ASSOCIATING MICROBIOME COMPOSITION WITH ENVIRONMENTAL COVARIATES USING GENERALIZED UNIFRAC DISTANCES

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ABSTRACT

Motivation: The human microbiome plays an important role in human disease and health. Identification of factors that affect the microbiome composition can provide insights into disease mechanism as well as suggest ways to modulate the microbiome composition for therapeutic purposes. Distance-based statistical tests have been applied to test the association of microbiome composition with environmental or biological covariates. The unweighted and weighted UniFrac distances are the most widely used distance measures. However, these two measures assign too much weight either to rare lineages or to most abundant lineages, which can lead to loss of power when the important composition change occurs in moderately abundant lineages.

Results: We develop generalized UniFrac distances that extend the weighted and unweighted UniFrac distances for detecting a much wider range of biologically relevant changes. We evaluate the use of generalized UniFrac distances in associating microbiome composition with environmental covariates using extensive Monte Carlo simulations. Our results show that tests using the unweighted and weighted UniFrac distances are less powerful in detecting abundance change in moderately abundant lineages. In contrast, the generalized UniFrac distance is most powerful in detecting such changes, yet it retains nearly all its power for detecting rare and highly abundant lineages. The generalized UniFrac distance also has an overall better power than the joint use of unweighted/weighted UniFrac distances. Application to two real microbiome datasets has demonstrated gains in power in testing the associations between human microbiome and diet intakes and habitual smoking.

Availability: http://cran.r-project.org/web/packages/GUniFrac

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1 INTRODUCTION

Understanding the compositional differences of microbial communities (microbiomes) is essential in microbial ecology. With the development of next-generation sequencing technologies, microbiome composition can now be determined by direct DNA sequencing without the need for laborious cultivation (Dinsdale et al. 2007; Tett et al. 2009; Grace et al. 2009; Wu et al. 2011; Tringe et al. 2009; Tumbolari et al. 2009; von Mering et al. 2009). There has been a great interest in human microbiome studies in different body sites, ranging from skin (Grace et al. 2009; Arumugam et al. 2011; Muegge et al. 2011; Qin et al. 2010; Wu et al. 2011) and respiratory tract (Charlson et al. 2011). Important insights have been gained from analysis of large-scale human microbiome data, including the discovery of enterotypes (Arumugam et al. 2011) and discovery of the link between diet and these enterotypes (Wu et al. 2011). Although the metagenomic shotgun approach is potentially more powerful and unbiased, 16S rRNA gene targeted sequencing is routinely performed to determine the taxonomic composition. The generated 16S rRNA sequence tags are usually clustered into operational taxonomic units (OTUs) with a specified amount of variability allowed within each OTU (Charlson et al. 2010a; Wu et al. 2010). At 97% similarity, these OTUs represent ‘species’. Downstream analysis is then performed on the OTU abundance data.

Two central themes in human microbiome studies are to identify potential biological and environmental factors that are associated with microbiome composition, and to define the relationship between microbiome features and biological or clinical outcomes. The goal is to provide a better understanding of the factors that shape our microbiome and, potentially, contribute to the development of new therapeutic strategies to modulate the microbiome composition and affect human health (Spor et al. 2011; Virgin and Told 2011). Testing the association of microbiome composition with potential environmental factors using OTU abundances directly is difficult due to high dimensionality, non-normality and phylogenetic structure of the OTU data. Instead, distance-based non-parametric testing, in which a distance measure is defined between any two microbiomes, is usually used to achieve this goal (Charlson et al. 2010; Fukuyama et al. 2010; Kuczynski et al. 2010; Wu et al. 2010). The power of the distance-based test depends on a proper choice of the distance measure. Numerous distance measures have been proposed to compare microbial communities (Kuczynski et al. 2010b; Swenson 2011). Phylogenetic distance measures, which account for the phylogenetic relationship among the species, provide far more power because they exploit the degree of divergence between different sequences. Among these, the UniFrac distances are the most popular ones (Lozupone and
Consider two microbiome communities.

