Looking for a Signal in the Noise: Revisiting Obesity and the Microbiome

Running Title: The Human Microbiome and Obesity

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Abstract

Two recent studies have re-analyzed published data and found that when datasets are analyzed independently there was limited support for the widely accepted hypothesis that changes in the microbiome are associated with obesity. This hypothesis was reconsidered by increasing the number of datasets and pooling the results across the individual datasets. The Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines were applied to identify 10 studies for an updated and more synthetic analysis. Alpha diversity metrics and the relative risk of obesity based on those metrics were used to identify a limited number of significant associations with obesity; however, when the results of the studies were pooled using a random effects model significant associations were 10 observed between Shannon diversity, number of observed OTUs, and Shannon evenness 11 and obesity status. They were not observed for the ratio of Bacteroidetes and Firmicutes 12 or their individual relative abundances. Although these tests yielded small P-values, the difference between the Shannon diversity index of non-obese and obese individuals was 2.07%. A power analysis demonstrated that only one of the studies had sufficient power to detect a 5% difference in diversity. When Random Forest machine learning models were trained on one dataset and then tested using the other 9 datasets, the median accuracy varied between 33.01 and 64.77% (median=56.68%). Although there was support for a relationship between the microbial communities found in human feces and obesity status, this association is relatively weak and its detection is confounded by large interpersonal variation and insufficient sample sizes.

22 Importance

As interest in the human microbiome grows there is an increasing number of studies that can be used to test numerous hypotheses across human populations. The hypothesis

that variation in the gut microbiota can explain or be used to predict obesity status has received considerable attention and is frequently mentioned as an example for the role of the microbiome in human health. Here we assess this hypothesis using ten independent studies and find that although there is an association, it is smaller than can be detected by most microbiome studies. Furthermore, we directly tested the ability to predict obesity status based on the composition of an individual's microbiome and find that the median classification accuracy is between 33.01 and 64.77%. This type of analysis can be used to design future studies and expanded to explore other hypotheses.

3 Introduction

Obesity is a growing health concern with approximately 20% of the youth (aged 2-19) in
the United States classified as either overweight or obese (1). This number increases
to approximately 35% in adults (aged 20 or older) and these statistics have seen little
change since 2003 (1). Traditionally, the body mass index (BMI) has been used to classify
individuals as non-obese or obese (2). Recently, there has been increased interest in
the role of the microbiome in modulating obesity (3, 4). If the microbiome does affect
obesity status, then manipulating the microbiome could have a significant role in the future
treatment of obesity and in helping to stem the current epidemic.

There have been several studies that report observing a link between the composition of microbiome and obesity in animal models and in humans. The first such study used genetically obese mice and observed the ratio of the relative abundances of Bacteroidetes to Firmicutes (B:F) was lower in obese mice than lean mice (5). Translation of this result to humans by the same researchers did not observe this effect, but did find that obese individuals had a lower alpha-diversity than lean individuals (6). They also showed that the relative abundance of *Bacteroidetes* and *Firmicutes* increased and decreased, respectively, as obese individuals lost weight while on a fat or carbohydrate restricted diet (7). Two re-analysis studies by Walters et al. (8) and Finucane et al. (9) interrogated previously 50 published microbiome and obesity data and concluded that the previously reported 51 differences in community diversity and B:F among non-obese and obese individuals could not be generalized. Regardless of the results using human populations, studies using animal models where the community was manipulated with antibiotics or established 54 by colonizing germ-free animals with varied communities appear to support the association 55 since these manipulations yielded differences in animal weight (10-13). The purported association between the differences in the microbiome and obesity have been widely repeated with little attention given to the lack of a clear signal in human cohort studies.

The recent publication of additional studies that collected BMI data for each subject as well as other studies that were not included in the earlier re-analysis studies offered the opportunity to revisit the question relating the structure of the human microbiome to obesity. 61 One critique of the prior re-analysis studies is that the authors did not aggregate the results 62 across studies to increase the effective sample size. It is possible that there were small 63 associations within each study that were not statistically significant because the individual studies lacked sufficient power. Alternatively, diversity metrics may mask the appropriate 65 signal and it is necessary to measure the association at the level of microbial populations. The Walters re-analysis study demonstrated that Random Forest machine learning models 67 were capable of predicting obesity status within a single cohort, but did not attempt to test the models on other cohorts. The purpose of this study was to perform a meta-analysis of 69 the association between differences in the microbiome and obesity status by analyzing and applying a more systematic and synthetic approach than was used previously.

