridia(92.9);Clostridiales(92.9);Ruminococcaceae(92.8);Pseudoflavonifractor(52.1);
192 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Sporobacter(48.9);
193 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(100);
194 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(93.7);Clostridium_XIVa(60.2);
195 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(99.9);Anaerotruncus(57.9);
196 Bacteria(100);Firmicutes(94.6);Clostridia(93.4);Clostridiales(92.4);Clostridiales_Incertae_Sedis_XII(26.6);Guggenheimella(16.5);
197 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(92.9);Lachnospiraceae_Incertae_Sedis(37.7);
198 Bacteria(100);"Proteobacteria"(100);Betaproteobacteria(100);Burkholderiales(100);Burkholderiaceae(100);Ralstonia(99.9);
199 Bacteria(99.9);"Bacteroidetes"(51);"Bacteroidia"(36);"Bacteroidales"(36);"Porphyromonadaceae"(30.8);Tannerella(14.1);
200 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Lachnospiraceae_Incertae_Sedis(36.4);
201 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.8);Blautia(31.8);
202 Bacteria(100);Firmicutes(77.8);Clostridia(77.7);Clostridiales(77.7);Peptostreptococcaceae(73);Sporacetigenium(56.5);
203 Bacteria(100);Firmicutes(66.3);Clostridia(64.7);Clostridiales(64.6);Ruminococcaceae(63.6);Ethanoligenens(51.8);
204 Bacteria(100);"Proteobacteria"(100);Betaproteobacteria(100);Burkholderiales(100);Comamonadaceae(100);Delftia(99.3);
205 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Lachnospiraceae(97.9);Marvinbryantia(44.5);
206 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Actinomycetales(100);Nocardiaceae(100);Rhodococcus(100);
207 Bacteria(100);"Bacteroidetes"(89.2);"Bacteroidia"(87.8);"Bacteroidales"(87.8);"Porphyromonadaceae"(75.2);Tannerella(45.5);
208 Bacteria(100);Firmicutes(92.5);Clostridia(90.2);Clostridiales(88.4);Ruminococcaceae(46.9);Clostridium_IV(32.2);
209 Bacteria(100);Firmicutes(97.2);Erysipelotrichia(96.9);Erysipelotrichales(96.9);Erysipelotrichaceae(96.9);Turicibacter(96.9);
210 Bacteria(100);Firmicutes(68.4);Clostridia(65.2);Clostridiales(65);Ruminococcaceae(31.7);Butyricoccus(20.1);
211 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Coriobacteriales(100);Coriobacteriaceae(100);Enterorhabdus(100);
212 Bacteria(100);Firmicutes(77);Clostridia(76.1);Clostridiales(76);Clostridiaceae_1(61);Anaerobacter(58.6);
213 Bacteria(100);Firmicutes(85.6);Clostridia(85.1);Clostridiales(84.7);Ruminococcaceae(44.2);Clostridium_IV(32.4);
214 Bacteria(100);Firmicutes(56.4);Clostridia(56.4);Clostridiales(56.4);Lachnospiraceae(56.3);Johnsonella(48.1);
215 Bacteria(100);"Proteobacteria"(100);Alphaproteobacteria(100);Rhizobiales(100);Hyphomicrobiaceae(100);Devosia(100);
216 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(98.8);"Bacteroidales"(98.8);"Porphyromonadaceae"(95.7);Barnesiella(43.7);
217 Bacteria(100);Firmicutes(99.8);Erysipelotrichia(99.7);Erysipelotrichales(99.7);Erysipelotrichaceae(99.7);Allobaculum(85.4);
218 Bacteria(100);Firmicutes(70.4);Clostridia(69.9);Clostridiales(69.9);Ruminococcaceae(69);Anaerotruncus(37);
219 Bacteria(100);Firmicutes(96.7);Clostridia(96.5);Clostridiales(96.5);Lachnospiraceae(97.3);Marvinbryantia(39.2);
220 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(62.1);Oribacterium(10.7);
221 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(95.6);
222 Bacteria(100);"Proteobacteria"(100);Alphaproteobacteria(100);Sphingomonadales(100);Sphingomonadaceae(100);Sphingomonas(99.9);
223 Bacteria(99.9);Firmicutes(96.7);Clostridia(96.5);Clostridiales(96.5);Lachnospiraceae(86.5);Robinsoniella(14.5);
224 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Butyrivibrio(49);
225 Bacteria(100);"Proteobacteria"(100);Gammaproteobacteria(100);Xanthomonadales(100);Xanthomonadaceae(100);Stenotrophomonas(94.6);
226 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Lachnobacterium(24);
227 Bacteria(100);Firmicutes(100);Bacilli(75.8);Lactobacillales(100);Streptococcaceae(100);Streptococcus(100);
228 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Lachnospiraceae_Incertae_Sedis(47.6);
229 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Coriobacteriales(100);Coriobacteriaceae(100);Adlercreutzia(4.6);
230 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);Bacteroidaceae(100);Bacteroides(100);
231 Bacteria(100);Firmicutes(99.8);Bacilli(96.5);Lactobacillales(96.4);Streptococcaceae(96.3);Lactococcus(94.9);
232 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Lachnospiraceae_Incertae_Sedis(66.7);
233 Bacteria(100);"Proteobacteria"(100);Alphaproteobacteria(100);Rhizobiales(100);Rhizobiaceae(100);Rhizobium(100);
234 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(94.7);Syntrophococcus(35);
235 Bacteria(100);Firmicutes(59.8);Clostridia(59.3);Clostridiales(59.3);Clostridiaceae_1(52.2);Anaerobacter(49);
236 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);"Prevotellaceae"(100);Prevotella(92.1);
237 Bacteria(100);"Bacteroidetes"(86.8);"Bacteroidia"(60.8);"Bacteroidales"(60.8);"Porphyromonadaceae"(52.9);Tannerella(22.4);
238 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Flavonifractor(63);
239 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Flavonifractor(59.1);
240 Bacteria(100);Firmicutes(99.7);Bacilli(75.8);Lactobacillales(72.7);Enterococcaceae(17.3);Melissococcus(15.2);
241 Bacteria(100);"Proteobacteria"(100);Betaproteobacteria(100);Burkholderiales(100);Alcaligenaceae(100);Achromobacter(97.4);
242 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);"Porphyromonadaceae"(100);Parabacteroides(100);
243 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Actinomycetales(100);Propionibacteriaceae(100);Propionibacterium(100);
244 Bacteria(100);"Bacteroidetes"(100);"Sphingobacteria"(100);"Sphingobacteriales"(100);Sphingobacteriaceae(100);Sphingobacterium(100);
245 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Lachnospiraceae(97);Clostridium_XIVa(56.7);
246 Bacteria(100);Firmicutes(98.9);Clostridia(98.9);Clostridiales(98.9);Ruminococcaceae(98.6);Clostridium_IV(81.6);
247 Bacteria(100);Firmicutes(85.9);Clostridia(85.9);Clostridiales(85.9);Lachnospiraceae(82.5);Syntrophococcus(22.9);
248 Bacteria(100);"Bacteroidetes"(97.8);"Bacteroidia"(92.8);"Bacteroidales"(92.8);"Porphyromonadaceae"(84.9);Tannerella(37.6);
249 Bacteria(100);Firmicutes(92.2);Clostridia(87);Clostridiales(79.9);Ruminococcaceae(56.8);Saccharofermentans(21.6);
250 Eukaryota(11.9);Streptophyta(9.9);Liliopsida(9.9);Poales(9.9);Poaceae(9.9);Zea(9.9);
251 Bacteria(100);Firmicutes(100);Clostridia(94);Clostridiales(94);Ruminococcaceae(94);Flavonifractor(84.9);
252 Bacteria(87.6);"Elusimicrobia"(4.1);Elusimicrobia(4.1);Elusimicrobiales(4.1);Elusimicrobiaceae(4.1);Elusimicrobium(4.1);
253 Bacteria(100);Firmicutes(83.7);Clostridia(83.7);Clostridiales(83.7);Ruminococcaceae(83.7);Oscillibacter(64.1);
254 Bacteria(100);"Bacteroidetes"(75.4);"Bacteroidia"(70.5);"Bacteroidales"(70.5);"Porphyromonadaceae"(64.9);Tannerella(49.9);
255 Bacteria(100);"Proteobacteria"(100);Alphaproteobacteria(100);Rhizobiales(100);Phyllobacteriaceae(100);Mesorhizobium(99.3);