2 METHODS

2.1 Generalized UniFrac distances between two microbial communities

Consider two microbiome communities A and B and suppose that we have a rooted phylogenetic tree with $n$ branches. Let $h_i$ be the length of the branch $i$ and $p_A^i$ and $p_B^i$ are the taxa proportions descending from the branch $i$ for community $A$ and $B$, respectively. The unique fraction metric, or UniFrac, measures the phylogenetic distance between sets of taxa in a phylogenetic tree as the fraction of the branch length of the tree that leads to descendants from either one environment or the other, but not both. The original definition refers to unweighted UniFrac (Lozupone and Knight, 2005; Lozupone et al., 2011), which is mathematically defined as

$$d_{U} = \frac{\sum_{i=1}^{n} h_i |p_A^i > 0 > p_B^i|}{\sum_{i=1}^{n} h_i}$$

where $h_i$ is the indicator function and only presence/absence of species of branch $i$, $p_A^i > 0$ and $p_B^i > 0$, are used in the definition. The distance definition $d_U$ completely ignores the taxa abundance information. In contrast, the (normalized) weighted UniFrac distance $d_{UW}$ weights the branch length with abundance difference and is defined as

$$d_{UW} = \frac{\sum_{i=1}^{n} h_i |p_A^i - p_B^i|}{\sum_{i=1}^{n} h_i}$$

Note that $d_{UW}$ cannot be reduced to $d_U$ even if we convert abundance data into presence/absence data. Also note that $d_U$ uses the absolute proportion difference $|p_A^i - p_B^i|$ in its formulation. The consequence of using the absolute difference is that the value of $d_U$ is determined mainly by branches with large proportions and is less sensitive to the abundance changes on the branches with small proportions. To attenuate the weight on branches with large proportions, we may instead use the relative difference $|p_A^i - p_B^i|/p_A^i + p_B^i$ ($\in [0, 1]$) in the formulation. We denote this distance measure as

$$d_{UWR} = \frac{\sum_{i=1}^{n} h_i |p_A^i - p_B^i|/p_A^i + p_B^i}{\sum_{i=1}^{n} h_i}$$

where $\sum_{i=1}^{n} h_i$ in the denominator is the normalizing factor so that $d_{UWR}$ $\in [0, 1]$. Now if we dichotomize the abundance data using the indication function $h_i$, $d_{UWR}$ is reduced to $d_U$. So $d_{UWR}$ can be seen as a ‘weighted version’ of $d_U$. Using the relative differences, we place equal emphasis on every branch and the distance is not dominated by the branches with large proportions, since the relative difference does not depend on the magnitude of $p_A^i, p_B^i$. However, the low-abundance branches may be more noisy and the relative difference may amplify such noises. To strike a balance between relative difference and absolute difference, we weight the branch length both by the relative difference and its importance indicated by the branch proportion. We propose the following generalized UniFrac distances

$$d_{UW} = \frac{\sum_{i=1}^{n} h_i |p_A^i - p_B^i|}{\sum_{i=1}^{n} h_i}$$

where $\alpha \in [0, 1]$ controls the contribution from high-abundance branches, and $\sum_{i=1}^{n} h_i |p_A^i + p_B^i|$ is the normalizing factor so that $d_{UWR}^\alpha$ $\in [0, 1]$. Branches with zero proportions for both communities will not be included in the calculation. As $\alpha$ changes from 0 to 1, more emphasis is placed on high-abundance branches. When $\alpha = 1$, $d_{UWR}^\alpha$ is reduced to $d_{UWR}$. When $\alpha = 0$, we get $d_U$ defined above.

Therefore, by varying $\alpha$ from 1 to 0 , we achieve a series of distances ranging from $d_{U}$ to $d_{UWR}^{\alpha}$. Note that $d_{UWR}^{\alpha}$ is obtained by dichotomizing the abundance in $d_{UWR}$, but is different from $d_{UWR}^{\alpha}$. We are particularly interested in $d_{UWR}^{\alpha}$, the distance in the middle of the distance series

$$d_{UWR}^{\alpha} = \frac{\sum_{i=1}^{n} h_i \sqrt{p_A^i + p_B^i}}{\sum_{i=1}^{n} h_i}$$

We also compare $d_{U}, d_{UWR}^{\alpha}, d_{UWR}^{\alpha}$ and $d_{UWR}^{\alpha}$ with VAW-UniFrac distance $d_{UWR}^{\alpha}$, which is defined as:

$$d_{UWR}^{\alpha} = \frac{\sum_{i=1}^{n} h_i |p_A^i - p_B^i|}{\sum_{i=1}^{n} h_i}$$

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$$d_{UWR}^{\alpha} = \frac{\sum_{i=1}^{n} h_i |p_A^i - p_B^i|}{\sum_{i=1}^{n} h_i}$$


where $n_i$ is the total number of individuals reads from both communities on the $i$th branch and $m$ is total number of individuals reads.