72 Results

Literature Review and Study Inclusion. To perform a robust meta-analysis and limit inclusion bias, we followed the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines to identify the studies that we analyzed (14). A detailed description of our selection process and the exact search terms are provided in the Supplemental Text and in Figure 1. Briefly, we searched PubMed for original research studies that involved studying obesity and the human microbiome. The initial search yielded 187 studies. We identified ten additional studies that were not designed to explicitly test for an association between the microbiome and obesity. We then manually curated the 196 studies to select those studies that included BMI and 16S rRNA gene sequence data. This yielded 10 eligible studies. An additional study was removed from our analysis because no individuals in the study had a BMI over 30. Among the final 10 studies, 3 were

from identified from our PubMed search (10, 15, 16), 5 were originally identified from the 10 studies that did not explicitly investigate obesity but included BMI data (17-21), and two datasets were used (22, 23) because these publications did not specifically look for any metabolic or obesity conditions but had control populations and enabled us to help mitigate 87 against publication biases associated with the bacterial microbiome and obesity. The ten 88 studies are summarized in Table 1. For comparison, two of these studies were included in the Finucane re-analysis study (10, 21) and four of these studies were included in the Walters re-analysis study (10, 15, 20, 21). The 16S rRNA gene sequence data from each 91 study was re-analyzed using a similar approach based on previously described methods for reducing the number of chimeric sequences and sequencing errors for 454 and Illumina MiSeq data (24, 25). The sequences were either clustered into operational taxonomic units (OTUs) using the average neighbor approach (26) or taxonomic groupings based on their 95 classification using a naive Bayesian classifier (27).

Alpha diversity analysis. We calculated the Shannon diversity index, observed richness, and Shannon evenness, the relative abundance of Bacteroidetes and Firmicutes, and the ratio of their relative abundance (B:F) for each sample. Once we transformed each of the six alpha diversity metrics to make them normally distributed, we used a t-test 100 to identify significant associations between the alpha diversity metric and whether an 101 individual was obese for each of the ten studies. The B:F and the relative abundance 102 of Firmicutes were not significantly associated with obesity in any study. We identified 103 seven P-values that were less than 0.05: three studies indicated obese individuals had 104 a lower richness, two studies indicated a significantly lower diversity, one study indicated 105 a significantly lower evenness, and one study indicated a significantly higher relative 106 abundance of Bacteroidetes (Figures 2 and S1). These results largely match those of the 107 Walters and Finucane re-analysis studies. Interestingly, although only two of the ten studies observed the previously reported association between lower diversity and obesity, the other studies appeared to have the same trend, albeit the differences were not statistically

significant. We used a random effects linear model to combine the studies using the study as the random effect and found statistical support for decreased richness, evenness, and diversity among obese individuals (all P<0.011). Although there was a significant 113 relationship between these metrics and obesity status, the effect size was quite small. 114 The obese individuals averaged 7.47% lower richness, 0.88% lower evenness, and 2.07% 115 lower diversity. There were no significant associations when we pooled the phylum-level 116 metrics across studies. These results indicate that obese individuals do have a statistically 117 significant lower diversity than non-obese individuals; however, it is questionable whether 118 the difference is biologically significant. 119

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Relative risk. Building upon the alpha diversity analysis we calculated the relative risk of being obese based on an individual's alpha diversity metrics relative to the median metric for that study. Inspection of funnel plots for each of the metrics suggested that the studies included in our analysis were not biased (Figure S2). The results using relative risk largely matched those of using the raw alpha diversity data. Across the ten studies and six metrics, the only significant relative risk values were the richness, evenness, and diversity values from the Goodrich study (Figures 3 and S3). Again, although the relative risk values were not significant for other studies, the values tended to be above one. When we pooled the data using a random effects model, the relative risk associated with having a richness, evenness, or diversity below the median for the population was significantly associated with obesity (all P<0.0044). The relative risks associated with alpha diversity were small. The relative risk of having a low richness was 1.30 (95% CI: 1.13-1.49), low evenness was 1.20 (95% CI: 1.06-1.37), and low diversity was 1.27 (95% CI: 1.09-1.48). There were no significant differences in the phylum-level metrics. Again, the relative risk results indicate that individuals with a lower richness, evenness, or diversity are at statistically significant increased risk of being obese, it is questionable whether that risk is biologically or clinically relevant.

