The power of microbiome studies: some considerations on which alpha and beta metrics to use and how to report analysis the results

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The power of microbiome studies: some considerations on which alpha and beta metrics to use and how to report analysis the results

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ABSTRACT

Since sequencing techniques become less expensive, larger sample sizes are applicable for microbiota studies. The aim of this study is to show how, and to what extent, different diversity metrics and different compositions of the microbiota influence the needed sample size to observed dissimilar groups. Empirical 16S rRNA amplicon sequence data obtained from animal experiments, observational human data, and simulated data was used to perform retrospective power calculations. A wide variation of alpha diversity and beta diversity metrics were used to compare the different microbiota data sets and the effect on the sample size. Our data showed that beta diversity metrics are most sensitive to observe differences compared to alpha diversity metrics. The structure of the data influenced which alpha metrics are most sensitive. Regarding beta diversity, the Bray-Curtis metric is in general most sensitive to observe differences between groups, resulting in lower sample size and potential publication bias. We recommend to perform power calculations and to use multiple diversity metrics as an outcome measure. To improve microbiota studies awareness needs to be raised on the sensitivity and bias for microbiota research outcomes created by the used metrics rather than biological differences. We have seen that different alpha and beta diversity metrics lead to different study power: on the basis of this observation, one could be naturally tempted to try all possible metrics until one or more are found that give a statistically significant test result, i.e. $p$-value $< \alpha$. This way of proceeding is one of the many forms of the so-called $p$-value hacking. To this end, in our opinion, the only way to protect ourselves from (the temptation of) $p$-hacking would be to publish, and we stress here the word publish, a statistical plan before experiments are initiated: this practice is customary for clinical trials where a statistical plan describing the endpoints and the corresponding statistical analyses must be disclosed before the start of the study.

Keywords: Microbiota, power analysis, multivariate analysis, sample size
INTRODUCTION

For a few decades now, researchers have left culture-based methods and used molecular technologies, and more recently mostly sequencing-based approaches, to characterize microbial communities within a certain environment, referred to as the microbiome. In humans and animals, the microbiome has an important role in health and disease. For example, animals raised without or fewer microbes showed an underdeveloped immune system and are more susceptible to diseases [1-3]. Microbiome studies have as goal to investigate, characterize and understand the compositional and functional variability of microorganisms. The question “What is different?” between different groups of interests can be translated into an hypothesis testing procedure.

Hypothesis testing rests on the definition and choice of four parameters: i) the effect size, i.e. the quantification of the outcome of interest (in the simple case the difference between two groups), ii) the sample size \( n \), i.e. the number of samples (to be) collected; iii) the power of tests \( 1-\beta \), i.e. the probability of the test of rejecting the Null hypothesis when actually false and iv) the confidence level \( \alpha \), i.e. the probability of rejecting the Null hypothesis when actually true.

It is necessary to perform power analysis before performing experiments. This is well acknowledged in all fields of research, however, microbiome studies are challenged with conflicting results [4]. Under-powering and the failure to correct for false positives are among the causes underlying the lack of reproducibility of many biological findings [5, 6]. The power of a test is linked to the probability \( \beta \) of accepting the Null hypothesis when actually false (False negative error or Type II error), and \( \alpha \) describes the False positive error or Type I error. Once acceptable error rates \( \alpha \) (usually 0.05 or 0.01) and \( \beta \) (usually 0.2, although context-dependent), and the effect that one is interested to assess statistically are chosen, it is possible, at least in principle, to determine the optimal sample size, i.e. the number of samples that one need to collect/analyze to obtain, with probability \( 1-\beta \), a statistically significant result with confidence \( \alpha \).

Given the nature of microbiome data it is possible to quantify differences between groups at two levels: the alpha (within-sample) and beta (between samples) diversity (Figure 1). Alpha diversity metrics summarize the structure of a microbial community with respect to its richness (number of taxonomic groups), evenness (distribution of abundances of the
groups), or both [7]. Commonly used alpha metrics are Phylogenetic diversity [8], Observed number of amplicon sequence variants (ASV) [9], Chao1 [10], Simpson [11, 12] and Shannon indices [12, 13]. Beta diversity metrics summarize which samples differ from one another by considering sequence abundances or considering only presence-absence of sequences. Commonly used beta metrics are Bray-Curtis dissimilarity [14], Jaccard [15], unweighted UniFrac [16] and weighted UniFrac [17]. The choice of the diversity metrics affect the subsequent statistical testing and, as a result, how, and to which extent, power analysis can be performed.

Using an alpha diversity metric a single diversity value is obtained for each sample containing measurement over \( m \) taxa; thus the problem of assessing differences between two (or more) groups can be addressed with a univariate test, like \( t \)-test, ANOVA or a non-parametric version. The use of a beta diversity metric implies that all samples are to be considered simultaneously, and several methods to compare groups of samples measured on \( m > 1 \) have been proposed like ANOSIM [18] and PERMANOVA [19] to replace classical multivariate tests like the Hotelling \( T^2 \) or Multivariate ANOVA, which are in general not applicable because microbiome data does not satisfy basic assumptions. These assumptions include the independence of the sample units, the multivariate normality of errors, homogeneity of variance–covariance matrices among the groups or because the number of variables is larger than the number of samples making it impossible to apply the test [20] [6, 21-23].