### 2.2 Statistical test based on UniFrac distances

We study the power of generalized UniFrac distances using the distance-OTU abundances. Each replication consists of drawing 400 points from Generalized UniFrac distances are then calculated based on the NJ tree and rooted by midpoint rooting method. (Neighbor Joining, 'nj' function in R) and using a permutation test with pseudo-F ratios to obtain the P-values. The pseudo-F statistics is defined as:

$$F = \frac{\nu \langle H \rangle_{(m-1)}}{\nu \langle H \rangle_{(m-1)} - \langle H \rangle_{(m-2)}},$$

where $(\cdot)$ is the trace function of a matrix. $H = XX'X'$ of the hat (projection) matrix of the design matrix $X$. $G$ is Gower's centered matrix and $m$ and $n$ is the number of samples and the number of predictors, respectively. Let $d_r$ be the generalized UniFrac distance between community $r$ and $s$ and denote $A = (a_{i,j}) = (1 - d_{i,j}^2)$. The Gower's matrix is defined as $G = (1 - H/n, 1 - H/n)$, where $1$ is a vector of 1's.

Since $d^0$ and $d^θ$ reflect the abundance change in either rare lineages or abundant lineages, combining $d^0$ and $d^θ$ may potentially increase the overall power. Instead of applying Bonferroni correction to the P-values from separate PERMANOVA tests using $d^0$ or $d^θ$ to control the family-wise Type I error rate, a more powerful approach is to take the maximum of pseudo-F statistics for $d^0$ and $d^θ$ as a new test statistic. The significance of the pseudo-F statistics is assessed based on permutations.

### 2.3 Simulation strategies

We use two simulation strategies to evaluate the power of the generalized UniFrac distances under various conditions. The first strategy is a modification of the simulation method proposed by Schloss (2008), where we draw points (16S rRNA sequences) from a 2D circle with known densities (Fig. 2B). This strategy facilitates simulations of different community characteristics such as species evenness and species richness. The Euclidean distance between points is analogous of the genetic distance between the sequences. The diameter of the circle represents the maximum genetic variation, fits linear models to distance matrices and uses a permutation test with pseudo-F ratios to obtain the P-values. The pseudo-F statistics is defined as:

$$F = \frac{\nu \langle H \rangle_{(m-1)}}{\nu \langle H \rangle_{(m-1)} - \langle H \rangle_{(m-2)}},$$

where $(\cdot)$ is the trace function of a matrix. $H = XX'X'$ of the hat (projection) matrix of the design matrix $X$. $G$ is Gower's centered matrix and $m$ and $n$ is the number of samples and the number of predictors, respectively. Let $d_r$ be the generalized UniFrac distance between community $r$ and $s$ and denote $A = (a_{i,j}) = (1 - d_{i,j}^2)$. The Gower's matrix is defined as $G = (1 - H/n, 1 - H/n)$, where $1$ is a vector of 1's.

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### 3 RESULTS

#### 3.1 Comparison of the power of different UniFrac variants using 2D circle-based simulations

We use PERMANOVA to test for environmental effect and compare the power of $d^0$, $d^{0.5}$, $d^{θ}$, $d^θ$ and $d^{θθ}$. Specifically, we simulate two environmental conditions (e.g. smoking versus non-smoking) under which we draw 10 samples each. We then vary the degree of community difference under these two conditions and produce the power curve over a grid of 10 for each UniFrac distance. We investigate six scenarios, where the environmental factor affects the community membership, species evenness, species richness, most abundant lineages moderately abundant lineages and rare lineages (Fig. 2B–G). For each scenario, we vary one community characteristic (Supplementary Table S1). Suppose $x_1$ and $x_2$ are the mean values of the community characteristic for Conditions 1 and 2. We simulate 10 communities for each condition with community
Two simulation strategies to evaluate the generalized UniFrac distances. (A–G), 2D circle-based simulation of microbial communities with different characteristics. (A) The microbial community is represented by a 2D circle. Points are drawn from the circle to simulate the 16S-based sampling process. These points are further binned into small hexagons as OTUs. UPGMA or NJ method is used to build the OTU phylogenetic tree. Six scenarios are investigated, where the difference occurs in: community membership (B), evenness (C), richness (D), most abundant lineages (E), moderately abundant lineages (F) and rare lineages (G). The affected lineages are indicated by a red circle or ring. H, tree-based simulation of microbial communities based on the phylogenetic tree and DM model. A real OTU phylogenetic tree from a throat microbial community dataset is used. These OTUs are roughly divided into 20 clusters (lineages) by performing PAM method using the OTU patristic distance matrix. Each cluster is subjected to abundance change in response to the environment. Counts are generated from a DM model.