Beta diversity analysis. Following the approach used by the Walters and Finucane re-analysis studies, for each dataset we calculated a Bray-Curtis distance matrix to measure the difference in the membership and structure of the individuals from each study. We then used AMOVA to test for significant differences between the structure 140 of non-obese and obese individuals (Table 1). The Escobar, Goodrich, and Turnbaugh 141 datasets indicated a significant difference in community structure (all P<0.05). Because 142 it was not possible to ascertain the directionality of the difference in community structure 143 because the samples are arrayed in a non-dimensional space. nor perform a pooled 144 analysis using studies that had non-overlapping 16S rRNA gene sequence regions it is 145 unclear whether these differences reflect a broader, but perhaps small, shift in community 146 structure between non-obese and obese individuals. 147

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Development of a microbiome-based classifier of obesity. The Walters re-analysis study suggested that it was possible to classify individuals as being non-obese or obese 149 based on the composition of their microbiota. We repeated this analysis with additional 150 datasets using OTU and genus-level phylotype data. For each study we developed a 151 Random Forest machine learning model to classify individuals. Using ten-fold cross 152 validation, the observed AUC values varied between 0.52 and 0.69 indicating a relatively 153 poor ability to classify individuals (Figure 4A). So that we could test models on other 154 datasets, we trained models using genus-level phylotype data for each dataset. The 155 cross-validated AUC values for the models applied to the training datasets varied between 156 0.51 and 0.65, again indicating a relatively poor ability to classify individuals from the 157 original dataset (Figure 4B). For each model we identified the probability where the sum 158 of the sensitivity and specificity was the highest. We then used this probability to define 159 a threshold for calculating the accuracy of the models when applied to the other nine 160 datasets (Figure 5). Although there was considerable variation in accuracy values for each model, the median accuracy for each model varied between 0.33 (Turnbaugh) and 0.65 (HMP) (median=0.57). When we considered the number of samples, balance of non-obese and obese individuals, and region within the 16S rRNA gene it was not possible to identify factors that predictably affected model performance. The ability to predict obesity status using the relative abundance of OTUs and genera in the communities is only marginally better than random. These results suggest that given the large diversity of microbiome compositions it is difficult to identify a taxonomic signal that can be associated with obesity.

Power and Sample Size Estimate Simulations. The inability to detect a difference 169 between non-obese and obese individuals could be due to the lack of a true effect or 170 because the study had insufficient statistical power to detect a difference because of 171 insufficient sampling, large interpersonal variation, and unbalanced sampling of non-obese and obese individuals. To assess this, we calculated the power to detect differences of 1, 5, 10, and 15% in each of the alpha diversity metrics using the sample sizes used in each of the studies (Figures 6, S4-S9). Although there is no biological rationale for these effect sizes, they represent a range that includes effect sizes that would be generally considered to be biologically significant. Only the Goodrich study had power greater than 0.80 to 177 detect a 5% difference in Shannon diversity and six of the studies had enough power to 178 detect a 10% difference (Figure 6). None of the studies had sufficient power to detect a 179 15% difference between B:F values (Figure S4). In fact, the maximum power among any of 180 the studies to detect a 15% difference in B:F values was 0.25. Among the tests for relative 181 risk, none of the studies had sufficient power to detect a Cohen's d of 0.10 and only two 182 studies had sufficient power to detect a Cohen's d of 0.15. We next estimated how many 183 individuals would need to have been sampled to have sufficient power to detect the four 184 effect sizes assuming the observed interpersonal variation from each study and balanced 185 sampling between the two groups. To detect a 1, 5, 10, or 15% difference in Shannon 186 index, the median required sampling effort per group was approximately 3,400, 140, 35, 187 or 16 individuals, respectively. To detect a 1, 5, 10, and 15% difference in B:F values, the median required sampling effort per group was approximately 160,000, 6,300, 1,600, or 700 individuals, respectively. To detect a 1, 5, 10, and 15% difference in relative risk values

using Shannon diversity, the median required sampling effort per group was approximately 39,000, 1,500, 380, or 170 individuals, respectively. These estimates indicate that most microbiome studies are underpowered to detect modest effect sizes using either metric. In the case of obesity, the studies were underpowered to detect the 0.90 to 6% difference in diversity that was observed across the studies.