256 Bacteria(100);Firmicutes(95.3);Clostridia(95.3);Clostridiales(95.3);Ruminococcaceae(95.2);Flavonifractor(57.8);
257 Bacteria(86.5);"Elusimicrobia"(3.9);Elusimicrobia(3.9);Elusimicrobiales(3.9);Elusimicrobiaceae(3.9);Elusimicrobium(3.9);
258 Bacteria(100);Firmicutes(80.2);Clostridia(80);Clostridiales(80);Peptostreptococcaceae(79.5);Sporacetigenium(72.1);
259 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);Bacteroidaceae(100);Bacteroides(100);
260 Bacteria(100);Firmicutes(83.8);Clostridia(83.8);Clostridiales(83.5);Lachnospiraceae(82);Clostridium_XIVb(81.8);
261 Bacteria(100);Firmicutes(86.2);Clostridia(78.7);Clostridiales(78.5);Gracilibacteraceae(32.3);Gracilibacter(30);
262 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(98.1);"Bacteroidales"(98.1);"Porphyromonadaceae"(97.7);Barnesiella(90.3);
263 Bacteria(88.1);"Bacteroidetes"(35.1);"Bacteroidia"(12);"Bacteroidales"(12);"Porphyromonadaceae"(6);Petrimonas(4.4);
264 Bacteria(87.6);"Bacteroidetes"(22.6);Flavobacteria(5.8);"Flavobacteriales"(5.8);Flavobacteriaceae(5.1);Ornithobacterium(1.5);
265 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Lachnospiraceae_incertae_sedis(38.7);
266 Eukaryota(12.3);Arthropoda(8.5);Insecta(8.5);Coleoptera(8.5);Tenebrionidae(8.5);Tribolium(8.5);
267 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);Bacteroidaceae(100);Bacteroides(100);
268 Bacteria(100);Firmicutes(62.9);Clostridia(62.9);Clostridiales(62.9);Ruminococcaceae(62.8);Anaerotruncus(54.9);
269 Eukaryota(13);Streptophyta(11.1);Liliopsida(11.1);Poales(11.1);Poaceae(11.1);Zea(11.1);
270 Archaea(17.5);"Euryarchaeota"(11.7);Thermoplasmata(6.6);Thermoplasmatales(6.6);Thermoplasmatales_incertae_sedis(6.4);Thermogymnomonas(6.4);
271 Bacteria(100);Firmicutes(100);Bacilli(100);Lactobacillales(100);Carnobacteriaceae(100);Atopostipes(99.6);
272 Eukaryota(28.2);Chordata(15.9);Mammalia(15.9);Primates(2.8);Hominidae(2.8);Homo(2.8);
273 Bacteria(88.8);"Bacteroidetes"(24.7);"Sphingobacteria"(14.4);"Sphingobacteriales"(14.4);Sphingobacteriaceae(3.6);Pseudosphingobacterium(2.7);
274 Eukaryota(28.9);Streptophyta(19.7);Liliopsida(19.7);Poales(19.7);Poaceae(19.7);Zea(19.7);
275 Bacteria(100);Firmicutes(96.8);Erysipelotrichia(94);Erysipelotrichales(94);Erysipelotrichaceae(94);Holdemania(49.7);
276 Eukaryota(13.1);Streptophyta(4.9);Liliopsida(4.9);Poales(4.9);Poaceae(4.9);Zea(4.9);
277 Eukaryota(23.1);Chordata(13.3);Mammalia(13.3);Primates(3.8);Hominidae(3.8);Homo(3.8);
278 Bacteria(80.8);Firmicutes(23.5);Clostridia(20);Clostridiales(16.4);Clostridiales_Incertae_Sedis_XI(7.7);Parvimonas(7.5);
279 Bacteria(100);Firmicutes(99.8);Clostridia(99.3);Clostridiales(99.3);Lachnospiraceae(63);Lactonifactor(15.5);
280 Bacteria(100);Firmicutes(95.4);Clostridia(94.8);Clostridiales(94.8);Ruminococcaceae(44.9);Saccharofermentans(22.1);
281 Bacteria(100);Firmicutes(64.3);Erysipelotrichia(63.8);Erysipelotrichales(63.8);Erysipelotrichaceae(63.8);Holdemania(49.6);
282 Bacteria(100);Firmicutes(99.5);Clostridia(99.5);Clostridiales(99.5);Ruminococcaceae(99.5);Oscilibacter(92.8);
283 Eukaryota(8.1);Streptophyta(7.1);Liliopsida(7.1);Poales(7.1);Poaceae(7.1);Zea(7.1);
284 Bacteria(78.9);"Proteobacteria"(29);Deltaproteobacteria(15.6);Myxococcales(7.8);Nannocystaceae(7.5);Enhygromyxa(7.1);
285 Bacteria(100);"Tenericutes"(76.7);Mollicutes(76.7);Anaeroplasmatales(76);Anaeroplasmataceae(76);Anaeroplasmata(76);
286 Bacteria(100);"Proteobacteria"(90.1);Alphaproteobacteria(90.1);Rhizobiales(90.1);Hyphomicrobiaceae(90.1);Gemmiger(90.1);
287 Bacteria(100);Firmicutes(89.4);Bacilli(89.3);Lactobacillales(87.1);Streptococcaceae(86.3);Lactococcus(82.5);
288 Bacteria(100);Firmicutes(76.1);Clostridia(71.7);Clostridiales(69.8);Ruminococcaceae(22.3);Butyricoccus(10.3);
289 Eukaryota(16.8);Streptophyta(13.7);Liliopsida(13.7);Poales(13.7);Poaceae(13.7);Zea(13.7);
290 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Oribacterium(26.4);
291 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);"Prevotellaceae"(100);Prevotella(85.1);
292 Bacteria(87.5);Firmicutes(28.7);Clostridia(22.2);Clostridiales(20.2);Clostridiales_Incertae_Sedis_XI(4.4);Parvimonas(3.4);
293 Bacteria(88.1);"Proteobacteria"(39.2);Gammaproteobacteria(11.8);Legionellales(4.5);Coxiellaceae(4.5);Coxiella(4.5);
294 Bacteria(82.8);"Bacteroidetes"(27.7);"Sphingobacteria"(15.2);"Sphingobacteriales"(15.2);Cytophagaceae(13.3);Litoribacter(5.2);
295 Bacteria(100);"Spirochaetes"(96.8);Spirochaetes(96.8);Spirochaetales(96.8);Spirochaetaceae(96);Spirochaeta(92.6);
296 Archaea(22.1);"Euryarchaeota"(16.4);Halobacteria(11.9);Halobacteriales(11.9);Halobacteriaceae(11.9);Halonotius(5.1);
297 Bacteria(100);Firmicutes(64.5);Clostridia(62.2);Clostridiales(61.1);Ruminococcaceae(20.3);Butyricoccus(12.4);
298 Eukaryota(13.6);Chordata(12.3);Mammalia(12.3);Rodentia(11.4);Muridae(11.4);Mus(11.4);
299 Eukaryota(22.2);Chordata(15.4);Mammalia(15.4);Primates(3.8);Hominidae(3.8);Homo(3.8);
300 Eukaryota(12.8);Streptophyta(11.5);Liliopsida(11.5);Poales(11.5);Poaceae(11.5);Zea(11.5);
301 Bacteria(90.1);"Deferribacteres"(6.9);Deferribacteres(6.9);Deferribacteriales(6.9);Deferribacteraceae(6.9);Mucispirillum(6.8);
302 Bacteria(100);"Proteobacteria"(100);Alphaproteobacteria(100);Rhizobiales(100);Bradyrhizobiaceae(100);Bradyrhizobium(98.7);
303 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Dorea(76.1);
304 Eukaryota(12.2);Streptophyta(10.7);Liliopsida(10.7);Poales(10.7);Poaceae(10.7);Zea(10.7);
305 Eukaryota(23.3);Chordata(21.6);Mammalia(21.6);Rodentia(15.8);Muridae(15.8);Mus(15.8);
306 Bacteria(100);Firmicutes(99.6);Clostridia(99.3);Clostridiales(99.3);Lachnospiraceae(97.1);Marvinbryantia(78.8);
307 Bacteria(88.1);"Bacteroidetes"(31.2);"Bacteroidia"(13.4);"Bacteroidales"(13.4);"Porphyromonadaceae"(7.6);Proteiniphilum(3.9);
308 Bacteria(86.5);Firmicutes(26.4);Clostridia(20.9);Clostridiales(20.2);Clostridiales_Incertae_Sedis_XI(6.3);Parvimonas(4.5);
309 Bacteria(100);Firmicutes(100);Negativicutes(100);Selenomonadales(100);Acidaminococcaceae(100);Phascolarctobacterium(100);
310 Bacteria(100);Firmicutes(100);Erysipelotrichia(100);Erysipelotrichales(100);Erysipelotrichaceae(100);Clostridium_XVIII(100);
311 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(99.9);Oscilibacter(82.2);
312 Bacteria(91.9);Firmicutes(18.4);Bacilli(6.4);Lactobacillales(5.4);Enterococcaceae(4.9);Pilibacter(4.5);
313 Bacteria(84.8);"Proteobacteria"(28.1);Alphaproteobacteria(11.6);Caulobacteriales(5.8);Hyphomonadaceae(5.8);Woodsholea(4.7);
314 Bacteria(100);Firmicutes(100);Clostridia(50.1);Clostridiales(50.1);Lachnospiraceae(50);Johnsonella(44);
315 Eukaryota(4.7);Chordata(3.9);Mammalia(3.9);Rodentia(3.9);Muridae(3.9);Mus(3.9);
316 Bacteria(100);"Proteobacteria"(100);Gammaproteobacteria(100);"Enterobacteriales"(100);Enterobacteriaceae(100);Klebsiella(98.8);
317 Bacteria(100);"Bacteroidetes"(99.9);"Bacteroidia"(95);"Bacteroidales"(95);"Porphyromonadaceae"(89.9);Barnesiella(49.2);
318 Bacteria(100);Firmicutes(95.2);Clostridia(95.2);Clostridiales(95.2);Lachnospiraceae(88.5);Robinsoniella(34);
319 Bacteria(100);Firmicutes(58.7);Bacilli(58.4);Lactobacillales(55.3);Carnobacteriaceae(40.1);Desemzia(16.8);
320 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.3);Clostridium_XIVa(78.4);
321 Bacteria(99.9);"Tenericutes"(58);Mollicutes(58);Anaeroplasmatales(57.8);Anaeroplasmataceae(57.8);Asteroleplasma(57.8);
322 Bacteria(100);"Bacteroidetes"(14.1);"Bacteroidia"(13.9);"Bacteroidales"(13.9);Bacteroidaceae(13.9);Anaerorhabdus(13.9);
323 Eukaryota(12.4);Streptophyta(11.9);Liliopsida(11.9);Poales(11.9);Poaceae(11.9);Zea(11.9);
324 Eukaryota(28);Chordata(27.6);Mammalia(27.6);Rodentia(21.6);Muridae(21.6);Mus(21.6);
325 Eukaryota(11.7);Chordata(8.4);Mammalia(8.4);Rodentia(8.4);Muridae(8.4);Mus(8.4);
326 Archaea(22.3);"Euryarchaeota"(18.8);Halobacteria(10.1);Halobacteriales(10.1);Halobacteriaceae(10.1);Halonotius(5.5);
327 Eukaryota(12.1);Arthropoda(7.6);Insecta(7.6);Coleoptera(7.6);Tenebrionidae(7.6);Tribolium(7.6);
328 Eukaryota(19.4);Streptophyta(13.4);Liliopsida(13.4);Poales(13.4);Poaceae(13.4);Zea(13.4);
329 Eukaryota(24.3);Chordata(12.3);Mammalia(12.3);Rodentia(12.3);Muridae(12.3);Mus(12.3);
330 Bacteria(90);Firmicutes(26);Clostridia(20.8);Clostridiales(19.1);Clostridiales_Incertae_Sedis_XI(4.2);Parvimonas(3);
331 Eukaryota(9.6);Streptophyta(8);Liliopsida(8);Poales(8);Poaceae(8);Zea(8);
332 Bacteria(87.2);"Bacteroidetes"(30.8);"Bacteroidia"(11.3);"Bacteroidales"(11.3);"Porphyromonadaceae"(6.9);Petrimonas(4.1);
333 Bacteria(100);Firmicutes(100);Erysipelotrichia(100);Erysipelotrichales(100);Erysipelotrichaceae(100);Turicibacter(100);
334 Bacteria(100);Firmicutes(100);Negativicutes(100);Selenomonadales(100);Veillonellaceae(100);Veillonella(100);
335 Bacteria(85.3);Firmicutes(19.9);Erysipelotrichia(7.8);Erysipelotrichales(7.8);Erysipelotrichaceae(7.8);Solobacterium(7.7);
336 Eukaryota(9.8);Streptophyta(8.4);Liliopsida(8.4);Poales(8.4);Poaceae(8.4);Zea(8.4);
337 Bacteria(100);"Tenericutes"(91.9);Mollicutes(91.9);Anaeroplasmatales(91.9);Anaeroplasmataceae(91.9);Anaeroplasmata(91.9);
338 Eukaryota(18.4);Chordata(7.2);Mammalia(7.2);Primates(2.3);Hominidae(2.3);Homo(2.3);
339 Bacteria(100);Firmicutes(100);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(87.4);Oscilibacter(86.7);
340 Bacteria(90.4);"Proteobacteria"(30.5);Alphaproteobacteria(14.6);Rhizobiales(5.2);Bradyrhizobiaceae(3.2);Agromonas(3);
341 Bacteria(84.4);"Bacteroidetes"(32);"Bacteroidia"(11.7);"Bacteroidales"(11.7);"Porphyromonadaceae"(6.4);Petrimonas(5);

342 Bacteria(86.6);"Bacteroidetes"(33.4);"Bacteroidia"(13.7);"Bacteroidales"(13.7);"Porphyromonadaceae"(6.9);Petrimonas(2.6);
343 Bacteria(87.2);"Proteobacteria"(24.8);Deltaproteobacteria(13.9);Desulfovibrionales(5.6);Desulfovibrionaceae(5.3);Bilophila(4.5);
344 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.8);Lachnospiraceae_incertae_sedis(54.7);
345 Eukaryota(11.3);Streptophyta(10.2);Liliopsida(10.2);Poales(10.2);Poaceae(10.2);Zea(10.2);
346 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(99.4);
347 Bacteria(100);Firmicutes(99.7);Clostridia(99.7);Clostridiales(99.7);Lachnospiraceae(97.7);Johnsonella(38.9);
348 Eukaryota(18);Chordata(7.6);Mammalia(7.6);Primates(1.7);Hominidae(1.7);Homo(1.7);
349 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Pseudoflavonifractor(64.8);
350 Bacteria(100);Firmicutes(99.5);Clostridia(99.5);Clostridiales(99.5);Lachnospiraceae(95.4);Acetitomaculum(30);
351 Bacteria(100);Firmicutes(43.2);Erysipelotrichia(43);Erysipelotrichales(43);Erysipelotrichaceae(43);Turicibacter(43);
352 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(96);Lachnospiraceae_incertae_sedis(26.8);
353 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Clostridium_XIVa(36.6);
354 Bacteria(81.5);"Proteobacteria"(36.1);Gammaproteobacteria(23.5);Chromatiales(8.8);Halothiobacillaceae(7.8);Thioalkalibacter(7.8);
355 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.3);Clostridium_XIVa(53.1);
356 Bacteria(100);"Bacteroidetes"(69.6);"Bacteroidia"(64.8);"Bacteroidales"(64.8);"Porphyromonadaceae"(61.5);Tannerella(27.9);
357 Bacteria(100);Firmicutes(77.5);Clostridia(77.3);Clostridiales(77.3);Lachnospiraceae(75.7);Lachnobacterium(47.9);
358 Bacteria(86.7);"Actinobacteria"(10.4);Actinobacteria(10.4);Actinomycetales(9.1);Micromonosporaceae(7.1);Catelliglobospora(6.4);
359 Bacteria(100);Firmicutes(100);Erysipelotrichia(100);Erysipelotrichales(100);Erysipelotrichaceae(100);Clostridium_XVIII(100);
360 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.5);Marvinbryantia(35.4);
361 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(94);Clostridium_XIVa(42.2);
362 Bacteria(100);Firmicutes(100);Bacilli(100);Lactobacillales(100);Streptococcaceae(100);Streptococcus(100);
363 Eukaryota(13.2);Streptophyta(11);Liliopsida(11);Poales(11);Poaceae(11);Zea(11);
364 Bacteria(100);Firmicutes(74.8);Clostridia(74.8);Clostridiales(74.8);Lachnospiraceae(73.7);Lachnobacterium(23.8);
365 Eukaryota(10.1);Streptophyta(8.2);Liliopsida(8.2);Poales(8.2);Poaceae(8.2);Zea(8.2);
366 Eukaryota(15.7);Chordata(7.2);Mammalia(7.2);Primates(2);Hominidae(2);Homo(2);
367 Bacteria(87.8);Firmicutes(31);Clostridia(26.4);Clostridiales(24.7);Clostridiales_Incertae_Sedis_XI(5.3);Parvimonas(4.2);
368 Eukaryota(13.8);Chordata(5.8);Mammalia(5.8);Rodentia(5.8);Muridae(5.8);Mus(5.8);
369 Archaea(26.2);"Euryarchaeota"(23.6);Halobacteria(12.3);Halobacteriales(12.3);Halobacteriaceae(12.3);Halonotius(4);
370 Archaea(20.5);"Euryarchaeota"(17.5);"Methanomicrobia"(7.9);Methanosarcinales(6);Methanosarcinaceae(5.7);Methanomicrococcus(5.6);
371 Archaea(25.1);"Euryarchaeota"(21.8);Halobacteria(11.3);Halobacteriales(11.3);Halobacteriaceae(11.3);Halonotius(3.5);
372 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);"Porphyromonadaceae"(100);Barnesiella(100);
373 Bacteria(89.4);"Deferribacteres"(7.4);Deferribacteres(7.4);Deferribacterales(7.4);Deferribacteraceae(7.4);Mucispirillum(6.7);
374 Bacteria(84.3);"Proteobacteria"(38.4);Alphaproteobacteria(13.9);Rhodobacterales(9.8);Rhodobacteraceae(9.8);Antarctobacter(0.9);
375 Bacteria(100);Firmicutes(100);Erysipelotrichia(77.5);Erysipelotrichales(77.5);Erysipelotrichaceae(77.5);Turicibacter(77.4);
376 Bacteria(100);Firmicutes(99.6);Clostridia(99.3);Clostridiales(99.3);Ruminococcaceae(91.1);Butyricoccus(87.6);
377 Bacteria(100);Firmicutes(99.1);Clostridia(99.1);Clostridiales(99.1);Lachnospiraceae(90.9);Syntrophococcus(18.7);
378 Bacteria(100);"Bacteroidetes"(91.6);Flavobacteria(91.5);"Flavobacteriales"(91.5);Flavobacteriaceae(90.8);Dokdonia(38);
379 Eukaryota(25.4);Chordata(15.6);Mammalia(15.6);Rodentia(15.6);Muridae(15.6);Mus(15.6);
380 Bacteria(100);Firmicutes(97.6);Clostridia(96.9);Clostridiales(96.7);Ruminococcaceae(35.8);Anaerotruncus(30.2);
381 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Coproccoccus(32.4);
382 Eukaryota(12.4);Streptophyta(11.2);Liliopsida(11.2);Poales(11.2);Poaceae(11.2);Zea(11.2);
383 Bacteria(100);"Bacteroidetes"(56.9);Flavobacteria(13.6);"Flavobacteriales"(13.6);Cryomorphaceae(12.4);Fluviicola(12.4);
384 Bacteria(100);Firmicutes(97.4);Clostridia(96.2);Clostridiales(96.1);Ruminococcaceae(90.5);Pseudoflavonifractor(52.2);
385 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(97);
386 Bacteria(82.6);"Proteobacteria"(14.2);Deltaproteobacteria(7.1);Syntrophobacterales(6.2);Syntrophobacteraceae(6.2);Thermodesulforhabdus(6.2);
387 Eukaryota(25.1);Chordata(24.3);Mammalia(24.3);Rodentia(18.1);Muridae(18.1);Mus(18.1);
388 Bacteria(100);Firmicutes(99.5);Clostridia(99.5);Clostridiales(99.5);Ruminococcaceae(99.5);Faecalibacterium(93.1);
389 Bacteria(86);"Bacteroidetes"(35.5);"Bacteroidia"(16.8);"Bacteroidales"(16.8);"Porphyromonadaceae"(8.3);Petrimonas(3.7);
390 Bacteria(100);Firmicutes(94.8);Erysipelotrichia(70.7);Erysipelotrichales(70.7);Erysipelotrichaceae(70.7);Turicibacter(70.5);
391 Eukaryota(14.5);Chordata(10.2);Mammalia(10.2);Rodentia(10.2);Muridae(10.2);Mus(10.2);
392 Eukaryota(29.3);Chordata(25.2);Mammalia(25.2);Rodentia(19.7);Muridae(19.7);Mus(19.7);
393 Bacteria(100);Firmicutes(62.4);Clostridia(62.3);Clostridiales(62.3);Lachnospiraceae(56.8);Lactonifactor(31.8);
394 Eukaryota(12.2);Chordata(5.8);Mammalia(5.8);Primates(2.2);Hominidae(2.2);Homo(2.2);
395 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Lachnospiraceae_incertae_sedis(93.9);
396 Bacteria(100);Firmicutes(85.3);Clostridia(83.7);Clostridiales(83.6);Lachnospiraceae(76);Clostridium_XIVb(70.7);
397 Bacteria(100);Firmicutes(82.5);Clostridia(82.5);Clostridiales(82.5);Lachnospiraceae(78.3);Oribacterium(32.9);
398 Bacteria(100);"Bacteroidetes"(99.9);"Bacteroidia"(91.1);"Bacteroidales"(91.1);"Porphyromonadaceae"(74.1);Barnesiella(53.9);
399 Bacteria(87.5);"Bacteroidetes"(31.1);"Bacteroidia"(12.6);"Bacteroidales"(12.6);"Porphyromonadaceae"(5.7);Petrimonas(3.3);
400 Bacteria(100);"Bacteroidetes"(99.9);"Bacteroidia"(94.5);"Bacteroidales"(94.5);"Porphyromonadaceae"(91.8);Barnesiella(72.1);
401 Eukaryota(5.4);Streptophyta(4.7);Liliopsida(4.7);Poales(4.7);Poaceae(4.7);Zea(4.7);
402 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(99.8);Oscillibacter(81.9);
403 Bacteria(100);Firmicutes(83.5);Clostridia(78.1);Clostridiales(77.5);Clostridiales_Incertae_Sedis_XII(34.1);Acidaminobacter(17.7);
404 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Actinomycetales(100);Microbacteriaceae(100);Microbacterium(100);
405 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Clostridium_XIVa(75.6);
406 Bacteria(100);Firmicutes(79.6);Clostridia(67.4);Clostridiales(67.4);Ruminococcaceae(35.1);Saccharofermentans(19.1);
407 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(98.9);"Bacteroidales"(98.9);"Porphyromonadaceae"(97.1);Porphyromonas(46.5);
408 Eukaryota(15.3);Streptophyta(12);Liliopsida(12);Poales(12);Poaceae(12);Zea(12);
409 Bacteria(100);Firmicutes(90.9);Clostridia(74);Clostridiales(70.7);Ruminococcaceae(31.4);Saccharofermentans(21.4);
410 Bacteria(100);Firmicutes(99.1);Clostridia(99.1);Clostridiales(99.1);Ruminococcaceae(98.9);Flavonifractor(74.2);
411 Bacteria(100);Firmicutes(72.4);Clostridia(72.4);Clostridiales(72.4);Clostridiaceae_1(71);Sarcina(70);
412 Archaea(16.5);"Euryarchaeota"(15.2);Halobacteria(11.3);Halobacteriales(11.3);Halobacteriaceae(11.3);Halolamina(3);
413 Bacteria(87.6);"Deferribacteres"(6.7);Deferribacteres(6.7);Deferribacterales(6.7);Deferribacteraceae(6.7);Mucispirillum(6.2);
414 Bacteria(100);Firmicutes(73);Clostridia(52);Clostridiales(51.4);Gracilibacteraceae(15.7);Gracilibacter(14.8);
415 Eukaryota(11);Arthropoda(5.4);Insecta(5.4);Coleoptera(5.4);Tenebrionidae(5.4);Tribolium(5.4);
416 Eukaryota(7.5);Arthropoda(5.2);Insecta(5.2);Coleoptera(5.2);Tenebrionidae(5.2);Tribolium(5.2);
417 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(66.1);Oscillibacter(54.8);
418 Bacteria(87.7);"Proteobacteria"(27.2);Deltaproteobacteria(9.4);Desulfovibrionales(2.5);Desulfovibrionaceae(1.7);Lawsonia(1.7);
419 Bacteria(90.6);Firmicutes(19.1);Bacilli(7.6);Lactobacillales(6.3);Enterococcaceae(5.6);Piliibacter(5.5);
420 Eukaryota(24.6);Chordata(13);Mammalia(13);Primates(2.7);Hominidae(2.7);Homo(2.7);
421 Bacteria(100);Firmicutes(91.2);Erysipelotrichia(68.8);Erysipelotrichales(68.8);Erysipelotrichaceae(68.8);Coproccoccus(25.7);
422 Eukaryota(18.5);Streptophyta(14.1);Liliopsida(14.1);Poales(14.1);Poaceae(14.1);Zea(14.1);
423 Bacteria(100);Firmicutes(89.1);Clostridia(89.1);Clostridiales(89.1);Lachnospiraceae(85.8);Marvinbryantia(40.1);
424 Bacteria(86.5);"Proteobacteria"(42);Gammaproteobacteria(22.6);Chromatiales(10.8);Halothiobacillaceae(10.4);Thioalkalibacter(10.2);
425 Eukaryota(24.7);Chordata(24.2);Mammalia(24.2);Rodentia(18.1);Muridae(18.1);Mus(18.1);
426 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(97);Coproccoccus(30.2);
427 Archaea(18.7);"Euryarchaeota"(15.9);Halobacteria(10.4);Halobacteriales(10.4);Halobacteriaceae(10.4);Halonotius(2.4);