While sample size, Type I and Type II error are well defined concepts, the definition of effect size depends on the outcome quantity of interest and how this quantity is mathematically defined. A fundamental step when performing power analysis is to define the effect size: for a simple two-sample \( t \)-test the effect size can be expressed as the Cohen’s \( \delta \) [24].

\[
\delta = \frac{|\mu_1 - \mu_2|}{\sigma}
\]

(1)

where \( \mu_1 \) and \( \mu_2 \) are the population means of the two groups and \( \sigma^2 \) is the pooled variance. Since \( \mu_1 \) and \( \mu_2 \) are population parameters which are inaccessible and on which we want to perform inference, an \textit{a priori} estimation, or educated guess, is necessary. This can be
accomplished by taking the sample means \( m_1 \) and \( m_2 \) and pooled variance \( s \) from a pilot study or existing data to obtain estimates of the population parameters. A critical aspect that is not sufficiently acknowledged is that the effect size from Equation (1) is sensitive to the particular diversity metric used to the point that sample size calculation can be severely affected.

The aim of this study is to show how, and to what extent, different diversity metrics influence the sample size needed to assess statistical significance of dissimilarities between different microbial communities. Both simulated and empirical 16S rRNA gene amplicon sequence data sets are used to perform retrospective power calculations on microbiota studies. A broad selection of alpha and beta diversity metrics were used to compare the different microbiota data sets. This study generated insight into the sensitivity and bias of certain statistical methods used in microbial ecology on microbiota research outcomes. We conclude with some recommendation for the reporting of power analysis and sample size calculations for microbiome studies.

**MATERIAL AND METHODS**

**Literature search**

To support the choice of alpha and beta diversity metrics to consider in our comparison, we performed a literature search on PubMed (www.pubmed.org) with query: (microbiota[Title] OR microbiome[Title]) NOT Review[Publication Type] 2020/01:2020/02 [Date of Publication]). This strategy aimed to include a broad scope of microbiota studies. We limited our search to studies published in English in with free full text.

**Alpha diversity metrics**

*Richness*

Richness is the number of taxa, most often defined as operational taxonomic unit (OTU) or amplicon sequence variant (ASV) observed [9]. Where \( s \) is the number of observed taxa, calculated as:

\[
S_{rich} = \sum s \cdot 1_{(p_s>0)}. \tag{2}
\]

*Phylogenetic Diversity*
Phylogenetic diversity (PD) is a phylogenetically-weighted measure of richness. Although, the name suggests diversity, it does not take into account the abundance of taxa. The PD is defined as the sum of the lengths of all those branches on the tree that span the members of the set, given the phylogenetic tree spanning \( s \) taxa [8]:

\[
PD = \sum_i b_i ,
\]  

(3)

where \( s \) is the number of observed taxa and \( b_i \) is the length of the \( i \)-th branch in the tree. The index \( i \) runs on all branches.

### Chao1

The Chao1 index is an abundance-based non-parametric estimator of taxa richness [10]. Its defined as

\[
Chao1 = s + \frac{F_1 (F_1 - 1)}{2(F_2 - 1)},
\]  

(4)

where \( s \) is the number of observed taxa, \( F_1 \) and \( F_2 \) are the number of OTU/ASV with only one sequence (i.e. “singletons”) and two sequences (i.e. “doubletons”). This metric assumes the number of organisms identified for a taxon to follow a Poisson distribution. Its definition rests on the concept that rare taxa bring most information about the number of missing taxa. This index gives more weight to the low abundances taxa and only the singletons and doubletons are used to estimate the number of missing taxa [10]. This index is particularly useful for data sets skewed towards low-abundance taxa [25, 26]. However, singletons and doubletons are often removed from 16S rRNA amplicon sequence data sets because the difficulty in robustly differentiating singleton errors from real singleton sequences [27, 28].

### Shannon index

The Shannon index \( H \) is an estimator of taxa diversity, combining richness and evenness) [12, 13]. It is defined as

\[
H = -\sum_{i=1}^{s} p_i \log(p_i),
\]  

(5)
where \( s \) is the number of OTU/ASV and \( p_i \) is the proportion of the community represented by the \( i \)-th OTU/ASV. Basically this index is the entropy associated to a given sample and quantifies the uncertainty in predicting the taxa identity of an individual selected at random from the sample. The Shannon’s index uses the relative abundances of different taxa, thus diversity depends on both taxa richness and evenness with which organisms are distributed among the different taxa. This index places a greater weight on taxa richness [26].

**Simpson index**

The Simpson index \( D \) is an estimator of taxa diversity, combining richness and evenness [11, 12]. It is defined as

\[
D = \frac{1}{\sum_{i=1}^{s} p_i^2},
\]

where \( s \) is the number of OTU/ASV and \( p_i \) is the proportion of the community represented by the \( i \)-th OTU/ASV. This index considers taxa evenness more than taxa richness in its measurement [26]; it indicates the taxa dominance and gives the probability of two individuals that belong to the same taxa being randomly chosen. It varies from 0 to 1 and the index increases as the diversity decreases [26].