Discussion

Our meta-analysis helps to provide clarity to the ongoing debate of whether or not there are specific microbiome-based markers that can be associated with obesity. We performed an extensive literature review of the existing studies on the microbiome and obesity and performed a meta-analysis on the studies that remained based on our inclusion and exclusion criteria. By statistically pooling the data from ten studies, we observed significant, but small, relationships between richness, evenness, and diversity and obesity status as well as the relative risk of being obese based on these metrics. We also generated Random Forest machine learning models trained on each dataset and tested on the remaining datasets. This analysis demonstrated that the ability to reliably classify individuals as being obese based solely on the composition of their microbiome was limited. Finally, we assessed the ability of each study to detect defined differences in alpha diversity and observed that most studies were underpowered to detect modest effect sizes. Considering these datasets are among the largest published, it appears that most human microbiome studies are underpowered to detect differences in alpha diversity.

Alpha diversity metrics are attractive because they distill a complex dataset to a single value. For example, diversity is a measure of the entropy in a community and integrates richness and evenness information. Two communities with little taxonomic similarity can have the same diversity. Among ecologists the relevance of these metrics is questioned

because it is difficult to ascribe a mechanistic interpretation to their relationship with stability or disease. Regardless, the concept of a biologically significant effect size needs to be developed among microbiome researchers. Alternative metrics could include the ability to detect a defined difference in the relative abundance of an OTU representing a defined relative abundance. What makes for a biologically significant difference or relative abundance is an important point that has yet to be discussed in the microbiome field. The use of operationally defined effect sizes should be adequate until it is possible to decide upon an accepted practice.

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By selecting a range of possible effect sizes, we were able to demonstrate that most studies 223 are underpowered to detect modest differences in alpha diversity metrics and phylum-level relative abundances. Several factors interact to limit the power of microbiome studies. There is wide interpersonal variation in the diversity and structure of the human microbiome. Some factors such as relationship between subjects could potentially decrease the amount variation (6) and other factors such as whether one lives in a rural environment could increase the amount of variation (28). It was interesting that although there were ranges in 220 variation across the studies included in our analysis, the sample size calculations yielded 230 similar results. In addition, the common experimental designs limit their power. As we 231 observed, most of the studies included in our analysis were unbalanced for the variable that 232 we were interested in. This was also true of those studies that originally sought to identify 233 associations with obesity. Even with a balanced design, we showed that it was necessary 234 to obtain approximately 140 and 6,300 samples per group to detect a 5% difference in 235 Shannon diversity or B:F, respectively. It was interesting that these sample sizes agreed 236 across studies regardless of their sequencing method, region within the 16S rRNA gene, 237 or subject population (Figure 6). This suggests that regardless of the treatment or category, 238 these sample sizes represent a good starting point for subject recruitment when using stool samples. Unfortunately, few studies have been published with this level of subject recruitment. This is troubling since the positive predictive rate of a significant finding in an

underpowered study is small leading to results that cannot be reproduced (29). Future microbiome studies should articulate the basis for their experimental design.

Two previous re-analysis studies have stated that there was not a consistent association 244 between alpha diversity and obesity (8, 9); however, neither of these studies made an 245 attempt to pool the existing data together to try and harness the additional power that 246 this would give and they did not assess whether the studies were sufficiently powered 247 to detect a difference. Additionally, our analysis used 16S rRNA gene sequence data 248 from ten studies whereas the Finucane study used 16S rRNA gene sequence data from three studies (7, 10, 21) and a metagenomic study (30) and the Walters study used 250 16S rRNA gene sequence data from five studies (10, 15, 20, 21, 28); two studies were included in both analyses (10, 21). Our analysis included four of these studies (10, 15, 20, 21) and excluded three of the studies because they were too small (7), only utilized 253 metagenomic data (30), or used short single read Illumina HiSeq data that has a high error rate making it untractable for de novo OTU clustering (28). The additional seven 255 datasets were published after the two reviews were performed and include datasets with 256 more samples than were found in the original studies. Our collection of ten studies allowed 257 us to largely use the same sequence analysis pipeline for all datasets and relied heavily 258 on the availability of public data and access to metadata that included variables beyond 259 the needs of the original study. To execute this analysis, we created an automated data 260 analysis pipeline, which can be easily updated to add additional studies as they become 261 available (https://github.com/SchlossLab/Sze Obesity mBio 2016/). Similarly, it would be 262 possible to adapt this pipeline to other body sites and treatment or variables (e.g. subject's 263 sex or age). 264