428 Bacteria(100);Firmicutes(99.4);Clostridia(96.1);Clostridiales(96);Peptostreptococcaceae(74.1);Sporacetigenium(45.1);
429 Bacteria(100);Firmicutes(77.3);Clostridia(74.7);Clostridiales(74.5);Lachnospiraceae(44.6);Lactonifactor(13.7);
430 Bacteria(100);Firmicutes(36.4);Clostridia(36.4);Clostridiales(36.4);Clostridiaceae_1(35);Sarcina(34.4);
431 Bacteria(100);Firmicutes(99.6);Erysipelotrichia(52.8);Erysipelotrichales(52.8);Erysipelotrichaceae(52.8);Turicibacter(51.5);
432 Eukaryota(11);Chordata(6.6);Mammalia(6.6);Rodentia(6.6);Muridae(6.6);Mus(6.6);
433 Bacteria(100);Firmicutes(81.6);Clostridia(81.6);Clostridiales(81.6);Lachnospiraceae(71.4);Robinsoniella(38.4);
434 Archaea(24.9);"Euryarchaeota"(23.6);"Methanomicrobia"(10.9);Methanosarcinales(9.8);Methanosarcinaceae(9.6);Methanimicrococcus(9.4);
435 Bacteria(100);Firmicutes(96);Clostridia(96);Clostridiales(96);Ruminococcaceae(95.9);Anaerotruncus(40.5);
436 Bacteria(100);Firmicutes(82.9);Clostridia(82.9);Clostridiales(82.9);Lachnospiraceae(74.7);Dorea(63.8);
437 Bacteria(100);Firmicutes(100);Negativicutes(100);Selenomonadales(100);Veillonellaceae(100);Dialister(99.9);
438 Eukaryota(15.4);Arthropoda(7.5);Insecta(7.5);Coleoptera(7.5);Tenebrionidae(7.5);Tribolium(7.5);
439 Bacteria(100);Firmicutes(97.8);Clostridia(97.8);Clostridiales(97.8);Ruminococcaceae(97.7);Flavonifractor(37.7);
440 Bacteria(87.4);Firmicutes(34.3);Clostridia(23.1);Clostridiales(15.4);Lachnospiraceae(5.6);Cellulosilyticum(3);
441 Archaea(9.9);"Euryarchaeota"(7.8);Thermoplasmata(5.2);Thermoplasmatales(5.2);Ferropasmaceae(5.1);Ferroplasma(5.1);
442 Bacteria(100);"Proteobacteria"(100);Betaproteobacteria(100);Neisseriales(100);Neisseriaceae(100);Neisseria(100);
443 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(98.9);Coproccoccus(40.4);
444 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);"Prevotellaceae"(100);Prevotella(100);
445 Eukaryota(10.1);Arthropoda(6.4);Insecta(6.4);Coleoptera(6.4);Tenebrionidae(6.4);Tribolium(6.4);
446 Bacteria(100);"Bacteroidetes"(77);"Bacteroidia"(73.9);"Bacteroidales"(73.9);"Porphyromonadaceae"(72.9);Tannerella(35.8);
447 Eukaryota(25.5);Chordata(19.6);Mammalia(19.6);Rodentia(17.3);Muridae(17.3);Mus(17.3);
448 Bacteria(100);Firmicutes(96);Clostridia(96);Clostridiales(96);Lachnospiraceae(98.4);Coproccoccus(13.7);
449 Bacteria(100);"Bacteroidetes"(99.1);"Bacteroidia"(96.8);"Bacteroidales"(96.8);"Porphyromonadaceae"(83.3);Tannerella(49.5);
450 Eukaryota(14.5);Chordata(11.8);Mammalia(11.8);Rodentia(11.8);Muridae(11.8);Mus(11.8);
451 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(99.8);"Bacteroidales"(99.8);"Porphyromonadaceae"(98.5);Barnesiella(79.9);
452 Eukaryota(14.6);Arthropoda(9.3);Insecta(9.3);Coleoptera(9.3);Tenebrionidae(9.3);Tribolium(9.3);
453 Bacteria(100);"Bacteroidetes"(99.8);"Bacteroidia"(98.5);"Bacteroidales"(98.5);"Porphyromonadaceae"(96.2);Barnesiella(71.8);
454 Bacteria(100);Firmicutes(73.6);Clostridia(73.1);Clostridiales(73.1);Clostridiaceae_1(69.5);Anaerobacter(68.7);
455 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(98.7);"Bacteroidales"(98.7);"Porphyromonadaceae"(96.3);Barnesiella(73.4);
456 Bacteria(99.9);"Tenericutes"(41.2);Mollicutes(41.2);Anaeroplasmatales(40.9);Anaeroplasmataceae(40.9);Asteroleplasma(40.9);
457 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(77.5);Roseburia(31.4);
458 Archaea(17.3);"Euryarchaeota"(14.5);"Methanomicrobia"(7.6);Methanosarcinales(5.8);Methanosarcinaceae(5.6);Methanimicrococcus(5.6);
459 Eukaryota(6.6);Arthropoda(4.6);Insecta(4.6);Coleoptera(4.6);Tenebrionidae(4.6);Tribolium(4.6);
460 Bacteria(86.7);"Thermodesulfobacteria"(5.8);Thermodesulfobacteria(5.8);Thermodesulfobacteriales(5.8);Thermodesulfobacteriaceae(5.8);Caldimicrobium(5.8);
461 Eukaryota(26.5);Chordata(25.2);Mammalia(25.2);Rodentia(19.4);Muridae(19.4);Mus(19.4);
462 Bacteria(100);"Bacteroidetes"(99.9);"Bacteroidia"(97.5);"Bacteroidales"(97.5);"Porphyromonadaceae"(94.2);Barnesiella(71.3);
463 Bacteria(85.3);"Bacteroidetes"(37.6);"Bacteroidia"(18.1);"Bacteroidales"(18.1);"Porphyromonadaceae"(11.8);Proteiniphilum(8.8);
464 Archaea(20.3);"Euryarchaeota"(19);Thermoplasmata(7.3);Thermoplasmatales(7.3);Ferropasmatales(7.3);Ferroplasma(5.7);
465 Eukaryota(11.8);Streptophyta(10.3);Liliopsida(10.3);Poales(10.3);Poaceae(10.3);Zea(10.3);
466 Bacteria(99.6);"Actinobacteria"(70.3);Actinobacteria(70.3);Coriobacteriales(63.6);Coriobacteriaceae(63.6);Asaccharobacter(28.5);
467 Eukaryota(24.2);Chordata(22);Mammalia(22);Rodentia(16.6);Muridae(16.6);Mus(16.6);
468 Eukaryota(8.7);Arthropoda(4.9);Insecta(4.9);Coleoptera(4.9);Tenebrionidae(4.9);Tribolium(4.9);
469 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.1);Coproccoccus(28.4);
470 Bacteria(83.6);"Proteobacteria"(36.5);Gammaproteobacteria(23.4);Chromatiales(9.9);Halothiobacillaceae(9.7);Thioalkalibacter(9.6);
471 Eukaryota(23.2);Chordata(23);Mammalia(23);Rodentia(17.1);Muridae(17.1);Mus(17.1);
472 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Flavonifractor(62.6);
473 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Clostridium_XlVa(58.6);
474 Eukaryota(15.7);Arthropoda(11.5);Insecta(11.5);Coleoptera(11.5);Tenebrionidae(11.5);Tribolium(11.5);
475 Bacteria(100);Firmicutes(97.7);Clostridia(97.2);Clostridiales(97);Ruminococcaceae(49.5);Anaerotruncus(20.7);
476 Bacteria(81.1);"Proteobacteria"(27.7);Gammaproteobacteria(12.5);"Vibrionales"(5.9);Vibrionaceae(5.9);Grimontia(5.9);
477 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(82.5);Ruminococcus(82.2);
478 Bacteria(86);"Bacteroidetes"(30.4);"Bacteroidia"(9.9);"Bacteroidales"(9.9);"Porphyromonadaceae"(5.7);Petrimonas(3.2);
479 Eukaryota(7.2);Streptophyta(6.1);Liliopsida(6.1);Poales(6.1);Poaceae(6.1);Zea(6.1);
480 Bacteria(90.2);"Proteobacteria"(31.4);Betaproteobacteria(9.9);Neisseriales(7.8);Neisseriaceae(7.8);Stenoxybacter(3.5);
481 Eukaryota(13.4);Streptophyta(12.4);Liliopsida(12.4);Poales(12.4);Poaceae(12.4);Zea(12.4);
482 Eukaryota(6.2);Arthropoda(3.8);Insecta(3.8);Coleoptera(3.8);Tenebrionidae(3.8);Tribolium(3.8);
483 Bacteria(78.6);Firmicutes(22.7);Clostridia(17.8);Clostridiales(15.6);Clostridiales_Incertae_Sedis_XI(7.1);Parvimonas(6.4);
484 Bacteria(100);Firmicutes(97.6);Clostridia(95.2);Clostridiales(94.9);Lachnospiraceae(62.1);Marvinbryantia(21.9);
485 Eukaryota(14.5);Streptophyta(12.2);Liliopsida(12.2);Poales(12.2);Poaceae(12.2);Zea(12.2);
486 Bacteria(100);Firmicutes(97.5);Clostridia(97.5);Clostridiales(97.5);Lachnospiraceae(97.3);Robinsoniella(78.5);
487 Eukaryota(13.5);Chordata(7.1);Mammalia(7.1);Primates(2.3);Hominidae(2.3);Homo(2.3);
488 Bacteria(100);Firmicutes(98.7);Clostridia(98.7);Clostridiales(98.7);Lachnospiraceae(90.7);Syntrophococcus(27.3);
489 Archaea(17);"Euryarchaeota"(10.8);Thermoplasmata(8);Thermoplasmatales(8);Thermoplasmatales_incertae_sedis(7.4);Thermogymnomonas(7.4);
490 Bacteria(81);"Lentisphaerae"(7.3);"Lentisphaeria"(7.3);Lentisphaerales(7.3);"Lentisphaeraceae"(7.3);Lentisphaera(7.3);
491 Eukaryota(16.5);Streptophyta(14.6);Liliopsida(14.6);Poales(14.6);Poaceae(14.6);Zea(14.6);
492 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(98.6);"Bacteroidales"(98.6);"Porphyromonadaceae"(97.6);Barnesiella(64.3);
493 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);Bacteroidaceae(100);Bacteroides(100);
494 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(94);
495 Bacteria(100);Firmicutes(90.6);Erysipelotrichia(79.5);Erysipelotrichales(79.5);Erysipelotrichaceae(79.5);Sharpea(20.3);
496 Archaea(19.8);"Euryarchaeota"(17);"Methanomicrobia"(9.9);Methanosarcinales(7.7);Methanosarcinaceae(7.7);Methanolobium(4.5);
497 Eukaryota(13.3);Streptophyta(10.6);Liliopsida(10.6);Poales(10.6);Poaceae(10.6);Zea(10.6);
498 Eukaryota(22.3);Chordata(12.5);Mammalia(12.5);Primates(3.1);Hominidae(3.1);Homo(3.1);
499 Bacteria(85.2);"Proteobacteria"(37.6);Gammaproteobacteria(16.5);Legionellales(5.5);Coxiellaceae(5.5);Coxiella(5.5);
500 Eukaryota(11.3);Streptophyta(9.8);Liliopsida(9.8);Poales(9.8);Poaceae(9.8);Zea(9.8);
501 Eukaryota(19.6);Streptophyta(18);Liliopsida(18);Poales(18);Poaceae(18);Zea(18);
502 Eukaryota(13.7);Chordata(9.9);Mammalia(9.9);Rodentia(9.9);Muridae(9.9);Mus(9.9);
503 Bacteria(82.1);"Bacteroidetes"(30);"Sphingobacteria"(16.4);"Sphingobacteriales"(16.4);"Cyclobacteriaceae"(8.4);Fontibacter(7.2);
504 Eukaryota(6);Arthropoda(4);Insecta(4);Coleoptera(4);Tenebrionidae(4);Tribolium(4);
505 Bacteria(84.3);"Bacteroidetes"(21.6);"Sphingobacteria"(14.9);"Sphingobacteriales"(14.9);Cytophagaceae(9.9);Leadbetterella(4.8);
506 Bacteria(100);Firmicutes(99.9);Clostridia(69.9);Clostridiales(69.9);Lachnospiraceae(67.6);Syntrophococcus(28.2);
507 Bacteria(91.2);"Proteobacteria"(27.7);Alphaproteobacteria(13.1);Rhizobiales(4.4);Bradyrhizobiaceae(2.7);Agromonas(2.7);
508 Bacteria(88);"Bacteroidetes"(26.1);"Sphingobacteria"(15.8);"Sphingobacteriales"(15.8);Sphingobacteriaceae(4.3);Pseudosphingobacterium(3.2);
509 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(53.4);Robinsoniella(23.2);
510 Eukaryota(26.6);Chordata(16.2);Mammalia(16.2);Rodentia(13.4);Muridae(13.4);Mus(13.4);
511 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(75);
512 Bacteria(85.2);"Proteobacteria"(40.9);Gammaproteobacteria(25.3);Chromatiales(13.2);Halothiobacillaceae(11.8);Thioalkalibacter(11.7);
513 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Pseudoflavonifractor(74.5);