**Metrics for beta diversity**

**Bray-Curtis dissimilarity**

The Bray-Curtis index (BC) [14] measures the compositional dissimilarity between the microbial communities of two samples \( i \) and \( j \) based on counts on each samples. It is defined as

\[
BC = 1 - \frac{2C_{ij}}{S_i + S_j},
\]

where \( C_{ij} \) is the sum of the smallest values for only those taxa in common between the sample \( i \) and \( j \), \( S_i \) and \( S_j \) are the total number of taxa counted in sample \( i \) and \( j \), respectively. This index ranges between 0 (the two samples share all taxa) and 1 (the two samples do not share any
taxa). It gives more weight to common taxa [29]. The Bray-Curtis dissimilarity is computed pairwise between all samples.

**Jaccard distance**

The Jaccard distance \( J \) between two samples \( i \) and \( j \) is defined as \( J = 1 - J(i,j) \) where \( J(i,j) \) is the Jaccard index which is defined as

\[
J(i, j) = \frac{|i \cap j|}{|i \cup j|},
\]

which is the ratio between the number of members that are common between the two samples and the number of members that are distinct; it is a measure of similarity for the two communities, and ranges between 0 (the communities are different) and 1 (the two communities are identical).

**UniFrac distances**

Unweighted UniFrac (UF) and weighted UniFrac distances between two samples take into account the phylogenetic tree, and thus phylogenetic distances between community members [17]. In unweighted UniFrac, the distance is calculated as the fraction of the branch length and in weighted UniFrac, branch lengths are weighted by the relative abundance of sequences. The sum of unshared branch lengths is divided by the sum of all tree branch lengths, which results in the fraction of total unshared branch lengths that is defined as

\[
\sum_i^n b_i \times \left[ \frac{A_i}{A_T} - \frac{B_i}{B_T} \right].
\]

Lozupone et al., [17] defined \( n \) as the total number of branches in the tree, \( b_i \) as the length of branch \( i \), \( A_i \) and \( B_i \) are the numbers of sequences that descend from branch \( i \) in communities \( A \) and \( B \), respectively, and \( A_T \) and \( B_T \) are the total numbers of sequences in communities \( A \) and \( B \), respectively. In order to control for unequal sampling effort, \( A_i \) and \( B_i \) are divided by \( A_T \) and \( B_T \) [17].
Experimental data sets

Chickdata data set: this data set contains 16S rRNA gene amplicon sequence data obtained from a broiler chicken experiment. The data set is described in detail in (Kers et al., 2019[30]); briefly, chickens were raised under three different housing conditions with the same medium chain fatty acid feed intervention. Between those housing conditions, bird management was kept as similar as possible. At the hatchery the chicks were randomly allocated to three different experimental facilities. Data set A contains 70 broilers from a grow-out feed trial facility, data set B contains 70 broilers raised at a floor stable and data set C contains 70 broilers raised in isolators. A feed intervention was used as a tool to create differences in cecal microbiota between broilers within the same housing condition.

HMP data set: this data set was obtained from the Human Microbiome Project (HMP) phase I [31]. It contains 16S rRNA gene amplicon sequence data of 169 stool samples, 150 oral samples, 86 vaginal samples and 69 skin samples. The bodyside microbiomes are all diverse in terms of community membership [31].

In all data sets ASVs were defined as unique sequences. All data was analysed using NG-Tax [32]. Taxonomy was assigned using the SILVA 128 16S rRNA gene reference [33]. An overview of data set characteristics (sample size, the number of ASVs, mean values of different alpha- and beta diversity metrics.) is shown in Table 1.

Simulated data set

We simulated two different scenarios described by Simulated data set 1 and 2.

Simulated data set 1, is built starting from 1995 microbial features observed in 169 stool samples from the Human Microbiome Project (HMP), indicated as $X_1$ in the following. Data sets (name $X_i$ for sake of simplicity) were created where 2%, 5%, 10%, 25%, 50% and 75% of bacterial features were randomly removed.

Simulated data set 2, is a case-control scenario ($X_1$: controls, $X_2$: cases, 1995 features and 169 samples each) where 1%, 2%, 5%, 10%, 15% and 20% of bacterial features were differentially abundant. Both simulated data sets have the same phylogenetic structure as the HMP data set. An overview of the characteristics of the simulated data sets is shown in Table 2.
**Statistical tests for group differences**

*Univariate statistical analysis*

Differences between groups using alpha diversity as determined by using Phylogenetic diversity, Richness (defined as Observed), Chao1, Simpson and Shannon, were assessed using the a Kruskal-Wallis test (setting $K = 1000$)[34]. A significance threshold $\alpha = 0.01$ was used in all calculations.

*PERMANOVA*

Differences between groups using beta diversity as determined by using Bray-Curtis, Jaccard, unweighted UniFrac and weighted UniFrac were assessed using the Permutational Multivariate Analysis of Variance (PERMANOVA) [19]. PERMANOVA is a robust approach to compare groups of samples measured on $m > 1$ variables. It construct ANOVA-like test statistics from a matrix of resemblances (distances, dissimilarities, or similarities) calculated among the sample units, and assesses significance of the observed differences using random permutations of observations among the groups [35]. The Null-hypothesis $H_0$ tested by PERMANOVA is that the centroids of the groups (in the space of the chosen resemblance measure), are the same for all groups. This test assumes that samples are exchangeable under the Null hypothesis, are independent and have similar multivariate dispersion. The PERMANOVA test statistic is a pseudo ANOVA $F$-ratio:

$$F = \frac{SS_B(g-1)}{SS_W(n-g)}$$

(10)

where $SS_B$ is the total sum of squares of the (diss)similarities between groups, $SS_W$ is the total sum of squares of the (diss)similarities within groups, $g$ is the number of groups and $n$ is the total number of samples.