Similar to our study, the Walters study generated Random Forest machine learning models to differentiate between non-obese and obese individuals (8). They obtained similar AUC values to our analysis; however, they did not attempt to test these models on the other

studies in their analysis. When we performed the inter-dataset cross validation the median accuracy across datasets was only 56.68% indicating that the models did a poor job when applied to other datasets. This could be due to differences in subject populations and methods. Considering the median AUC for models trained and tested on the same data with ten-fold cross validation only varied between 0.51 and 0.65 and there was not a strong signal in the alpha diversity data, we suspect that there is insufficient signal to reliably classify individuals.

Although we failed to find an effect this does not necessarily mean that there is no role for the microbiome in obesity. There is strong evidence in murine models of obesity that the microbiome and level of adiposity can be manipulated via genetic manipulation of the animal and manipulation of the community through antibiotics or colonizing germ free mice with diverse fecal material from human donors (5, 10–13). These studies appear to conflict with the observations using human subjects. Recalling the large interpersonal variation in the structure of the microbiome, it is possible that each individual has their own signatures of obesity. Alternatively, it could be that the involvement of the microbiome in obesity is not apparent based on the taxonomic information provided by 16S rRNA gene sequence data. Rather, the differences could become more apparent at the level of a common set of gene transcripts or metabolites that can be produced from different structures of the microbiome.

Methods

Sequence Analysis Pipeline. All sequence data were publicly available and were downloaded from the NCBI Sequence Read Archive, the European Nucleotide Archive, or the investigators' personal website (https://gordonlab.wustl.edu/TurnbaughSE/_10/_09/STM/_2009.html). In total seven studies used 454 (6, 15, 16, 18, 20–22) and three studies used Illumina sequencing (17, 19, 23). All of these studies used amplification-based

16S rRNA gene sequencing. Among the studies that sequenced the 16S rRNA gene, the researchers targeted the V1-V2 (20), V1-V3 (15, 16, 18), V3-V5 (21, 22), V4 [(19); (23);], and V3-4 (17) regions. For those studies where multiple regions were sequenced, we selected the region that corresponded to the largest number of subjects (6, 21). We 295 processed the 16S rRNA gene sequence data using a standardized mothur pipeline. Briefly, 296 our pipelines attempted to follow previously recommended approaches for 454 and Illumina 297 sequencing data (24, 25). All sequences were screened for chimeras using UCHIME and 298 assigned to operational taxonomic units (OTUs) using the average neighbor algorithm 299 using a 3% distance threshold (26, 31). All sequence processing was performed using 300 mothur (v.1.37.0) (32). 301

Data Analysis. We split the overall meta-analysis into three general strategies using R (3.3.0). First, we followed the approach employed by Finucane et al (9) and Walters et al 303 (8) where each study was re-analyzed separately to identify associations between BMI and the relative abundance of Bacteroidetes and Firmicutes, the ratio of Bacteroidetes 305 and Firmicutes relative abundances (B:F), Shannon diversity, observed richness, and 306 Shannon evenness (33). After each variable was transformed to fit a normal distribution 307 a two-tailed t-test was performed for comparison of non-obese and obese individuals 308 (i.e. BMI > 35.0). We performed a pooled analysis on these measured variables using 309 linear random effect models to correct for study effect to asses differences on the combined 310 dataset between non-obese and obese groups using the Ime4 (v.1.1-12) R package (34). 311 Next, we compared the community structure from non-obese and obese individuals using 312 analysis of molecular variance (AMOVA) with Bray-Curtis distance matrices (35). This 313 analysis was performed using the vegan (v.2.3-5) R package. For both analyses, the 314 datasets were rarefied (N=1000) so that each study had the same number of sequences. 315 Second, for each study we partitioned the subjects into a low or high group depending 316 on whether their alpha diversity metrics were below or above the median value for the 317 study. The relative risk (RR) was then calculated as the ratio of the number of obese