514 Bacteria(92.8);"Proteobacteria"(31.1);Alphaproteobacteria(13.3);Rhizobiales(5.5);Bradyrhizobiaceae(3.2);Agromonas(3.1);
515 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(93.2);Robinsoniella(29.5);
516 Bacteria(89.9);Firmicutes(31.4);Clostridia(25.8);Clostridiales(23.4);Clostridiales_Incertae_Sedis_XI(5.8);Parvimonas(4.6);
517 Bacteria(100);Firmicutes(99.7);Clostridia(99.7);Clostridiales(99.7);Ruminococcaceae(69.6);Oscillibacter(60.7);
518 Bacteria(100);Firmicutes(96.2);Clostridia(96);Clostridiales(96);Lachnospiraceae(80.8);Robinsoniella(30.8);
519 Bacteria(100);Firmicutes(61.6);Clostridia(61.5);Clostridiales(61.5);Lachnospiraceae(60.1);Shuttleworthia(30.2);
520 Bacteria(100);Firmicutes(56.2);Clostridia(48.6);Clostridiales(47.6);Ruminococcaceae(24);Acetivibrio(12.7);
521 Eukaryota(12);Streptophyta(7.4);Liliopsida(7.4);Poales(7.4);Poaceae(7.4);Zea(7.4);
522 Eukaryota(23.9);Chordata(16.1);Mammalia(16.1);Rodentia(16.1);Muridae(16.1);Mus(16.1);
523 Eukaryota(20.3);Chordata(18.9);Mammalia(18.9);Rodentia(15.8);Muridae(15.8);Mus(15.8);
524 Bacteria(87.7);"Proteobacteria"(28.5);Deltaproteobacteria(7.9);Desulfobacterales(3.5);Desulfobacteraceae(2);Desulfatiferula(1.8);
525 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);"Prevotellaceae"(99.9);Prevotella(82.1);
526 Bacteria(100);"Tenericutes"(98);Mollicutes(98);Anaeroplasmatales(65.5);Anaeroplasmataceae(65.5);Anaeroplasma(65.5);
527 Eukaryota(9.7);Arthropoda(6.3);Insecta(6.3);Coleoptera(6.3);Tenebrionidae(6.3);Tribolium(6.3);
528 Eukaryota(27.2);Chordata(24.5);Mammalia(24.5);Rodentia(18.6);Muridae(18.6);Mus(18.6);
529 Bacteria(100);Firmicutes(100);Clostridia(94.5);Clostridiales(94.5);Lachnospiraceae(90.6);Coprococcus(46.1);
530 Bacteria(100);Firmicutes(98.3);Erysipelotrichia(98.3);Erysipelotrichales(98.3);Erysipelotrichaceae(98.3);Clostridium_XVIII(94.2);
531 Bacteria(100);Firmicutes(60.7);Clostridia(60.7);Clostridiales(60.7);Lachnospiraceae(55);Marvinbryantia(12.2);
532 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(91.5);Clostridium_XIVa(31.2);
533 Bacteria(100);Firmicutes(99.7);Clostridia(99.7);Clostridiales(99.7);Ruminococcaceae(99.7);Oscillibacter(87.5);
534 Bacteria(88.4);"Bacteroidetes"(31.6);"Bacteroidia"(15.3);"Bacteroidales"(15.3);"Porphyromonadaceae"(8.4);Proteiniphilum(4.2);
535 Bacteria(100);"Bacteroidetes"(99.1);"Bacteroidia"(95.4);"Bacteroidales"(95.4);"Porphyromonadaceae"(83.4);Tannerella(38.5);
536 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(99.7);Clostridium_IV(63.1);
537 Eukaryota(6.1);Arthropoda(4.2);Insecta(4.2);Coleoptera(4.2);Tenebrionidae(4.2);Tribolium(4.2);
538 Eukaryota(15.4);Arthropoda(8.5);Insecta(8.5);Coleoptera(8.5);Tenebrionidae(8.5);Tribolium(8.5);
539 Bacteria(100);Firmicutes(99.8);Clostridia(99.8);Clostridiales(99.8);Ruminococcaceae(85.9);Anaerotruncus(35.1);
540 Bacteria(100);Firmicutes(99.7);Clostridia(99.6);Clostridiales(99.6);Lachnospiraceae(86.8);Moryella(23.6);
541 Bacteria(100);Firmicutes(86.7);Clostridia(86.7);Clostridiales(86.7);Lachnospiraceae(86.7);Syntrophococcus(30.2);
542 Bacteria(100);Firmicutes(85.6);Clostridia(85.5);Clostridiales(85.5);Lachnospiraceae(78.1);Marvinbryantia(32.2);
543 Bacteria(100);Firmicutes(93.9);Clostridia(84.8);Clostridiales(84.7);Peptococcaceae_1(48.6);Dehalobacter(45.9);
544 Archaea(16.4);"Euryarchaeota"(14.4);Halobacteria(7.4);Halobacteriales(7.4);Halobacteriaceae(7.4);Halonotus(3);
545 Bacteria(92.3);"Proteobacteria"(33.5);Alphaproteobacteria(14.5);Rhizobiales(5);Bradyrhizobiaceae(2.7);Agromonas(2.7);
546 Eukaryota(14.4);Chordata(11.5);Mammalia(11.5);Rodentia(11.5);Muridae(11.5);Mus(11.5);
547 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.6);Pseudobutyrvibrio(40.2);
548 Bacteria(100);Firmicutes(100);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(76.6);Clostridium_IV(51.1);
549 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(99.8);Oscillibacter(86.4);
550 Bacteria(100);Firmicutes(99.8);Clostridia(99.8);Clostridiales(99.6);Ruminococcaceae(92.9);Oscillibacter(76.1);
551 Bacteria(100);"Bacteroidetes"(49.1);"Bacteroidia"(44.8);"Bacteroidales"(44.8);"Porphyromonadaceae"(44.1);Barnesiella(42.3);
552 Bacteria(86.9);"Actinobacteria"(9.7);Actinobacteria(9.7);Actinomycetales(8.6);Micromonosporaceae(6.9);Catelliglobospora(6.5);
553 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);"Rikenellaceae"(100);Alistipes(100);
554 Bacteria(100);Firmicutes(53.5);Clostridia(53.5);Clostridiales(53.5);Ruminococcaceae(53.5);Clostridium_IV(34.6);
555 Eukaryota(26.8);Chordata(23.9);Mammalia(23.9);Rodentia(17.9);Muridae(17.9);Mus(17.9);
556 Bacteria(90.3);"Proteobacteria"(31.9);Alphaproteobacteria(13.4);Rhizobiales(7.2);Bradyrhizobiaceae(4.8);Agromonas(4.8);
557 Bacteria(100);Firmicutes(88.3);Clostridia(83.1);Clostridiales(82.6);Clostridiales_Incertae_Sedis_XII(41.2);Acidaminobacter(32.9);
558 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Acetanaerobacterium(82.7);
559 Eukaryota(8);Streptophyta(5.9);Liliopsida(5.9);Poales(5.9);Poaceae(5.9);Zea(5.9);
560 Bacteria(85.6);"Bacteroidetes"(32.4);"Bacteroidia"(15);"Bacteroidales"(15);"Porphyromonadaceae"(6.3);Petrimonas(4.4);
561 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(97.9);"Bacteroidales"(97.9);"Porphyromonadaceae"(97);Barnesiella(73.1);
562 Bacteria(100);"Bacteroidetes"(96);"Bacteroidia"(87.9);"Bacteroidales"(87.9);"Porphyromonadaceae"(87.1);Barnesiella(64.3);
563 Eukaryota(15.4);Streptophyta(5.7);Liliopsida(5.7);Poales(5.7);Poaceae(5.7);Zea(5.7);
564 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(95.8);Lachnospiraceae_incertain_sedis(52.6);
565 Bacteria(100);Firmicutes(98.6);Clostridia(98.5);Clostridiales(98.4);Ruminococcaceae(85.7);Anaerotruncus(34.9);
566 Bacteria(81.7);"Proteobacteria"(40.6);Gammaproteobacteria(23.7);Chromatiales(11.2);Halothiobacillaceae(11.2);Thioalkalibacter(11);
567 Bacteria(100);"Bacteroidetes"(75.3);"Bacteroidia"(73.3);"Bacteroidales"(73.3);"Porphyromonadaceae"(67.8);Tannerella(35.1);
568 Eukaryota(27.7);Arthropoda(14.7);Insecta(14.7);Coleoptera(14.7);Tenebrionidae(14.7);Tribolium(14.7);
569 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(95.4);Papillibacter(28.6);
570 Bacteria(100);Firmicutes(98.8);Clostridia(98.8);Clostridiales(98.8);Ruminococcaceae(98.7);Flavonifractor(84.9);
571 Bacteria(100);Firmicutes(96);Clostridia(58.3);Clostridiales(58.2);Ruminococcaceae(49.6);Oscillibacter(33.8);
572 Bacteria(100);Firmicutes(100);Clostridia(98.2);Clostridiales(98.2);Lachnospiraceae(97);Coprococcus(38.2);
573 Eukaryota(8.2);Streptophyta(6.2);Liliopsida(6.2);Poales(6.2);Poaceae(6.2);Zea(6.2);
574 Bacteria(90.3);"Proteobacteria"(37.2);Alphaproteobacteria(7.5);Rickettsiales(4.5);Anaplasmataceae(4.5);Wolbachia(4.4);
575 Bacteria(100);Firmicutes(96.9);Clostridia(83.9);Clostridiales(83);Ruminococcaceae(48.4);Pseudoflavonifractor(14.1);
576 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(98.8);Oscillibacter(87.6);
577 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(98);Ruminococcus(52.1);
578 Bacteria(100);"Bacteroidetes"(64.3);"Bacteroidia"(58);"Bacteroidales"(58);"Porphyromonadaceae"(51);Tannerella(29.1);
579 Eukaryota(22.2);Streptophyta(19.7);Liliopsida(19.7);Poales(19.7);Poaceae(19.7);Zea(19.7);
580 Bacteria(100);Firmicutes(99.8);Clostridia(99.8);Clostridiales(99.8);Lachnospiraceae(97.5);Marvinbryantia(38.5);
581 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Lachnospiraceae(99.9);Roseburia(60.8);
582 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Clostridiaceae_1(100);Clostridium_sensu_stricto(87.9);
583 Eukaryota(11.1);Streptophyta(8.9);Liliopsida(8.9);Poales(8.9);Poaceae(8.9);Zea(8.9);
584 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(74.8);Lachnospira(19.5);
585 Bacteria(82.2);"Proteobacteria"(25.2);Deltaproteobacteria(11.3);Myxococcales(8.1);Nannocystaceae(7.7);Enhygromyxa(7.3);
586 Bacteria(100);Firmicutes(82);Erysipelotrichia(79.3);Erysipelotrichales(79.3);Erysipelotrichaceae(79.3);Holdemania(61.3);
587 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.4);Clostridium_XIVa(40.1);
588 Bacteria(100);Firmicutes(95.3);Erysipelotrichia(95.3);Erysipelotrichales(95.3);Erysipelotrichaceae(95.3);Holdemania(65);
589 Bacteria(100);Firmicutes(99.9);Clostridia(99.8);Clostridiales(99.8);Clostridiaceae_1(99.8);Clostridium_sensu_stricto(84);
590 Bacteria(87.7);"Bacteroidetes"(32.7);"Bacteroidia"(12.9);"Bacteroidales"(12.9);"Porphyromonadaceae"(6.4);Proteiniphilum(3.1);
591 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Lachnospiraceae(85.4);Coprococcus(33.5);
592 Bacteria(100);"Bacteroidetes"(99.9);"Bacteroidia"(99.2);"Bacteroidales"(99.2);"Porphyromonadaceae"(94.8);Tannerella(28.9);
593 Eukaryota(9.5);Arthropoda(5.3);Insecta(5.3);Coleoptera(5.3);Tenebrionidae(5.3);Tribolium(5.3);
594 Eukaryota(10.2);Streptophyta(6.3);Liliopsida(6.3);Poales(6.3);Poaceae(6.3);Zea(6.3);
595 Bacteria(100);Firmicutes(86);Clostridia(85.8);Clostridiales(85.8);Ruminococcaceae(81.9);Oscillibacter(41.1);
596 Bacteria(99.9);"Bacteroidetes"(64.2);"Bacteroidia"(46.2);"Bacteroidales"(46.2);"Porphyromonadaceae"(42.9);Barnesiella(35.8);
597 Bacteria(84.8);"Actinobacteria"(9.1);Actinobacteria(9.1);Actinomycetales(8.4);Micromonosporaceae(6.9);Catelliglobospora(6.5);
598 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Clostridiaceae_1(100);Clostridium_sensu_stricto(81.9);
599 Bacteria(100);Firmicutes(90.7);Clostridia(90.7);Clostridiales(90.7);Ruminococcaceae(90.7);Papillibacter(36.6);