The significance of the $F$ statistics is calculated by means of permutations ($k=9999$). The distribution of $F$ under the Null hypothesis is generated by permuting $g$ times the sample group labels and recalculating $F$ on the permuted data. Significance is expressed as $p$-value calculated as the fraction of permuted $F$-statistics, which are equal to or greater than the pseudo $F$-ratio observed on the original data.
Calculation of the empirical power

From \( N_1 \times m \) and \( N_2 \times m \) data matrices \( \mathbf{X}_1 \) and \( \mathbf{X}_2 \) we random sampled with replacements \( K \) \( n_1 \times m \) and \( n_2 \times m \) data sets and applied a statistical test to assess the difference between \( \mathbf{X}_1 \) and \( \mathbf{X}_2 \) (as quantified by any of the alpha and beta metrics) at significance level \( \alpha = 0.01 \) under the assumption of the Null hypothesis being false. The empirical power of the test is defined as the empirical probability \( EPr \) of \( H_0 \) being rejected calculated as

\[
EPr = \frac{\#(H_0 \text{ rejected}|H_0 \text{ false})}{K} \tag{11}
\]

where \( \#() \) indicate the number of times that \( H_0 \) is rejected. For sake of simplicity we consider \( n_1 = n_2 = n \) and we varied \( n \) between \( X \) and \( Y \).

Software

All statistical analyses were performed in R version 4.0.2 (R Foundation for Statistical Computing, Austria [36]), using the packages: Phyloseq, Microbiome, and Vegan [37-39]. PERMANOVA was performed using the \textit{adonis} function for the Vegan package. Other power calculations were performed using the G*power software [40] using the “Means: Difference between two independent means (two groups)” as Statistical test and “a priori” and “post hoc” option for the Type of Power analysis. Differentially abundant microbiota profiles were simulated with the microbiomeDASim R package (Williams et al., 2019[41]) using the \textit{gen_norm_microbiome} function. The R scripts can be found on the Github page: https://github.com/mibwurrepo/KersSaccenti-Power.

Data availability


RESULTS
**Motivation example**

We begin with a motivational example to show how the choice of the diversity metrics affects the power of a microbiome study, and how the same study may be underpowered if a different metric is used.

Let’s suppose we want to plan an experiment to assess whether gut and oral microbial communities are different. A very simple and basic study design would be to collect $n_1 = n_2$ gut and oral samples and compare the alpha diversity between the two conditions (gut vs oral) using a two-sample Kruskal-Wallis $t$-test.

We can base our estimation $d$ of a very similar effect size $\delta$ on data from HMP (Table 1). Using four different alpha metrics and Equation (1) we obtained $d = 1.27$ (Phylogenetic diversity), $d = 0.3621$ (Shannon), $d = 0.58$ (Chao1), $d = 0$ (Simpson). These values are markedly different: fixing the power to 0.8 ($\beta = 0.2$) and confidence $\alpha = 0.05$, they will lead to dramatically different required total sample size (Figure 2A). This clearly indicates that microbiome studies may be severely underpowered depending on which alpha metric was used to compare two (or more) groups.

We also explored the achievable power by fixing the sample size ($n = n_1 + n_2 = 50 + 50 = 100$) and using different effect sizes (Figure 2B). Consistently with what observed in Figure 2A results vary strongly, providing a clear indication of the risk of underpowering when Shannon diversity is used.

Note that with the use of beta diversity metrics, performing a priori power analysis becomes much more complicated. The classical tools for power analysis cannot be applied since the statistical tools are not parametric: solutions have been proposed in the literature: see, for instance [42-44].

**Literature search**

Our literature search returned 632 papers matching the search criteria. We selected randomly 100 papers, and of those the material and methods or full text was investigated to obtain an overview of the most frequently used alpha and beta diversity metrics and of the sample sizes used. Of the 100 full text papers, 92(%) papers contained alpha metrics and 83(%) papers contained beta metrics.
In 58% of the papers more than one alpha metric was used. In 21% of the papers more than one beta metric was used. An overview of the frequency of the different metrics showed that the Shannon index and Bray-Curtis dissimilarity are the most common metrics (Table 3). There was a wide variance in the used sample size, defined as the smallest number per group: 46(%) papers had a sample size between ≤10 samples, 34(%) papers used between 11-50 samples, 7(%) papers between 51-100 samples, 10(%) papers between 101-1000 samples, and three papers more than >1000 samples.

The power of microbiome studies
As shown in the Motivational example, the choice of a particular alpha (and beta) diversity metric determines the number of samples required to achieve a pre-determined power. Based on this observation we examined two simulated data sets using both alpha and beta diversity metrics to understand the relationship between the sample size, the observed power and the diversity metrics, together with two experimental data sets (Chicken and HMP data sets).