individuals in the low group to the number of obese individuals in the high group. We then performed a Fisher exact-test to investigate whether the RR was significantly different from 1.0 within each study and across all of the studies using the epiR (0.9-77) and metafor (1.9-8) packages. Third, we used the AUCRF (1.1) R package to generate Random Forest 322 models (36). For each study we developed models using either OTUs or genus-level 323 phylotypes. The quality of each model was assessed by measuring the area under the 324 curve (AUC) of the Receiver Operating Characteristic (ROC) using ten-fold cross validation. 325 Because the genus-level phylotype models were developed using a common reference, it 326 was possible to use one study's model (i.e. the training set) to classify the samples from 327 the other studies (i.e. the testing sets). The optimum threshold for the training set was set 328 as the probability threshold that had the highest combined sensitivity and specificity. This 329 threshold was then used to calculate the accuracy of the model applied to the test studies. 330 To generate ROC curves and calculate the accuracy of the models we used the pROC (1.8) 331 R package (37). Finally, we performed power and sample number simulations for different 332 effect sizes for each study using the pwr (1.1-3) R package and base R functions. We also 333 calculated the actual sample size needed based on the effect size of each individual study. 334 Reproducible methods. A detailed and reproducible description of how the data were 335 processed and analyzed can be found at https://github.com/SchlossLab/Sze_Obesity_ 336 mBio_2016/. 337

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- Figure 1: PRISMA flow diagram of total records searched (38).
- Figure 2: Individual and combined comparison of obese and non-obese groups for

 Shannon diversity (A) and B:F (B).
- Figure 3: Meta analysis of the relative risk of obesity based on Shannon diversity
 (A) or B:F (B).
- Figure 4: ROC curves for each study based on classification of non-obese or obese groups using OTUs (A) or genus-level classification (B).
- Figure 5: Overall accuracy of each study to predict non-obese and obese individuals based on that study's Random Forest machine learning model applied to each of the other studies.
- Figure 6: Power (A) and sample size simulations (B) for Shannon diversity for differentiating between non-obese versus obese for effect sizes of 1, 5, 10, and 15%. Power calculations use the sampling distribution from the original studies and the sample size estimations assume an equal amount of sampling from each treatment group.

- Figure S1: Individual and Combined comparison of Obese and Non-Obese groups
 Based on Evenness (A), Richness (B), or the Relative Abundance of *Bacteroidetes*(C) and Firmictues (D).
- Figure S2: Funnel plots depicting the general lack of bias in the selection of datasets included in the analysis.
- Figure S3: Meta Analysis of the Relative Risk of Obesity Based on Evenness (A),
 Richness (B), or the Relative Abundance of *Bacteroidetes* (C) and Firmictues (D).
- Figure S4: Power (A) and sample size simulations (B) for B:F for differentiating between non-obese versus obese for effect sizes of 1, 5, 10, and 15%. Power calculations use the sampling distribution from the original studies and the sample size estimations assume an equal amount of sampling from each treatment group.
- Figure S5: Power (A) and sample size simulations (B) for richness for differentiating
 between non-obese versus obese for effect sizes of 1, 5, 10, and 15%. Power
 calculations use the sampling distribution from the original studies and the sample size
 estimations assume an equal amount of sampling from each treatment group.
- Figure S6: Power (A) and sample size simulations (B) for evenness for differentiating between non-obese versus obese for effect sizes of 1, 5, 10, and 15%. Power calculations use the sampling distribution from the original studies and the sample size estimations assume an equal amount of sampling from each treatment group.
- Figure S7: Power (A) and sample size simulations (B) for the relative abundance of

 Bacteroidetes for differentiating between non-obese versus obese for effect sizes

 of 1, 5, 10, and 15%. Power calculations use the sampling distribution from the original

 studies and the sample size estimations assume an equal amount of sampling from each

- 383 treatment group.
- Figure S8: Power (A) and sample size simulations (B) for the relative abundance of

 Firmicutes for differentiating between non-obese versus obese for effect sizes of

 1, 5, 10, and 15%. Power calculations use the sampling distribution from the original

 studies and the sample size estimations assume an equal amount of sampling from each

 treatment group.
- Figure S9: Power (A) and sample size simulations (B) for relative risk of obesity
 based on Shannon diversity. Power calculations use the sampling distribution from the
 original studies and the sample size estimations assume an equal amount of sampling
 from each treatment group.

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