600 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(98.7);Coproccoccus(19.5);
601 Eukaryota(10.8);Streptophyta(9.2);Liliopsida(9.2);Poales(9.2);Poaceae(9.2);Zea(9.2);
602 Bacteria(87.6);"Bacteroidetes"(28.8);"Bacteroidia"(11.8);"Bacteroidales"(11.8);"Porphyromonadaceae"(4.8);Petrimonas(3.2);
603 Bacteria(84.1);"Bacteroidetes"(26.2);"Sphingobacteria"(11.1);"Sphingobacteriales"(11.1);"Flammeovirgaceae"(6.4);Flexithrix(5.5);
604 Bacteria(100);Firmicutes(62.5);Clostridia(61.6);Clostridiales(61.2);Ruminococcaceae(50.4);Flavonifractor(18.9);
605 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Anaerotruncus(99.8);
606 Bacteria(100);Firmicutes(99.8);Clostridia(85.5);Clostridiales(85.5);Lachnospiraceae(84.5);Johnsonella(51.7);
607 Bacteria(87.6);"Bacteroidetes"(32.7);"Bacteroidia"(11.8);"Bacteroidales"(11.8);"Porphyromonadaceae"(7.5);Petrimonas(4.4);
608 Eukaryota(9);Streptophyta(8.5);Liliopsida(8.5);Poales(8.5);Poaceae(8.5);Zea(8.5);
609 Bacteria(100);Firmicutes(98.5);Clostridia(94.2);Clostridiales(94.2);Lachnospiraceae(81.4);Acetitomaculum(19.8);
610 Eukaryota(21.8);Chordata(21.3);Mammalia(21.3);Rodentia(16);Muridae(16);Mus(16);
611 Bacteria(86.5);"Bacteroidetes"(31.7);"Bacteroidia"(13.1);"Bacteroidales"(13.1);"Porphyromonadaceae"(5.4);Petrimonas(3.5);
612 Bacteria(86.4);"Proteobacteria"(39.5);Alphaproteobacteria(12.3);Rhodobacterales(8.9);Rhodobacteraceae(8.9);Nereida(3.5);
613 Bacteria(100);"Bacteroidetes"(58.2);"Bacteroidia"(48.5);"Bacteroidales"(48.5);"Porphyromonadaceae"(42.7);Tannerella(27.5);
614 Archaea(17.7);"Euryarchaeota"(13.6);Thermoplasmata(5.5);Thermoplasmatales(5.5);Thermoplasmatales_incertae_sedis(3.3);Thermogymnomonas(3.3);
615 Bacteria(100);"Bacteroidetes"(61);"Bacteroidia"(55.2);"Bacteroidales"(55.2);"Porphyromonadaceae"(45.8);Tannerella(34.9);
616 Bacteria(83.9);"Bacteroidetes"(27.3);Flavobacteria(19.9);"Flavobacteriales"(19.9);Flavobacteriaceae(19.4);Joostella(8.5);
617 Archaea(8.9);"Euryarchaeota"(7.7);Thermoplasmata(5.3);Thermoplasmatales(5.3);Ferroplasma(5.2);Ferroplasma(5.2);
618 Bacteria(100);Firmicutes(87.6);Clostridia(85.2);Clostridiales(85.1);Ruminococcaceae(80.3);Anaerotruncus(27.4);
619 Eukaryota(16.7);Streptophyta(9.5);Liliopsida(9.5);Poales(9.5);Poaceae(9.5);Zea(9.5);
620 Eukaryota(8.2);Arthropoda(7.3);Insecta(7.3);Coleoptera(7.3);Tenebrionidae(7.3);Tribolium(7.3);
621 Eukaryota(14.4);Chordata(10.1);Mammalia(10.1);Rodentia(10.1);Muridae(10.1);Mus(10.1);
622 Eukaryota(11.9);Arthropoda(8);Insecta(8);Coleoptera(8);Tenebrionidae(8);Tribolium(8);
623 Eukaryota(12.7);Arthropoda(8.2);Insecta(8.2);Coleoptera(8.2);Tenebrionidae(8.2);Tribolium(8.2);
624 Bacteria(88);"Bacteroidetes"(31.6);"Bacteroidia"(11);"Bacteroidales"(11);"Porphyromonadaceae"(4.6);Petrimonas(2.2);
625 Archaea(26.4);"Euryarchaeota"(25.6);"Methanomicrobia"(12.4);Methanomicrobiales(12.4);Methanomicrobiaceae(7.8);Methanosphaerula(5.8);
626 Archaea(24.4);"Euryarchaeota"(19.8);Halobacteria(14);Halobacteriales(14);Halobacteriaceae(14);Halonotius(6.6);
627 Eukaryota(11.9);Streptophyta(7.1);Liliopsida(7.1);Poales(7.1);Poaceae(7.1);Zea(7.1);
628 Bacteria(88.2);Firmicutes(28.4);Clostridia(21.6);Clostridiales(19.1);Clostridiales_Incertae_Sedis_XI(4.3);Parvimonas(3);
629 Eukaryota(10.4);Chordata(6.2);Mammalia(6.2);Rodentia(6.2);Muridae(6.2);Mus(6.2);
630 Bacteria(100);Firmicutes(99.6);Clostridia(99.6);Clostridiales(99.3);Ruminococcaceae(71.4);Flavonifractor(30.4);
631 Eukaryota(21.2);Chordata(10.9);Mammalia(10.9);Rodentia(8.2);Muridae(8.2);Mus(8.2);
632 Eukaryota(8.3);Arthropoda(6.4);Insecta(6.4);Coleoptera(6.4);Tenebrionidae(6.4);Tribolium(6.4);
633 Bacteria(100);Firmicutes(94.1);Clostridia(94.1);Clostridiales(94.1);Lachnospiraceae(85.7);Parasporobacterium(52.2);
634 Bacteria(100);"Bacteroidetes"(58.5);"Bacteroidia"(43.3);"Bacteroidales"(43.3);"Porphyromonadaceae"(35.8);Petrimonas(15.2);
635 Bacteria(100);Firmicutes(100);Clostridia(72.1);Clostridiales(72.1);Ruminococcaceae(71.7);Acetanaerobacterium(17.7);
636 Bacteria(100);Firmicutes(25.5);Clostridia(100);Clostridiales(100);Ruminococcaceae(99.8);Oscillibacter(63.3);
637 Bacteria(100);Firmicutes(83.1);Clostridia(83.1);Clostridiales(83.1);Ruminococcaceae(81.7);Clostridium_IV(39.9);
638 Eukaryota(26.4);Chordata(23.2);Mammalia(23.2);Rodentia(19.8);Muridae(19.8);Mus(19.8);
639 Eukaryota(10.9);Streptophyta(9.6);Liliopsida(9.6);Poales(9.6);Poaceae(9.6);Zea(9.6);
640 Bacteria(100);"Bacteroidetes"(41.2);"Bacteroidia"(40.7);"Bacteroidales"(40.7);Bacteroidaceae(40.7);Anaerorhabdus(40.7);
641 Bacteria(85.7);Firmicutes(25.5);Clostridia(22.2);Clostridiales(20.8);Clostridiales_Incertae_Sedis_XI(7);Parvimonas(5.7);
642 Eukaryota(12.6);Chordata(7.7);Mammalia(7.7);Primates(2.9);Hominidae(2.9);Homo(2.9);
643 Bacteria(100);Firmicutes(85.8);Clostridia(80.7);Clostridiales(80.7);Ruminococcaceae(78.3);Acetivibrio(49.3);
644 Archaea(18.6);"Euryarchaeota"(15.3);"Methanomicrobia"(15.3);Methanosarcinales(12.3);Methanosarcinaceae(5.2);Methanohalobium(3.4);
645 Bacteria(100);Firmicutes(97);Clostridia(96.5);Clostridiales(96.5);Ruminococcaceae(52.5);Ethanoligenens(17);
646 Eukaryota(12.9);Streptophyta(11.7);Liliopsida(11.7);Poales(11.7);Poaceae(11.7);Zea(11.7);
647 Eukaryota(11.7);Arthropoda(8.1);Insecta(8.1);Coleoptera(8.1);Tenebrionidae(8.1);Tribolium(8.1);
648 Bacteria(86.3);"Bacteroidetes"(31.6);"Bacteroidia"(11.8);"Bacteroidales"(11.8);"Porphyromonadaceae"(7.4);Petrimonas(5.6);
649 Eukaryota(22.2);Chordata(8.7);Mammalia(8.7);Primates(1.5);Hominidae(1.5);Homo(1.5);
650 Eukaryota(15.8);Arthropoda(9.3);Insecta(9.3);Coleoptera(9.3);Tenebrionidae(9.3);Tribolium(9.3);
651 Eukaryota(11.7);Streptophyta(10.4);Liliopsida(10.4);Poales(10.4);Poaceae(10.4);Zea(10.4);
652 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Lachnospiraceae_incertae_sedis(60.6);
653 Eukaryota(10.3);Arthropoda(5.2);Insecta(5.2);Coleoptera(5.2);Tenebrionidae(5.2);Tribolium(5.2);
654 Bacteria(85.7);"Bacteroidetes"(22.7);"Sphingobacteria"(12.6);"Sphingobacteriales"(12.6);Sphingobacteriaceae(3.4);Pseudosphingobacterium(2.2);
655 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(85.4);Oscillibacter(85.3);
656 Bacteria(100);Firmicutes(99.3);Clostridia(99.3);Clostridiales(99.3);Lachnospiraceae(90.2);Anaerostipes(38.9);
657 Eukaryota(43.6);Chordata(39.8);Mammalia(39.8);Rodentia(30.8);Muridae(30.8);Mus(30.8);
658 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Hespellia(38.1);
659 Bacteria(84.9);"Elusimicrobia"(4.8);Elusimicrobiales(4.8);Elusimicrobiaceae(4.8);Elusimicrobium(4.8);
660 Archaea(25);"Euryarchaeota"(22.3);"Methanomicrobia"(16.5);Methanosarcinales(12.3);Methanosarcinaceae(12.3);Methanohalobium(9.1);
661 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(71.8);Anaerotruncus(66.6);
662 Eukaryota(18.9);Streptophyta(9.8);Liliopsida(9.8);Poales(9.8);Poaceae(9.8);Zea(9.8);
663 Bacteria(87.2);"Bacteroidetes"(35.7);"Bacteroidia"(16);"Bacteroidales"(16);"Porphyromonadaceae"(8.7);Petrimonas(4.9);
664 Eukaryota(12.4);Streptophyta(9.5);Liliopsida(9.5);Poales(9.5);Poaceae(9.5);Zea(9.5);
665 Eukaryota(12.5);Streptophyta(9.4);Liliopsida(9.4);Poales(9.4);Poaceae(9.4);Zea(9.4);
666 Eukaryota(17);Chordata(12.9);Mammalia(12.9);Rodentia(12.9);Muridae(12.9);Mus(12.9);
667 Eukaryota(7.4);Arthropoda(3.6);Insecta(3.6);Coleoptera(3.6);Tenebrionidae(3.6);Tribolium(3.6);
668 Bacteria(100);Firmicutes(99.1);Clostridia(99.1);Clostridiales(99.1);Lachnospiraceae(95.1);Marvinbryantia(16);
669 Bacteria(100);Firmicutes(93.3);Clostridia(93.3);Clostridiales(93.3);Lachnospiraceae(90.6);Lactonifactor(40);
670 Eukaryota(9.7);Streptophyta(8.1);Liliopsida(8.1);Poales(8.1);Poaceae(8.1);Zea(8.1);
671 Eukaryota(15);Streptophyta(13.2);Liliopsida(13.2);Poales(13.2);Poaceae(13.2);Zea(13.2);
672 Bacteria(100);Firmicutes(99.6);Clostridia(99.6);Clostridiales(99.6);Lachnospiraceae(98.6);Lactonifactor(16.2);
673 Bacteria(100);Firmicutes(98.6);Erysipelotrichia(92.5);Erysipelotrichales(92.5);Erysipelotrichaceae(92.5);Coproccoccus(63.9);
674 Eukaryota(11.8);Arthropoda(6.7);Insecta(6.7);Coleoptera(6.7);Tenebrionidae(6.7);Tribolium(6.7);
675 Bacteria(100);Firmicutes(98.4);Clostridia(96.3);Clostridiales(96.3);Ruminococcaceae(86.7);Ethanoligenens(26.9);
676 Eukaryota(9.8);Streptophyta(8.3);Liliopsida(8.3);Poales(8.3);Poaceae(8.3);Zea(8.3);
677 Bacteria(99.9);Firmicutes(65.8);Clostridia(60.9);Clostridiales(60.6);Clostridiales_Incertae_Sedis_XII(24);Fusibacter(17.9);
678 Eukaryota(11.6);Streptophyta(4.2);Liliopsida(4.2);Poales(4.2);Poaceae(4.2);Zea(4.2);
679 Bacteria(100);Firmicutes(99.8);Clostridia(99.8);Clostridiales(99.8);Ruminococcaceae(68.8);Sporobacter(39.2);
680 Eukaryota(12.4);Chordata(9.3);Mammalia(9.3);Rodentia(9.3);Muridae(9.3);Mus(9.3);
681 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.6);Lachnospiraceae_incertae_sedis(67.1);
682 Eukaryota(26.4);Chordata(25.8);Mammalia(25.8);Rodentia(19.2);Muridae(19.2);Mus(19.2);
683 Bacteria(86);"Elusimicrobia"(3.2);Elusimicrobia(3.2);Elusimicrobiales(3.2);Elusimicrobiaceae(3.2);Elusimicrobium(3.2);
684 Eukaryota(10.5);Chordata(8.5);Mammalia(8.5);Rodentia(7.6);Muridae(7.6);Mus(7.6);
685 Eukaryota(12.7);Chordata(9.2);Mammalia(9.2);Rodentia(9.2);Muridae(9.2);Mus(9.2);