As representatives of testing procedures using alpha and beta diversity measures to compare two groups we selected the Kruskal-Wallis test (for alpha metrics) and PERMANOVA (for beta metrics), respectively. The Kruskal-Wallis test is the non-parametric choice for comparing two groups when the normality assumption does not hold. When comparing two (or more) groups using beta diversity metrics PERMANOVA and ANOSIM (Analysis of Similarity, [18] are popular choices. The two approaches are equally popular (359 hits on Pubmed for PERMANOVA and 341 for ANOSIM), however, Anderson and Walsh [35] showed that while both approaches are sensitive to unbalanced designs and differences in variance within groups, PERMANOVA is a more robust approach: on this ground we based our choice for PERMANOVA.

Power analysis of simulated data sets
For the simulated data sets, the effect size is known a priori and it is expressed as the % of differentially abundant or present/absent microbial features (ASV) (Figure 3). The achievable power for Simulated data set 1 is shown as a function of the sample size (n) for different percentages of present/absent ASV. If 2%, 5% of the ASV are deleted from the data set, none of the alpha diversity metrics was able to capture the difference between data sets X1 and X2, irrespective of the sample size used (Figure 3A-B). When more than 10% of ASV were removed in data set X2 (Figure 3C-E) all measures were somehow able to capture the difference but the
resulting actual power was very different. Overall, Chao1 and observed diversity allowed higher power with the lower sample size (are more sensitive to observe differences), especially in the medium range of differences (10% to 25%, Figure 3C-D), whereas differences are minimal for >25%. Note that in contrast with the Motivation example, here the Phylogenetic diversity was not the metric resulting in the smallest sample size.

The same approach was used across different beta diversity metrics (Figure 4). The Jaccard diversity metric was most sensitive and weighted UniFrac was least sensitive to observe the differences in presence/absence between the data sets (Figure 4B-F). When more than 10% of ASV were removed in data set X2, no difference between data sets was observed by the UniFrac metric (Figure 4C), while with 25% removed, Bray-Curtis and unweighted UniFrac showed a comparable power and sample size (Figure 4D). In this simulated data set, Weighted UniFrac distance needed the highest sample size to observe the difference (Figure 4D-F).

The achievable power for Simulated data set 2 is also shown as function of the sample size for different percentages of differentially abundant ASV (Figure 5). If ≤ 5% of the ASV were differentially abundant in data set X2 as compared to X1, the Simpson metric needed the lowest sample size (is most sensitive) to observe differences between the data (Figure 5A-C). However, if 10% of the ASV were differentially abundant, the Phylogenetic diversity and Chao1 were more sensitive and the Simpson and Shannon metrics less sensitive (Figure 5D). With 15% of the ASV differentially abundant no differences were observed with the Simpson metrics (Figure 5E).

The same approach was used across different beta diversity metrics (Figure 6). Bray-Curtis distance was most sensitive to observe differences, whereas unweighted UniFrac needed the largest sample size. If 2% of the ASV were differentially abundant power of the beta metrics was totally different, for example, a sample size of 15 would result in power: 100 for Bray-Curtis, 50 for Weighted UniFrac, 40 for Jaccard distance, and just 10 with unweighted UniFrac (Figure 6B). However, if 10% of the ASV were differentially abundant, all metrics would result in a power higher than 0.80 (Figure 6D-F).

**Power analysis of experimental data sets: Chicken data set**

The Shannon index was the most sensitive alpha metric and Chao1 and Phylogenetic diversity were less sensitive metrics to observe a difference between the groups in data set B (Figure 7A). In data set B, Shannon index was also the most sensitive alpha metric but Simpson was
the least sensitive metric (Figure 7B). Unweighted UniFrac distance was the most sensitive beta
diversity metric to observe a difference between groups in data set A (Figure 8A). Jaccard
distance was the only metric that showed that H3 needed the smallest sample size, indicating
that in H3 specific ASV are differentially present between the groups (Figure 8B). Weighted
UniFrac was more sensitive compared to unweighted UniFrac to observe a difference between
the groups based on their microbial communities (Figure 8C, D). In general, the alpha diversity
measures were less sensitive to observe differences between the broilers compared to the
beta diversity (Figure 7A-B, Figure 8).

Although no difference in alpha diversity was observed between broilers fed with or
without MCFA raised in housing condition 1, the average daily gain and the average daily feed
intake were lower in MCFA broilers [30]. Therefore, the difference only observed based on the
beta diversity might already be biologically relevant and hence sufficient to draw conclusions
in this case. Based on this data set we observed that Shannon is the most sensitive alpha
diversity metric to observe differences between groups, resulting in the lowest needed sample
size. The sensitivity of the beta diversity, however, was different per data set. Based on this
retrospective power calculation two conclusions can be drawn on this study design. First, not
enough chickens were sampled to observe a difference in the alpha diversity between broilers
fed with or without MCFA raised in data set A. Second, 15 chicken samples instead of 35
samples per group would have resulted in the same conclusion.