Each community is sampled once. Initially, we let \( x_1 = x_2 \) (no difference) and then increase the difference between \( x_1 \) and \( x_2 \) to simulate stronger environmental effect. PERMANOVA is then performed on the distance matrices and the power curve is created over a grid of 10 using Type I error \( \alpha = 0.05 \). Figure 2 shows the power curves for different UniFrac distances under the six scenarios considered. When the environmental factor has no effect (\( x_1 = x_2 \)), PERMANOVA controls the Type I error at the nominal level of 0.05 for all five UniFrac distances. As the environmental effect becomes stronger, all the distances have better power. When the environmental factor affects the community membership or richness (Panels 1 and 3), all the distances give a similar power and their power curves are nearly identical. For the evenness change scenario (Panel 2), the power of \( d_{W} \) and \( d_{(0.5)} \) is very close and is more powerful than \( d_{(0)} \) and \( d_{U} \). \( d_{W} \) is the most powerful for detecting change in most abundant lineages (Panel 4) but is much less powerful for change in rare lineages (Panel 6). \( d_{(0)} \) shows an opposite trend: it is the most powerful for detecting change in rare lineages (Panel 6) but has almost no power for change in most abundant lineages (Panel 4). In contrast, \( d_{(0.5)} \) is the most powerful for detecting change in moderately abundant lineages (Panel 5). They are also the most robust among the distances investigated: its power is close to the best UniFrac distance under all scenarios. The performance of \( d_{(0)} \) lies between \( d_{(0.5)} \) and \( d_{U} \) and is also very robust. Finally, under the 2D circle simulations, the performance of \( d_{VAW} \) is almost identical to that of \( d_{(0.5)} \).

In the above simulations, we use a bin size of 0.015 to form ‘OTU’s (\( \sim 300 \) OTUs per sample). To study the effect of bin size, we compare the power curves of the generalized UniFrac distances using a smaller bin size of 0.01 (\( \sim 700 \) OTUs per sample) or a larger bin size of 0.03 (\( \sim 80 \) OTUs per sample). The bin size does not change the general conclusion (Supplementary Fig. S2). To study the effect of tree construction methods, we also construct the
A recently proposed variance adjusted UniFrac distance (difference) to cluster under Condition 1 is. Suppose that the proportion of the factor causes an increase of the abundance of a particular OTU in each replication and calculate the power over 2000 replications. We also report the power for the test combining \(d^W\) and \(d^{U}\) by taking the maximum of their pseudo-F statistics. We denote this method as \(d_{\text{MAX}}\). Figure 2 (left plot) shows that \(d^{W}\) has the lowest overall power and \(d^{(0.5)}\) and \(d_{\text{MAX}}\) have the best power, indicating \(d^{W}\) has a reciprocal relationship and neither of them is as robust as \(d^{(0.5)}\). Figure 2 shows the power curves of four representative cases. As the proportion of the affected cluster decreases from 19.7% to 0.9%, \(d^{W}\) becomes less powerful and the power of \(d^{U}\) has the opposite trend.

In the simulations presented above, the power is calculated assuming the affected cluster is known. Since the affected cluster can be abundant or rare, we randomly choose an affected OTU cluster in each replication and calculate the power over 2000 replications. We find that the power is higher for the tests combining \(d^{W}\) and \(d^{U}\) than the power of the tests combining \(d^{(0)}\) and \(d^{(10)}\). For Condition 2, the proportion of the environmental effect on most abundant lineages, \(d^{(0)}\) becomes less powerful and the power of \(d^{(10)}\) has the opposite trend.