686 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);"Prevotellaceae"(100);Prevotella(99.8);
687 Eukaryota(6.3);Arthropoda(3.7);Insecta(3.7);Coleoptera(3.7);Tenebrionidae(3.7);Tribolium(3.7);
688 Archaea(19.4);"Euryarchaeota"(16.4);"Methanomicrobia"(9.6);Methanosarcinales(6.3);Methanosarcinaceae(6.2);Methanohalobium(3.3);
689 Eukaryota(13.9);Streptophyta(5.1);Liliopsida(5.1);Poales(5.1);Poaceae(5.1);Zea(5.1);
690 Bacteria(100);Firmicutes(90.2);Clostridia(90.2);Clostridiales(90.2);Peptostreptococcaceae(84.8);Sporacetigenium(66.1);
691 Archaea(23.5);"Euryarchaeota"(18.2);"Methanomicrobia"(14.1);Methanosarcinales(9.3);Methanosarcinaceae(8.7);Methanohalobium(6.5);
692 Bacteria(100);Firmicutes(94.1);Clostridia(94);Clostridiales(94);Lachnospiraceae(92.7);Marvinbryantia(35.1);
693 Bacteria(86.5);Cyanobacteria_Chloroplast(5.9);Chloroplast(5.9);Chloroplast_order_incertae_sedis(5.9);Chloroplast(5.9);Chlorarachniophyceae(5.9);
694 Eukaryota(13.8);Streptophyta(11.4);Liliopsida(11.4);Poales(11.4);Poaceae(11.4);Zea(11.4);
695 Eukaryota(9.2);Streptophyta(8.4);Liliopsida(8.4);Poales(8.4);Poaceae(8.4);Zea(8.4);
696 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Actinomycetales(100);Corynebacteriaceae(99.9);Corynebacterium(99.8);
697 Eukaryota(15);Streptophyta(11.8);Liliopsida(11.8);Poales(11.8);Poaceae(11.8);Zea(11.8);
698 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(100);Johnsonella(65.6);
699 Eukaryota(24);Streptophyta(16.3);Liliopsida(16.3);Poales(16.3);Poaceae(16.3);Zea(16.3);
700 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);"Porphyromonadaceae"(100);Parabacteroides(91.3);
701 Bacteria(100);Firmicutes(96.3);Clostridia(96.3);Clostridiales(96.3);Peptostreptococcaceae(87.9);Sporacetigenium(53.4);
702 Archaea(16.4);"Euryarchaeota"(15.9);Thermoplasmata(8.4);Thermoplasmatales(8.4);Thermoplasmatales_incertae_sedis(6);Thermogymnomonas(6);
703 Bacteria(100);Firmicutes(69.2);Clostridia(61.6);Clostridiales(61.6);Lachnospiraceae(55.6);Oribacterium(21.6);
704 Bacteria(100);Firmicutes(99.9);Clostridia(98.1);Clostridiales(98);Lachnospiraceae(78.8);Clostridium_XIVb(55.2);
705 Eukaryota(16.8);Chordata(13.2);Mammalia(13.2);Rodentia(13.2);Muridae(13.2);Mus(13.2);
706 Eukaryota(11.6);Streptophyta(11.1);Liliopsida(11.1);Poales(11.1);Poaceae(11.1);Zea(11.1);
707 Eukaryota(10.9);Streptophyta(9.3);Liliopsida(9.3);Poales(9.3);Poaceae(9.3);Zea(9.3);
708 Bacteria(80.3);Firmicutes(25.6);Clostridia(19.2);Clostridiales(14.3);Peptococcaceae_2(5.2);Cryptanaerobacter(5.1);
709 Eukaryota(18.8);Arthropoda(12.2);Insecta(12.2);Coleoptera(12.2);Tenebrionidae(12.2);Tribolium(12.2);
710 Bacteria(89);Firmicutes(26.1);Clostridia(18.7);Clostridiales(15.5);Incertae_Sedis_XI(3.6);Anaerospaera(3.6);
711 Bacteria(99.9);"Bacteroidetes"(98.9);"Bacteroidia"(64.8);"Bacteroidales"(64.8);"Porphyromonadaceae"(58.3);Barnesiella(33.2);
712 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.7);Coproccoccus(50.1);
713 Bacteria(90);Firmicutes(29.9);Clostridia(25);Clostridiales(22.5);Clostridiales_Incertae_Sedis_XI(5.9);Parvimonas(4.8);
714 Bacteria(86.8);"Bacteroidetes"(23.4);Flavobacteria(9.5);Flavobacteriales(9.5);Cryomorphaceae(4.6);Crocinitomix(2.2);
715 Eukaryota(16.7);Streptophyta(11.4);Liliopsida(11.4);Poales(11.4);Poaceae(11.4);Zea(11.4);
716 Eukaryota(13.2);Streptophyta(9.9);Liliopsida(9.9);Poales(9.9);Poaceae(9.9);Zea(9.9);
717 Bacteria(100);Firmicutes(95.4);Clostridia(95.3);Clostridiales(95.3);Eubacteriaceae(28.6);Eubacterium(28.6);
718 Bacteria(100);Firmicutes(96.8);Clostridia(96.3);Clostridiales(96.3);Ruminococcaceae(95.9);Anaerotruncus(66.6);
719 Bacteria(100);Firmicutes(99.9);Clostridia(99.2);Clostridiales(99.2);Clostridiales_Incertae_Sedis_XIII(79.5);Anaerovorax(78.9);
720 Bacteria(100);Firmicutes(95.7);Clostridia(94.7);Clostridiales(94.7);Ruminococcaceae(91.3);Flavonifractor(45.2);
721 Bacteria(88.2);"Deferribacteres"(6.2);Deferribacteres(6.2);Deferribacterales(6.2);Deferribacteraceae(6.2);Mucispirillum(5.9);
722 Bacteria(88.4);Firmicutes(35.5);Clostridia(27.7);Clostridiales(24.8);Clostridiales_Incertae_Sedis_XI(8.3);Parvimonas(6.3);
723 Bacteria(100);Firmicutes(58.2);Clostridia(58.2);Clostridiales(58.2);Ruminococcaceae(58);Oscillibacter(40.8);
724 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Peptostreptococcaceae(100);Clostridium_XI(100);
725 Archaea(18.1);"Euryarchaeota"(14.6);Halobacteria(8.6);Halobacteriales(8.6);Halobacteriaceae(8.6);Halonotius(2.5);
726 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(83.5);Pseudoflavonifractor(62.4);
727 Archaea(24.5);"Euryarchaeota"(20.9);"Methanomicrobia"(16);Methanosarcinales(9.9);Methanosarcinaceae(9.8);Methanohalobium(6.8);
728 Bacteria(100);"Actinobacteria"(95.9);Actinobacteria(95.9);Coriobacteriales(95.9);Coriobacteriaceae(95.9);Enterorhabdus(89.3);
729 Eukaryota(24.9);Chordata(21);Mammalia(21);Rodentia(15.5);Muridae(15.5);Mus(15.5);
730 Eukaryota(27.1);Chordata(26.4);Mammalia(26.4);Rodentia(20);Muridae(20);Mus(20);
731 Eukaryota(19.2);Chordata(16.9);Mammalia(16.9);Rodentia(13);Muridae(13);Mus(13);
732 Bacteria(100);"Bacteroidetes"(10.8);"Bacteroidia"(5.1);"Bacteroidales"(5.1);Bacteroidaceae(5.1);Acetomicrobium(5.1);
733 Archaea(13.7);"Euryarchaeota"(11.5);Halobacteria(10.5);Halobacteriales(10.5);Halobacteriaceae(10.5);Halobaculum(5.6);
734 Bacteria(80.3);Cyanobacteria_Chloroplast(10.8);Chloroplast(10.8);Chloroplast_order_incertae_sedis(10.8);Chloroplast(10.8);Bangiophyceae(10.1);
735 Eukaryota(16.2);Arthropoda(6.8);Insecta(6.8);Coleoptera(6.8);Tenebrionidae(6.8);Tribolium(6.8);
736 Bacteria(100);Firmicutes(99.4);Erysipelotrichia(61.1);Erysipelotrichales(61.1);Erysipelotrichaceae(61.1);Clostridium_XVIII(58.6);
737 Bacteria(100);Firmicutes(92.4);Erysipelotrichia(65.8);Erysipelotrichales(65.8);Erysipelotrichaceae(65.8);Turicibacter(65.7);
738 Bacteria(100);Firmicutes(100);Erysipelotrichia(100);Erysipelotrichales(100);Erysipelotrichaceae(100);Coprobaillus(85.6);
739 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(55);Flavonifractor(21.4);
740 Bacteria(81.9);"Chloroflexi"(7.5);Anaerolineales(5.9);Anaerolineaceae(5.9);Bellilinea(4.3);
741 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(100);"Bacteroidales"(100);"Rikenellaceae"(100);Alistipes(100);
742 Eukaryota(13.3);Chordata(10.6);Mammalia(10.6);Rodentia(9.1);Muridae(9.1);Mus(9.1);
743 Bacteria(100);Firmicutes(99.7);Clostridia(99.7);Clostridiales(99.7);Lachnospiraceae(97.7);Marvinbryantia(26.5);
744 Bacteria(75.1);"Proteobacteria"(37.8);Gammaproteobacteria(25.9);Chromatiales(10.8);Halothiobacillaceae(9.9);Thioalkalibacter(9.8);
745 Bacteria(100);Firmicutes(99.8);Clostridia(99.7);Clostridiales(99.7);Clostridiaceae_1(18.5);Anaerospobacter(18.4);
746 Archaea(19.2);"Euryarchaeota"(16.8);"Methanomicrobia"(11.6);Methanosarcinales(7.1);Methanosarcinaceae(6.6);Methanohalobium(4.5);
747 Bacteria(83.4);"Bacteroidetes"(24.3);"Sphingobacteria"(14.1);"Sphingobacteriales"(14.1);Cytophagaceae(11.4);Litoribacter(4.5);
748 Bacteria(100);Firmicutes(85.1);Clostridia(85);Clostridiales(85);Lachnospiraceae(79.4);Johnsonella(38.5);
749 Eukaryota(8.1);Streptophyta(7.2);Liliopsida(7.2);Poales(7.2);Poaceae(7.2);Zea(7.2);
750 Eukaryota(14.8);Chordata(11.1);Mammalia(11.1);Rodentia(11.1);Muridae(11.1);Mus(11.1);
751 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Clostridium_XIVb(49.6);
752 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(98.4);"Bacteroidales"(98.4);"Porphyromonadaceae"(81.7);Barnesiella(71);
753 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.3);Lachnospiraceae_incertae_sedis(50);
754 Bacteria(100);Firmicutes(100);Clostridia(69);Clostridiales(69);Lachnospiraceae(67.6);Pseudobutyrvibrio(53.9);
755 Archaea(16.2);"Euryarchaeota"(13.8);"Methanomicrobia"(8.7);Methanosarcinales(7.9);Methanosarcinaceae(7.9);Methanimicrococcus(7.7);
756 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(93.7);Lachnospiraceae_incertae_sedis(36.4);
757 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(80.3);"Bacteroidales"(80.3);"Porphyromonadaceae"(64.1);Tannerella(14.8);
758 Eukaryota(10.7);Arthropoda(5.7);Insecta(5.7);Coleoptera(5.7);Tenebrionidae(5.7);Tribolium(5.7);
759 Bacteria(100);Firmicutes(95);Clostridia(95);Clostridiales(95);Ruminococcaceae(94.8);Clostridium_IV(40.8);
760 Bacteria(86.2);"Bacteroidetes"(28.4);"Bacteroidia"(10.1);"Bacteroidales"(10.1);"Porphyromonadaceae"(5.5);Petrimonas(3.7);
761 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Flavonifractor(70.6);
762 Bacteria(100);Firmicutes(99.3);Clostridia(99.3);Clostridiales(99.3);Ruminococcaceae(96.4);Clostridium_IV(38);
763 Bacteria(100);Firmicutes(99.5);Clostridia(99.5);Clostridiales(99.5);Lachnospiraceae(99.3);Shuttleworthia(27.4);
764 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.6);Marvinbryantia(36);
765 Bacteria(100);"Bacteroidetes"(97.2);"Bacteroidia"(87);"Bacteroidales"(87);"Porphyromonadaceae"(73.3);Tannerella(35.8);
766 Bacteria(87.8);"Bacteroidetes"(30.2);"Bacteroidia"(9.6);"Bacteroidales"(9.6);"Porphyromonadaceae"(5.9);Petrimonas(4.4);
767 Bacteria(100);"Bacteroidetes"(51.5);"Bacteroidia"(40.5);"Bacteroidales"(40.5);"Porphyromonadaceae"(37.7);Tannerella(27.9);
768 Bacteria(100);"Bacteroidetes"(92);"Bacteroidia"(75.5);"Bacteroidales"(75.5);"Porphyromonadaceae"(69.9);Petrimonas(25.4);
769 Bacteria(100);Firmicutes(99.8);Clostridia(99.8);Clostridiales(99.8);Lachnospiraceae(97.2);Coproccoccus(41.5);
770 Eukaryota(14.7);Arthropoda(8.6);Insecta(8.6);Coleoptera(8.6);Tenebrionidae(8.6);Tribolium(8.6);
771 Eukaryota(9.4);Arthropoda(6.9);Insecta(6.9);Coleoptera(6.9);Tenebrionidae(6.9);Tribolium(6.9);