Power analysis of experimental data: Human Microbiome Project data set
The samples in this data set were collected from different body sites and are known to have a
very distinct origin, and therefore expected to be different in microbial composition. The
comparison between different body sites showed wide variation in sample size across different
alpha diversities (Figure 7C-F). The difference in sample size was small in the comparison
between skin vs oral microbiome samples, all-around ten samples (threshold power 80, 1-
β)(Figure 7C). In the skin vs gut microbiome samples, the Simpson and Shannon alpha
diversities did not differ, and the phylogenetic diversity was the most sensitive to observe
differences (Figure 7D). In contrast, when comparing the gut vs the oral microbiome, the
phylogenetic diversity was least sensitive to observe differences, whereas the Shannon and
Simpson metrics were different between gut and oral samples (Figure 7E). In the skin vs vaginal
microbiome comparison, Simpson and Shannon's alpha diversities were more sensitive
compared to the Observed/Chao1 and phylogenetic diversity (Figure 7F). Based on the different beta diversity metrics, all comparisons between different body sites supported significant differences even when just five samples were compared (data not shown), due to the large difference between communities (Supplementary Figure 1). Therefore the retrospective power calculations were not informative for this data set.

Are microbiome studies underpowered?

Figure 9A shows the distribution of the sample size of the data sets that were analysed, using the Chao1 diversity measure (among others) in 28 of the 100 papers considered in the literature review. The distribution is highly skewed towards 0 with a median of 39 samples per group and a mode of 8 samples. Removing the two outlying studies with >300 samples, resulted in a median of 23 samples.

Even considering that Chao1 was one the best performing measures in both simulated and experimental data sets, these numbers appear to be worryingly low: on experimental data, which have a complicated structure that is impossible to replicate in simulations, it is rarely possible to attain a power of 80% with less than 40 samples per group. Similar considerations hold when PERMANOVA is applied (see Figure 9B), with a median group size of 22 and mode 3.

Reporting of power analysis

One of the studies we examined in the literary review reported that sample size and power analysis were performed: “Sample sizes were chosen on the basis of pilot experiments and on our experience with similar experiments.” This is commendable but we believe that the way forward is to employ and report in full a standardized summary of sample size calculations performed. The software G*power generates a Protocol for power analysis. For instance for a two group- comparison with a Mann-Whitney/Kruskal-Wallis test, one should report, once adapted

\[
\begin{array}{|c|c|}
\hline
\textbf{t tests} & \text{Means: Wilcoxon-Mann-Whitney test (two groups)} \\
\textbf{Options:} & \text{A.R.E. method} \\
\textbf{Analysis:} & \text{A priori: Compute required sample size} \\
\textbf{Input:} & \\
\text{Tail(s)} & = \text{One} \\
\text{Parent distribution} & = \text{Normal} \\
\text{Effect size d} & = 0.5 \\
\text{Alpha metric} & = \text{Shannon} \\
\alpha \text{ err prob} & = 0.05 \\
\hline
\end{array}
\]
Power (1−β err prob) = 0.8
Allocation ratio N2/N1 = 1

Output:
Noncentrality parameter δ = 2.51
Critical t = 1.66
Df = 99.2
Sample size group 1 = 53
Sample size group 2 = 53
Total sample size = 106
Actual power = 0.803

Together with this, information should be provided as to how effect size was determined, i.e. which pilot data were used and how the effect size was determined.

A similar reporting protocol could be devised if simulations are used in a PERMANOVA setting. Since simulation and/or pilot data must be used in this case, details on the simulations or pilot data should be reported. For instance, using the Chicken data 1 as pilot, one could report the following protocol, taking 100 re-samplings of size 6 to calculate the achievable power:

**Test – PERMANOVA**

**Options:**
9999 permutation
100 iterations

**Analysis:**
Compute achievable power

**Input:**
- Beta metics = Bary-Curtis
- α err prob = 0.05
- Number of groups = 2
- Number of taxa = 363
- Sample size group 1 = 6
- Sample size group 2 = 6

**Output:**
- Observed Effect size (average) σ² = 0.120682
- Min\Max effect size = 0.025922\0.3500687
- Observed Effect size (average) f = 0.2696886
- Min\Max effect size = 0.1319342\0.746349
- Numerator df = 1
- Denominator df = 10
- Power (1−β err prob) = 0.97

**DISCUSSION**
The aim of this study was to assess how, and to what extent, different diversity metrics and compositions of the microbiota influence the needed sample size to observed dissimilar groups. Based on our literature survey we observed that the Shannon and Bray-Curtis metrics are most published metrics. This might because they are often the most sensitive metrics to
observe differences between groups, resulting in a lower sample size. Our results are in line
with previous literature that showed that the choice of distance metric may significantly
influence the observed results [45].

A well-known phenomenon that can hamper progress in every research field concerns
publication biases in reporting mainly positive findings [46]. In microbiota research this might
even occur rather unintentionally, by using certain alpha and beta diversity metrics, but it
might also be that researchers selectively report only results for the metric that shows
significance even when other metrics had been assessed during the analyses. Our results lead
to the speculation that many microbiome studies may be underpowered or, conversely, only
reporting evidence of very large effects that can be assessed to be statistically significant also
with small sample size. However, since effect size and test statistic are not reported, it is
impossible to judge on the quality of the results. Not only, this also hampers the use of
published studies as pilot studies to perform power analysis and sample size calculations since,
as long as data is not de novo re-analysed.