3.2 Comparison of the power of different UniFrac variants using tree-based simulations

We also compare the power of different UniFrac distances for detecting environmental effect using tree-based simulations that mimic the throat microbiome data (see Section 3.2 for details).

A recently proposed variance adjusted UniFrac distance \(d_{\text{VAW}}\) is also compared in this setting. \(d_{\text{VAW}}\) was developed to moderate the branch proportion difference by its variance and was shown to increase the power of detecting the difference between two microbial communities. We use the phylogenetic tree of the 856 OTUs from the throat microbiome dataset and divide them into 20 clusters (Fig. 1H). The mean OTU proportions and the dispersion parameter are estimated from the real data by fitting a DM model. We assume that the environmental factor causes an increase of the abundance of a particular OTU cluster. Specifically, suppose that the proportion of the ith OTU cluster under Condition 1 is \(p_i\). For Condition 2, the proportion of ith OTU cluster is increased by \(k\) fold where \(k\) varies from 1 (no difference) to \(1/k\) (strong effect) on a grid of 10. The proportion vector is re-normalized to sum to 1. Next, 10 samples are simulated for each condition with their OTU counts generated by the DM model with the corresponding proportion vector and the common dispersion parameter. As expected, the five UniFrac distances differ in their power for detecting environmental effect for the 20 OTU clusters tested. Except for \(d^{(0)}\), all the UniFrac distances have their best-performance scenarios. \(d^{W}\), \(d^{(0.5)}\), \(d^{U}\) and \(d_{\text{VAW}}\) achieve the highest power in seven, six, three and one cases, respectively. For the remaining three cases, \(d^{W}\) and \(d^{(0.5)}\) are equally the most powerful (Supplementary Fig. S3).

The results are consistent with the 2D circle-based simulation: \(d^{W}\) is most powerful in detecting the environmental effect on most abundant lineages, \(d^{(0.5)}\) is most powerful for moderately abundant lineages and \(d^{U}\) is most powerful for rare lineages. In contrast, performance of the test with \(d^{(10)}\) and \(d_{\text{VAW}}\) is generally between \(d^{W}\) and \(d^{(0.5)}\). The power of \(d^{(10)}\) and \(d_{\text{VAW}}\) has a reciprocal relationship and neither of them is as robust as \(d^{(0.5)}\). Finally, we assume that the environmental factor affects a random set of 40 OTUs instead of a random OTU cluster. At this extreme where the phylogenetic relationship is no longer important, \(d^{(0.5)}\) has even higher power than the other distances, followed by \(d^{(10)}\) and \(d^{(0)}\) and \(d^{(10)}\) and \(d^{(0.5)}\) are equally the most powerful. In the simulations presented above, the power is calculated assuming the affected cluster is known. Since the affected cluster can be abundant or rare, we randomly choose an affected OTU cluster in each replication and calculate the power over 2000 replications. We also report the power for the test combining \(d^{W}\) and \(d^{U}\) by taking the maximum of their pseudo-F statistics. We denote this method as \(d_{\text{MAX}}\). Figure 2 (left plot) shows that \(d^{W}\) has the lowest overall power and \(d^{(0.5)}\) and \(d_{\text{MAX}}\) have the best power, indicating \(d^{W}\) has a reciprocal relationship and neither of them is as robust as \(d^{(0.5)}\). Overall, \(d^{(0.5)}\) has a better power than other UniFrac distances including the one that combines \(d^{W}\) and \(d^{U}\).
or replications. It might want to include all possible associations for further validation of nutrient-microbiome associations, there are situations when one test show the joint use of \( d^W \) and \( d^{UI} \), does not increase the power over \( d^W \), indicating most associations can be recovered by \( d^W \) alone.