772 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(86.6);Oscillibacter(77.3);
773 Bacteria(100);Firmicutes(99.8);Clostridia(99.8);Clostridiales(99.8);Lachnospiraceae(96.3);Oribacterium(15.9);
774 Bacteria(88.4);"Bacteroidetes"(32.5);Flavobacteria(12.8);"Flavobacteriales"(12.8);Flavobacteriaceae(11.3);Soonwooa(6.7);
775 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(91.4);Oscillibacter(64.6);
776 Bacteria(91.2);"Proteobacteria"(34.6);Alphaproteobacteria(5.4);Rickettsiales(4.1);Anaplasmataceae(4);Wolbachia(4);
777 Eukaryota(12.5);Arthropoda(8.7);Insecta(8.7);Coleoptera(8.7);Tenebrionidae(8.7);Tribolium(8.7);
778 Eukaryota(12.2);Chordata(8.5);Mammalia(8.5);Rodentia(8.5);Muridae(8.5);Mus(8.5);
779 Bacteria(100);"Proteobacteria"(34.1);Alphaproteobacteria(6.3);Rhodospirillales(2.8);Rhodospirillaceae(2.8);Tistrella(2.6);
780 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.8);Coproccoccus(20.2);
781 Bacteria(100);Firmicutes(84.5);Clostridia(84.2);Clostridiales(84.2);Lachnospiraceae(76.6);Johnsonella(24.4);
782 Bacteria(100);Firmicutes(95.5);Clostridia(95.4);Clostridiales(95.4);Ruminococcaceae(92.2);Anaerotruncus(67.3);
783 Bacteria(100);"Bacteroidetes"(51.1);"Bacteroidia"(46.2);"Bacteroidales"(46.2);"Porphyromonadaceae"(44.8);Barnesiella(40.5);
784 Bacteria(90.2);Firmicutes(30.8);Clostridia(25.6);Clostridiales(23.4);Clostridiales_Incertae_Sedis_XI(6.1);Parvimonas(4.9);
785 Bacteria(100);Firmicutes(97.2);Erysipelotrichia(97.2);Erysipelotrichales(97.2);Erysipelotrichaceae(97.2);Coproccillus(79.1);
786 Bacteria(100);Firmicutes(85.9);Clostridia(85.9);Clostridiales(85.9);Lachnospiraceae(80.9);Oribacterium(11.3);
787 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(59.8);Flavonifractor(30.1);
788 Bacteria(100);Firmicutes(76.1);Negativicutes(13.1);Selenomonadales(13.1);Veillonellaceae(13.1);Propionispira(2.8);
789 Bacteria(89.8);"Bacteroidetes"(33.8);"Bacteroidia"(11.1);"Bacteroidales"(11.1);"Porphyromonadaceae"(6.2);Petrimonas(3.1);
790 Archaea(26.1);"Euryarchaeota"(23.1);"Methanomicrobia"(7.7);Methanosarcinales(6.6);Methanosarcinaceae(6.6);Methanomicrococcus(6.5);
791 Bacteria(100);"Bacteroidetes"(99.9);"Bacteroidia"(98.6);"Bacteroidales"(98.6);"Porphyromonadaceae"(95.2);Tannerella(38.1);
792 Bacteria(100);Firmicutes(94.5);Clostridia(94.4);Clostridiales(94.4);Ruminococcaceae(92);Acetanaerobacterium(86.9);
793 Bacteria(100);Firmicutes(98);Clostridia(96.5);Clostridiales(96.5);Ruminococcaceae(56.1);Acetanaerobacterium(20.2);
794 Bacteria(100);Firmicutes(99.6);Clostridia(95.5);Clostridiales(95);Ruminococcaceae(43.8);Saccharofermentans(42.8);
795 Bacteria(100);"Bacteroidetes"(75.8);"Bacteroidia"(66.9);"Bacteroidales"(66.9);"Porphyromonadaceae"(43);Barnesiella(30.5);
796 Bacteria(100);"Bacteroidetes"(91);"Bacteroidia"(80.6);"Bacteroidales"(80.6);"Porphyromonadaceae"(66.9);Tannerella(43);
797 Bacteria(89.5);"Proteobacteria"(34.2);Alphaproteobacteria(14.2);Rhizobiales(5.9);Bradyrhizobiaceae(4.1);Agromonas(4.1);
798 Bacteria(100);Firmicutes(67.1);Clostridia(67.1);Clostridiales(67.1);Lachnospiraceae(62.3);Johnsonella(44.8);
799 Bacteria(85.7);"Bacteroidetes"(25.2);"Bacteroidia"(6.2);"Bacteroidales"(6.2);"Porphyromonadaceae"(3.4);Petrimonas(1.4);
800 Bacteria(100);Firmicutes(56.9);Clostridia(32.2);Clostridiales(30);Ruminococcaceae(19);Saccharofermentans(16.8);
801 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.9);Lactonifactor(14.6);
802 Bacteria(100);Firmicutes(98.9);Clostridia(98.9);Clostridiales(98.8);Lachnospiraceae(97.6);Marvinbryantia(32.7);
803 Bacteria(100);Firmicutes(99.7);Clostridia(99.4);Clostridiales(99.4);Clostridiaceae_1(76);Sarcina(55.9);
804 Bacteria(82.2);"Proteobacteria"(34.4);Gammaproteobacteria(19.8);Chromatiales(10.6);Halothiobacillaceae(9.9);Thioalkalibacter(9.9);
805 Bacteria(87.2);"Proteobacteria"(29.8);Deltaproteobacteria(10.1);Syntrophobacteriales(4.9);Syntrophaceae(3.3);Smithella(3.3);
806 Bacteria(100);Firmicutes(64.6);Clostridia(64.6);Clostridiales(64.6);Lachnospiraceae(62.5);Marvinbryantia(16.8);
807 Bacteria(100);Firmicutes(99.9);Clostridia(99.9);Clostridiales(99.9);Ruminococcaceae(99.9);Oscillibacter(99.3);
808 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(99.6);"Bacteroidales"(99.6);"Porphyromonadaceae"(93.2);Tannerella(71);
809 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.6);Moryella(38.5);
810 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Clostridium_IV(42.9);
811 Eukaryota(12.9);Arthropoda(8.5);Insecta(8.5);Coleoptera(8.5);Tenebrionidae(8.5);Tribolium(8.5);
812 Bacteria(100);"Bacteroidetes"(77.3);"Bacteroidia"(41.9);"Bacteroidales"(41.9);"Porphyromonadaceae"(39.7);Tannerella(22.8);
813 Bacteria(99.9);Firmicutes(88.9);Clostridia(88.9);Clostridiales(88.9);Ruminococcaceae(78);Anaerotruncus(45.2);
814 Eukaryota(10.6);Arthropoda(6);Insecta(6);Coleoptera(6);Tenebrionidae(6);Tribolium(6);
815 Eukaryota(15.5);Arthropoda(9.6);Insecta(9.6);Coleoptera(9.6);Tenebrionidae(9.6);Tribolium(9.6);
816 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(59);Oscillibacter(51.5);
817 Bacteria(76.6);"Proteobacteria"(21.6);Deltaproteobacteria(10.7);Myxococcales(4.8);Nannocystaceae(4.7);Enhygromyxa(4.7);
818 Bacteria(100);Firmicutes(97.8);Clostridia(80.7);Clostridiales(80.7);Lachnospiraceae(70.7);Catonella(12);
819 Bacteria(99.6);"Bacteroidetes"(81.5);Flavobacteria(22.8);"Flavobacteriales"(22.8);Cryomorphaceae(21.6);Fluviicola(20.9);
820 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(98.9);Lachnospiraceae_incertae_sedis(49.1);
821 Bacteria(100);Firmicutes(98.8);Clostridia(98.8);Clostridiales(98.8);Lachnospiraceae(98.1);Lactonifactor(39.7);
822 Bacteria(100);Firmicutes(82.8);Clostridia(80.6);Clostridiales(80.4);Ruminococcaceae(64);Flavonifractor(9.9);
823 Bacteria(86.1);"Thermodesulfobacteria"(5);Thermodesulfobacteriales(5);Thermodesulfobacteriaceae(5);Caldimicrobium(5);
824 Eukaryota(9.5);Arthropoda(6);Insecta(6);Coleoptera(6);Tenebrionidae(6);Tribolium(6);
825 Archaea(21.1);"Euryarchaeota"(18.7);Halobacteria(11.2);Halobacteriales(11.2);Halobacteriaceae(11.2);Halonotius(5.2);
826 Bacteria(100);"Bacteroidetes"(99.8);"Bacteroidia"(93.8);"Bacteroidales"(93.8);"Porphyromonadaceae"(65.2);Barnesiella(50.6);
827 Bacteria(99.9);Firmicutes(88.2);Clostridia(88);Clostridiales(88);Lachnospiraceae(83.4);Marvinbryantia(21.3);
828 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(94.2);
829 Eukaryota(9.7);Streptophyta(8.6);Liliopsida(8.6);Poales(8.6);Poaceae(8.6);Zea(8.6);
830 Bacteria(90);"Bacteroidetes"(24.3);"Bacteroidia"(8);"Bacteroidales"(8);"Porphyromonadaceae"(3.3);Petrimonas(1.7);
831 Bacteria(80.3);"Proteobacteria"(38.3);Gammaproteobacteria(24.7);Chromatiales(10.2);Halothiobacillaceae(9.4);Thioalkalibacter(9.3);
832 Bacteria(100);Firmicutes(96.8);Clostridia(87.3);Clostridiales(79.5);Ruminococcaceae(71.4);Papillibacter(22.1);
833 Bacteria(100);"Bacteroidetes"(99.9);"Bacteroidia"(55.1);"Bacteroidales"(55.1);"Rikenellaceae"(24.5);Rikenella(24.5);
834 Bacteria(100);Firmicutes(96.6);Clostridia(93.7);Clostridiales(93.3);Lachnospiraceae(71.6);Robinsoniella(21.4);
835 Bacteria(89.5);"Elusimicrobia"(2.9);Elusimicrobia(2.9);Elusimicrobiales(2.9);Elusimicrobiaceae(2.9);Elusimicrobium(2.9);
836 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(90.8);
837 Bacteria(89.7);"Proteobacteria"(40.1);Alphaproteobacteria(18.4);Rhizobiales(5.9);Bradyrhizobiaceae(3.8);Agromonas(3.7);
838 Eukaryota(12.8);Chordata(7.1);Mammalia(7.1);Primates(1.8);Hominidae(1.8);Homo(1.8);
839 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(99.4);Syntrophococcus(30.2);
840 Bacteria(100);"Bacteroidetes"(100);"Bacteroidia"(99.7);"Bacteroidales"(99.7);"Porphyromonadaceae"(98.2);Porphyromonas(30.4);
841 Bacteria(79.3);Firmicutes(28.6);Clostridia(21.7);Clostridiales(17.3);Incertae_Sedis_XI(5.9);Murdochiella(5.9);
842 Bacteria(100);Firmicutes(98.3);Clostridia(98);Clostridiales(98);Ruminococcaceae(91.1);Pseudoflavonifractor(53.9);
843 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(86.2);Clostridium_IV(68.5);
844 Eukaryota(13.2);Chordata(11.4);Mammalia(11.4);Rodentia(11.4);Muridae(11.4);Mus(11.4);
845 Bacteria(100);Firmicutes(93.2);Clostridia(89.2);Clostridiales(86.9);Ruminococcaceae(52.1);Saccharofermentans(31.7);
846 Bacteria(100);Firmicutes(56.9);Clostridia(56.1);Clostridiales(56.1);Ruminococcaceae(48.7);Flavonifractor(34.8);
847 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(98.3);Lachnospiraceae_incertae_sedis(42.9);
848 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Ruminococcaceae(100);Oscillibacter(85.1);
849 Eukaryota(14.1);Arthropoda(7);Insecta(7);Coleoptera(7);Tenebrionidae(7);Tribolium(7);
850 Eukaryota(20);Streptophyta(8);Liliopsida(8);Poales(8);Poaceae(8);Zea(8);
851 Bacteria(100);"Bacteroidetes"(92.8);"Bacteroidia"(91.5);"Bacteroidales"(91.5);"Porphyromonadaceae"(70.1);Barnesiella(39.2);
852 Bacteria(99.3);"Bacteroidetes"(93);"Bacteroidia"(69);"Bacteroidales"(69);"Porphyromonadaceae"(59.1);Tannerella(30.2);
853 Archaea(20.3);"Euryarchaeota"(17.2);Halobacteria(9.2);Halobacteriales(9.2);Halobacteriaceae(9.2);Halonotius(3.9);
854 Bacteria(100);Firmicutes(96);Clostridia(99.6);Clostridiales(99.6);Lachnospiraceae(93.7);Lactonifactor(27.4);
855 Bacteria(100);Firmicutes(55.6);Clostridia(53.7);Clostridiales(53.7);Clostridiaceae_1(46.2);Anaerobacter(44.6);
856 Bacteria(85.2);"Actinobacteria"(9.6);Actinobacteria(9.6);Actinomycetales(9);Micromonosporaceae(6.6);Catelliglobospora(6.3);
857 Bacteria(86.9);"Bacteroidetes"(28.5);"Bacteroidia"(10.1);"Bacteroidales"(10.1);"Porphyromonadaceae"(5.3);Petrimonas(2.8);