None of the 100 microbiome studies that we have considered reported the effect size. A
collaborative project aiming to investigate reproducibility of 100 high-profile psychological
studies reported the average effect size observed in the replication studies was approximately
half the magnitude of those given in the original studies, leading to a replication success of only
36% [47]. The lack of reported effects makes it impossible to analyse retrospectively
microbiome studies, to perform meta-analysis and, more importantly, makes it impossible to
check the consistency of the statistical analysis or detect errors.

On the basis of this, reporting of effects and test statistics should be made compulsory
in microbiome studies. For the highly used Kruskal-Wallis test, the $H$ test statistic is given by:

$$H = \frac{12}{N^2 - 1} \sum_i \frac{R_i^2}{n_i} - 3N + 1$$  \hspace{1cm} (13)

where $N$ is the total number of samples, $n_i$ is the number of samples in group $i$ and $R_i$ is the
total sum of ranks in the $i$-th group. Note that $H$ is easily obtainable from most software
packages.

For Kruskal-Wallis the most common effect is the $\eta^2$ which is defined as:
\[ \eta^2 = \frac{H - k + 1}{N - k} \]  

(14)

where \( \eta \) is the value obtained in the Kruskal-Wallis test, \( k \) is the number of groups. For instance, for the comparison of the two feedings (feed A and B, see Table 1) from the chicken data set using the Observed alpha diversity one could report:

Feed A \((n_1 = 35)\) and Feed B \((n_2 = 35)\) samples were compared with Kruskal-Wallis test using the Chao1 metric: \( H(df) = H(1) = 14.68, P\text{-value } 0.0001, \eta^2 = 0.66, \delta = 0.58. \)

where \( df \) indicates the degrees of freedom. Note here that for a two group comparison, the Kruskal-Wallis test is equivalent to the Wilcoxon-Mann-Whitney (WMW) test [48, 49]. For the WMW test the Cohen’s \( \delta \) effect size definition (Equation (1)) also applies [50]. This greatly simplifies power analysis and sample size calculations: we advise to also report \( \delta \) when two groups are considered.

In addition, performing power analysis for a Kruskal-Wallis is not a simple matter and requires the use of a rather advanced statistical machinery [51, 52]; for instance, such calculations are not included in G*power [40], which is the most complete software for power analysis. Kruskal-Wallis is the non-parametric counterpart of one-way ANOVA and as such is used in situations where there are more than two groups. However, whereas power analysis and sample size calculation for a one-way ANOVA with more than two groups are “easily” accessible within R or other software packages, this is not the case Kruskal-Wallis testing. We could locate a R package ‘MultNonParam’ [53] that performs power analysis for the Kruskal-Wallis test with more than two groups, however, it requires the specification of the offsets for the various populations, under the alternative hypothesis. Relating such tools to determine the effect size observed in microbiome data is a matter we believe to be worthy of exploration and brings us back to the problem that statistics and effect size are not easily available for microbiome studies.

The principle of reporting the effect size should also apply when testing is performed using beta diversity metrics, in which case the PERMANOVA pseudo F-statistics (see Equation
(10)) and the effect size should be reported. Typical effect measures in ANOVA are the Choen’s $f^2$ and $\eta^2$ and the $\omega^2$

Feed A ($n_1=35$) and Feed B ($n_2=35$) samples were compared with PERMANOVA test using the Unweighted UniFrac metric: $F(df_B, df_W) = F(1,68) = 6.27$, P-value = 0.0001, $f^2 = 0.092$, 1000 permutations.

where $df_B$, $df_W$ indicate the between-groups and within degrees of freedom. These notations follows guidelines of the American Psychological Association which provides standardized formats for the reporting of statistical analysis for statistical procedures [54]. For PERMANOVA the matter complicates considerably: to estimate statistical power and calculate sample size, one must quantify the expected within-group variance and the effect to be expected when comparing two or more groups. A package like micropower [43] in principle allows estimation of PERMANOVA effects quantified by the $\omega^2$ value (limited to the UniFrac measure): unfortunately, at the best of our knowledge, the package seems not to be maintained and lacks a proper manual. The original paper presents a table with effects calculated from different studies that could be used as guide, however this metric is not standard. It can be calculated from the PERMANOVA table as

$$\omega^2 = \frac{SS_{effeect} - df_{effect}MS_{residual}}{SS_{effect} + MS_{residual}}$$

For the chicken data we observed $\omega^2$ values in the range 0.04 – 0.1, depending on the beta metrics, and these are consistent with those reported in Table 1 from [43].

A more common effect measure is the Cohen’s $f^2$-value, i.e. the between group to within group ratio can be easily obtained by the ANOVA table provided by software like the R package Vegan by taking the ratio between the Treatment sum of squares and the Residual sum of squares. The $f^2$-value is used for power calculation in the ANOVA setting, however, it should not be used to perform sample size calculation for PERMANOVA, not even to obtain a rough indication, since the corresponding F-statistics do not follow an F distribution. For instance, when comparing Feed A and B from the first chicken data with PERMANOVA we can derive a Cohen’s $f^2 = 0.38$; if this value is used to perform power calculation for an ANOVA with
two groups with power 80% at $\alpha = 0.01$ we obtain that 42 sample per group are need. However, comparing with the results in Figure 8 we see that a 100% power can be obtained with 25 samples per group, no matter which measure is used: this is a clear indication that power analysis for PERMANOVA can be obtained only by mean of simulations. In this light Figure 8 can be viewed as a priori power calculations using pilot data.