### 3.4 Results from analysis of a throat microbiome dataset of smokers and non-smokers

Cigarette smokers have an increased risk of multiple diseases, including upper respiratory tract infections. Previous studies had linked smoking to specific respiratory tract bacteria, but the consequences of smoking for global airway microbial community composition had not been fully clarified. Charlson et al. (2010), investigated the smoking effect on the oropharyngeal and nasopharyngeal bacterial communities using 454 pyrosequencing of 16S sequence tags. Specifically, a total of 291 swab samples from the right and left nasopharynx and oropharynx of 29 smoking and 33 non-smoking healthy asymptomatic adults were collected. The variable region 1-2 (V1-V2) of the bacterial 16S rRNA gene was PCR-amplified using individually barcoded primer and subjected to multiplexed pyrosequencing. The pyrosequences were denoised by Quince et al. (2008) prior to taxonomic assignment and yielded an average of 1335 ± 603 (SD) reads per airway sample. The denoised sequences were then analyzed using the QIIME pipeline (Caporaso et al., 2010) with default parameter setting. We use the left oropharyngeal samples in this study. After removing two samples with read number < 500 and discarding singleton OTUs, i.e. OTU with only one read, we finally have an OTU table of 60 samples (28 smokers versus 32 non-smokers) and 856 OTUs.

We test the smoking effect on the throat microbial community composition by applying PERMANOVA (10000 permutations). All the five UniFrac distances achieve statistical significance at \( \alpha = 0.05 \) level, indicating smoking alters the community composition. However, test using \( d^{(0.5)} \) produces the smallest \( P \)-value of 0.006, followed by 0.008 from \( d^{(0.2)} \). The \( P \)-values based on \( d^W \), \( d^{UI} \) and \( d^{UIW} \) are 0.012, 0.019 and 0.043, respectively. We also perform a principle coordinate analysis using the distance matrices and plot the samples on the first two principle coordinates (Fig. 5). The distance \( d^{(0.5)} \) separates the samples better than the other

Fig. 3. Power comparison of different UniFrac variants for detecting environmental effect using tree-based simulation. PERMANOVA is used for testing hypotheses. 214 nutrients are included in the testing. The curves are generated by varying the \( P \)-value cutoffs.

![Diagram](https://example.com/diagram.jpg)

**Legend:**
- \( d^W \)
- \( d^{UI} \)
- \( d^{UIW} \)
- \( d^{MAX} \)

**Note:** The power curves are created by varying the \( P \)-value cutoffs to detect the joint use of \( d^W \) and \( d^{UI} \), does not increase the power over \( d^W \), indicating most associations can be recovered by \( d^W \) alone.
Testing association of microbiome composition with environmental covariates is considered. In addition, the generalized UniFrac distances are very comparable to the best UniFrac distances among the scenarios we considered. In addition, the generalized UniFrac distances are very robust to tree constructing methods. We suggest the use of $d^{(0.5)}$ for testing association of microbiome composition with environmental covariates to avoid missing important findings.

Both weighted and unweighted UniFrac distances are sensitive to sampling depth (Lozupone et al., 2011). Inflated distances at a low sampling depth are caused by sampling variation especially for the rare lineages. The generalized UniFrac distances are also sensitive to sampling depth (Supplementary Fig. S5). However, as the sampling depth increases, the distance stabilizes. For the gut microbiome dataset, we found a sequencing depth of $\sim 1000$ reads is sufficient to stabilize the generalized UniFrac distances. To overcome the potential adverse effects of uneven sampling, rarefaction is usually used to subsample the samples to the same depth. However, when the sampling depth varies greatly across the samples, rarefaction throws away a significant portion of the 16S reads and increases the sampling variation. We found that rarefaction is not necessary, at least, in the context of testing the association of the microbiome composition with covariates (Supplementary Fig. S6).

The power of UniFrac variants can also be compared in the context of testing whether two microbial communities differ significantly as in (Cham et al., 2011; Schloss, 2009). Instead of comparing power for detecting the difference between two communities, we focus our evaluations on the performance of UniFrac distances for associating microbiome composition to environmental covariates by collecting multiple independent samples. The rationale is that as the sequence depth increases, two sample comparison will have increased power to detect differences due to sources that we are not interested in (random noises), such as the individual-to-individual variability, day-to-day variability, sampling location variability or even technical variability (e.g. sample preparation). Multiple samples from a population coupled with multivariate statistical methods such as the distance-based PERMANOVA provide powerful design and analysis methods to overcome these potential random noises (Lozupone et al., 2011). As more and more large-scale microbiome datasets are being collected, we expect that our generalized UniFrac distances can help to identify important covariates that are associated with the microbiomes that could be missed using the commonly used UniFrac distances. In addition to identifying environmental covariates that may be determinants of microbiome composition, our approach would be equally suited to identifying microbiome features associated with biological or clinical outcomes, which is needed to begin to understand the impact of the microbiome on health.

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