858 Bacteria(100);"Actinobacteria"(100);Actinobacteria(100);Coriobacteriales(100);Coriobacteriaceae(100);Enterorhabdus(100);
859 Eukaryota(5);Chordata(4.1);Mammalia(4.1);Rodentia(4.1);Muridae(4.1);Mus(4.1);
860 Bacteria(87.6);"Proteobacteria"(28.9);Deltaproteobacteria(9.2);Desulfobacterales(5.3);Desulfobacteraceae(2.6);Desulfatiferula(2.2);
861 Bacteria(100);Firmicutes(100);Bacilli(100);Lactobacillales(100);Lactobacillaceae(100);Lactobacillus(100);
862 Eukaryota(12.8);Chordata(9.7);Mammalia(9.7);Rodentia(9.7);Muridae(9.7);Mus(9.7);
863 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(99.8);Ruminococcaceae(97.1);Oscillibacter(82.3);
864 Eukaryota(25.9);Chordata(24.4);Mammalia(24.4);Rodentia(18.8);Muridae(18.8);Mus(18.8);
865 Bacteria(100);Firmicutes(100);Clostridia(86.3);Clostridiales(86.3);Ruminococcaceae(86);Flavonifractor(57.2);
866 Archaea(18.7);"Euryarchaeota"(15.7);"Methanomicrobia"(8.1);Methanosarcinales(6.5);Methanosarcinaceae(6.5);Methanimicrococcus(6.5);
867 Bacteria(100);Firmicutes(99.3);Clostridia(97.1);Clostridiales(97.1);Ruminococcaceae(66.7);Oscillibacter(23.4);
868 Archaea(18.5);"Euryarchaeota"(15.1);"Methanomicrobia"(7.2);Methanosarcinales(4.3);Methanosarcinaceae(4.2);Methanohalobium(2.2);
869 Eukaryota(18.8);Chordata(14);Mammalia(14);Rodentia(14);Muridae(14);Mus(14);
870 Archaea(10.5);"Euryarchaeota"(9.7);Thermoplasmata(6.7);Thermoplasmatales(6.7);Ferroplasmaceae(6.5);Ferroplasma(6.5);
871 Eukaryota(16.9);Arthropoda(10.8);Insecta(10.8);Coleoptera(10.8);Tenebrionidae(10.8);Tribolium(10.8);
872 Eukaryota(12.5);Arthropoda(8);Insecta(8);Coleoptera(8);Tenebrionidae(8);Tribolium(8);
873 Archaea(18.3);"Euryarchaeota"(14.8);Halobacteria(10.1);Halobacteriales(10.1);Halobacteriaceae(10.1);Halonotius(3.8);
874 Bacteria(100);"Bacteroidetes"(99.9);"Bacteroidia"(94.2);"Bacteroidales"(94.2);"Porphyromonadaceae"(93.6);Barnesiella(83.2);
875 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Lachnospiraceae(96.9);Clostridium_XIVa(82.9);
876 Eukaryota(11.9);Streptophyta(10.8);Liliopsida(10.8);Poales(10.8);Poaceae(10.8);Zea(10.8);
877 Eukaryota(24.4);Chordata(23.6);Mammalia(23.6);Rodentia(18.2);Muridae(18.2);Mus(18.2);
878 Bacteria(83.2);"Thermodesulfobacteria"(4.4);Thermodesulfobacteria(4.4);Thermodesulfobacteriales(4.4);Thermodesulfobacteriaceae(4.4);Caldimicrobium(4.4);
879 Bacteria(89);"Proteobacteria"(36.7);Alphaproteobacteria(5.2);Rickettsiales(3.3);Anaplasmataceae(3.3);Wolbachia(3.3);
880 Eukaryota(17.1);Chordata(10.9);Mammalia(10.9);Rodentia(10.9);Muridae(10.9);Mus(10.9);
881 Bacteria(100);Firmicutes(100);Negativicutes(100);Selenomonadales(100);Acidaminococcaceae(96.8);Phascolarctobacterium(84.2);
882 Eukaryota(10.8);Arthropoda(6.1);Insecta(6.1);Coleoptera(6.1);Tenebrionidae(6.1);Tribolium(6.1);
883 Bacteria(86.4);Firmicutes(32.4);Clostridia(27.7);Clostridiales(24.9);Clostridiales_Incertae_Sedis_XI(10.3);Parvimonas(8.3);
884 Bacteria(100);Firmicutes(99.4);Clostridia(99.4);Clostridiales(99.4);Lachnospiraceae(89.7);Marvinbryantia(28.6);
885 Bacteria(89);"Proteobacteria"(34.8);Alphaproteobacteria(15);Rhizobiales(6.2);Bradyrhizobiaceae(4.2);Agromonas(4.1);
886 Bacteria(90);"Proteobacteria"(33.6);Epsilonproteobacteria(6.4);Campylobacteriales(6.2);"Hydrogenimonaceae"(5.6);Hydrogenimonas(5.6);
887 Bacteria(100);Firmicutes(100);Clostridia(100);Clostridiales(100);Peptostreptococcaceae(100);Clostridium_XI(92);

Supplemental Table S1. Significant effects of diet and exercise at different levels of taxonomy.

| | Main effects 2-way ANOVA | | | post-hoc testing | | | | |
|---------------------|--------------------------|------|----------|------------------|-----------|-----------------|-------------|---------------|
| | High fat diet | | exercise | interaction | ND vs HFD | ND+ex vs HFD+ex | ND vs ND+ex | HFD vs HFD+ex |
| Phylum | | | | | | | | |
| Bacteroidetes | ↓ | **** | ↓ | ** | **** | **** | * | * |
| Firmicutes | ↑ | **** | ↑ | ** | **** | **** | * | * |
| Tenericutes | ↓ | ** | | | | | * | |
| Proteobacteria | ↑ | * | ↑ | * | | | | |
| Actinobacteria | | | ↑ | * | | | | |
| Class | | | | | | | | |
| Bacteroidia | ↓ | **** | ↓ | *** | **** | **** | * | * |
| Clostridia | ↑ | **** | ↑ | ** | **** | **** | * | * |
| Erysipelotrichia | ↓ | **** | | | ** | *** | | |
| Mollicutes | ↓ | ** | | * | ** | | * | |
| Bacilli | ↑ | ** | | | | | | |
| Gammaproteobacteria | ↑ | * | | | | | | |
| Negativicutes | ↑ | * | | | | | | |
| Betaproteobacteria | | | ↑ | ** | | | | * |
| Actinobacteria | | | ↑ | * | | | | |
| Sphingobacteria | | | ↑ | * | | * | | * |
| Flavobacteria | | | ↑ | * | | | | * |
| Alphaproteobacteria | | | ↑ | * | | | | * |
| Order | | | | | | | | |
| Bacteroidales | ↓ | **** | ↓ | *** | **** | **** | * | * |
| Clostridiales | ↑ | **** | ↑ | ** | **** | **** | * | * |
| Erysipelotrichales | ↓ | **** | | | ** | *** | | |
| Lactobacillales | ↑ | *** | | | | * | | |
| Anaeroplasmatales | ↓ | ** | | * | ** | | * | |
| Bacillales | ↓ | ** | | | | | | |
| Pseudomonadales | ↑ | * | ↑ | * | | * | | * |
| Enterobacteriales | ↑ | * | | | | | | |
| Selenomonadales | ↑ | * | | | | | | |
| Burkholderiales | | | ↑ | ** | | | | * |
| Actinomycetales | | | ↑ | * | | | | |
| Coriobacteriales | | | ↑ | * | | | * | |
| Sphingobacteriales | | | ↑ | * | | * | | * |
| Flavobacteriales | | | ↑ | * | | | | * |
| Caulobacterales | | | ↑ | * | | | | * |
| Rhizobiales | | | ↑ | * | | | | * |

Supplemental Figure S2

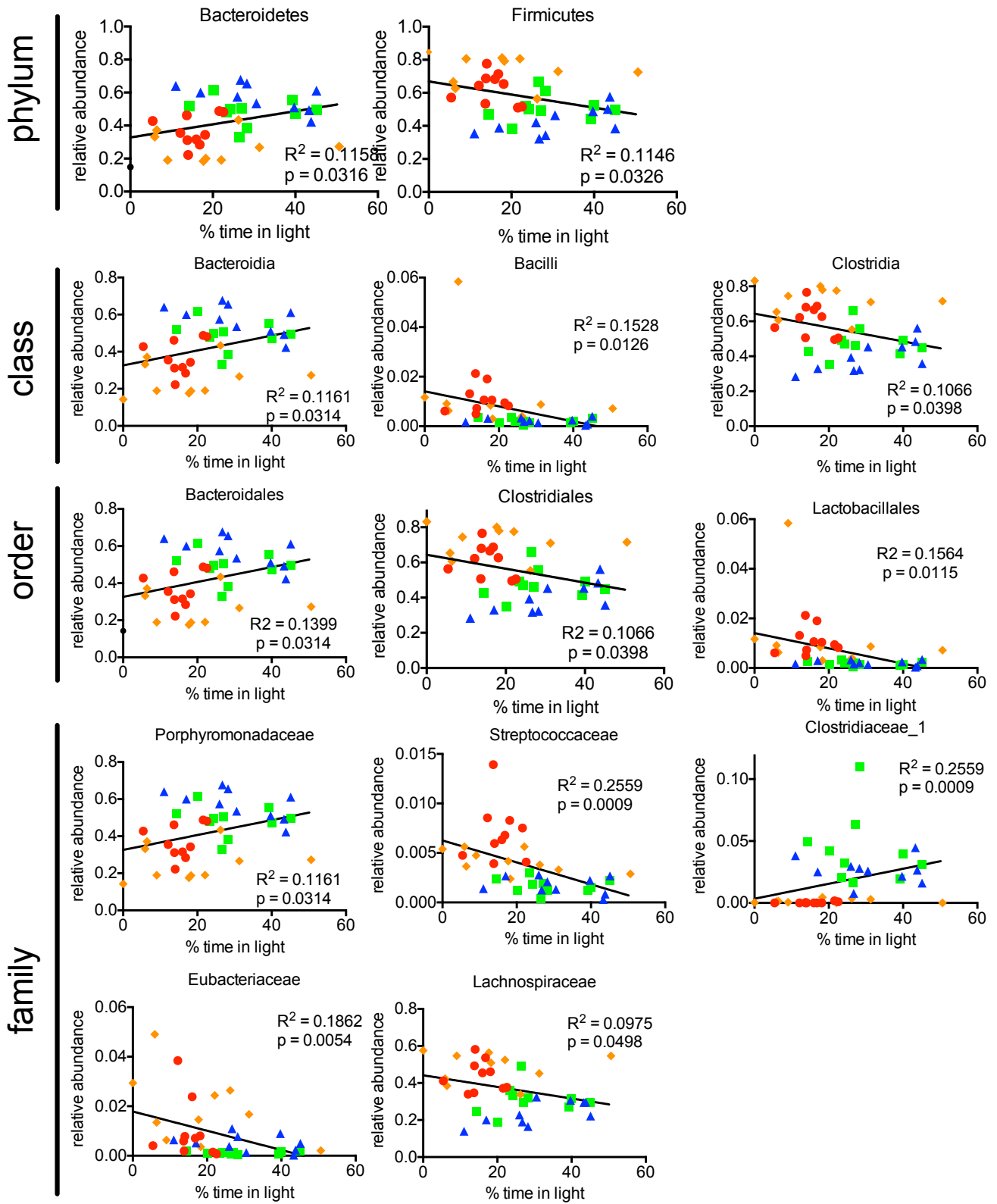


Figure S2. Anxiety levels were associated with several bacteria at the level of phyla, class, order, and family. Color scheme of data points follows the format as in Figure 1 (ND = blue, ND + exercise = green, HFD = red, and HFD + exercise in orange). Linear regression analysis for individual mice was performed with R^2 values indicating goodness of fit and uncorrected p values for slope calculated by F test.

Supplemental Table S3. Associations of bacterial abundances with body weight.

| taxa | Slope | R square | p value |
|-----------------------------------|--------------------------|-----------------|----------------|
| Bacteroidetes | -0.002756 ± 0.0006099 | 0.3495 | 0.0001 |
| Firmicutes | 0.002759 ± 0.0006103 | 0.3498 | 0.0001 |
| Bacteroidia | -0.002776 ± 0.0006155 | 0.3486 | 0.0001 |
| Bacilli | 0.0001799 ± 4.064e-005 | 0.3403 | 0.0001 |
| Clostridia | 0.002896 ± 0.0006277 | 0.3591 | 0.0001 |
| Erysipelotrichia | -0.0003175 ± 7.042e-005 | 0.3485 | 0.0001 |
| Bacteroidales | -0.002776 ± 0.0006155 | 0.3486 | 0.0001 |
| Lactobacillales | 0.0001825 ± 4.060e-005 | 0.3471 | 0.0001 |
| Clostridiales | 0.002896 ± 0.0006277 | 0.3591 | 0.0001 |
| Erysipelotrichales | -0.0003175 ± 7.042e-005 | 0.3485 | 0.0001 |
| Streptococcaceae | 6.102e-005 ± 1.012e-005 | 0.489 | 0.0001 |
| Clostridiaceae_1 | -0.0004780 ± 9.020e-005 | 0.4249 | 0.0001 |
| Lachnospiraceae | 0.002492 ± 0.0005078 | 0.388 | 0.0001 |
| Porphyromonadaceae | -0.002776 ± 0.0006156 | 0.3486 | 0.0001 |
| Erysipelotrichaceae | -0.0003175 ± 7.042e-005 | 0.3485 | 0.0001 |
| Ruminococcaceae | 0.0005949 ± 0.0001923 | 0.2012 | 0.0037 |
| Lactobacillaceae | 0.0001214 ± 3.995e-005 | 0.1955 | 0.0043 |
| Clostridiales_Incertae_Sedis_XIII | 8.926e-007 ± 3.416e-007 | 0.1558 | 0.0129 |
| Microbacteriaceae | 7.868e-008 ± 3.026e-008 | 0.1544 | 0.0133 |
| Bacillales | -2.547e-006 ± 1.005e-006 | 0.1446 | 0.0155 |
| Gammaproteobacteria | 1.005e-005 ± 3.993e-006 | 0.143 | 0.0161 |
| Proteobacteria | 1.269e-005 ± 5.423e-006 | 0.1259 | 0.0246 |
| Staphylococcaceae | -2.073e-006 ± 8.873e-007 | 0.1256 | 0.0248 |
| Bacteroidaceae | 2.696e-007 ± 1.184e-007 | 0.123 | 0.0286 |
| Xanthomonadaceae | 1.076e-006 ± 4.842e-007 | 0.1178 | 0.0324 |
| Enterobacteriales | 2.106e-006 ± 9.845e-007 | 0.1075 | 0.0389 |
| Enterobacteriaceae | 2.106e-006 ± 9.845e-007 | 0.1075 | 0.0389 |
| Rhizobiales | 5.527e-007 ± 2.618e-007 | 0.105 | 0.0414 |

Benjamini-Hochberg**p value**

0.00280

0.00140

0.00093

0.00070

0.00056

0.00047

0.00040

0.00035

0.00031

0.00028

0.00025

0.00023

0.00022

0.00020

0.00019

0.00648

0.00708

0.02007

0.01960

0.02170

0.02147

0.03131

0.03019

0.03337

0.03629

0.04189

0.04034

0.04140