Furthermore, there is a convention, more or less widely accepted, of classifying Cohen’s $\delta$ effect which we have used in the power calculation for the Kruskal-Wallis into trivial ($\delta < 0.2$) small ($\delta = 0.2$), medium ($\delta = 0.5$) and large ($\delta > 0.8$). However, this classification is based on what is observed in psychology and does not apply automatically to other fields of research [55]. In microbiome studies the effects may be in same order of magnitude that may be considered large or very large using the standard convention.

In sum, we have seen that different alpha and beta diversity metrics lead to different study power: on the basis of this observation, one could be naturally tempted to try all possible metrics until one or more are found that give a statistically significant test result, i.e. $p$-value < $\alpha$. This way of proceeding is one of the many forms of the so-called $p$-value hacking ($p$-hacking) [56]. $P$-hacking (also called data dredging, significance chasing, significance questing, or selective inference [57]) is the improper use of data (like adding or removing observations) or statistical procedures (like applying many different tests) until a configuration is found that produces a statistically significant result at the desired confidence level [58]. $P$-hacking is an illegitimate practice that promotes unreproducible results, polluting literature and adding to publication bias [59-61].

To this end, in our opinion, the only way to protect ourselves from (the temptation of) $p$-hacking would be to publish, and we stress here the word publish, a statistical plan before experiments are initiated: this practice is customary for clinical trials where a statistical plan describing the endpoints and the corresponding statistical analyses must be disclosed before the start of the study and must be adhered to if results are going to be published [62]. This is the only guarantee that data analysis is not manipulated towards artificially inflated significant results. We appreciate that clinical trials are inherently different from microbiome (and other omics) studies which are often exploratory in nature, but as far as statistics is concerned, they are prey of the same traps and pitfalls. It is obvious that such a change in the approach to
microbiome studies requires the concerted cooperation of researchers, journal editors, reviewers and publishers.
FUNDING
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CONFLICT OF INTEREST
The authors declare that the research was conducted in the absence of any commercial or financial relationship that could be construed as a potential conflict of interest.
Differences between Group 1 and Group 2 in microbiome data can be assessed using either alpha (within sample diversity) or beta (between sample diversity) metrics. The use of alpha metrics allows the use of classical univariate testing, either parametric or non-parametric. The use of beta metrics leads to the use of permutation-based testing approaches like PERMANOVA (see Material and Methods).
Figure 2

Total sample size \((n)\) required to assess the statistical significance of an effect \(d\) using a two-sample Kruskal-Wallis test with a power equal to 0.8 and confidence \(\alpha = 0.01.\) B) Achievable power attainable by a Kruskal-Wallis test using a total sample size \(n = n_1 + n_2 = 50 + 50 = 100\) using different alpha metrics. Note that for a null effect \((d=0)\) the achievable power coincides with \(\alpha.\)
Figure 3

Empirical power for the statistical comparison of two groups $X_1$ and $X_2$ (Simulated data set 1). The Empirical power is calculated using Equation (11) as a function of the sample size $n_1$ of group 1 (with $n_2 = n_1$ and total sample size $n = n_1 + n_2$). Alpha diversity metrics and Kruskal-Wallis test are used to test for differences.
Empirical power for the statistical comparison of two groups $X_1$ and $X_2$ (Simulated data set 1). The Empirical power is calculated using Equation (11) as a function of the sample size $n_1$ of group 1 (with $n_2 = n_1$ and total sample size $n = n_1 + n_2$). Beta diversity metrics and PERMANOVA test are used to test for differences ($B = \text{Bray-Curtis}, J = \text{Jaccard}, UF= \text{unweighted UniFrac}, WUF = \text{Weighted UniFrac}$).
Figure 5

Empirical power for the statistical comparison of two groups $X_1$ and $X_2$ (Simulated data set 2). The Empirical power is calculated using Equation (11) as a function of the sample size $n_1$ of group 1 (with $n_2 = n_1$ and total sample size $n = n_1 + n_2$). Alpha diversity metrics and Kruskal-Wallis test are used to test for differences.
Empirical power for the statistical comparison of two groups $X_1$ and $X_2$ (Simulated data set 2). The Empirical power is calculated using Equation (11) as a function of the sample size $n_1$ of group 1 (with $n_2 = n_1$ and total sample size $n = n_1 + n_2$). Beta diversity metrics and PERMANOVA test are used to test for differences.

**Figure 6**

Empirical power for the statistical comparison of two groups $X_1$ and $X_2$ (Simulated data set 2). The Empirical power is calculated using Equation (11) as a function of the sample size $n_1$ of group 1 (with $n_2 = n_1$ and total sample size $n = n_1 + n_2$). Beta diversity metrics and PERMANOVA test are used to test for differences.
Figure 7

Empirical power for the statistical comparison of the skin, gut and oral microbiome data sets from the Human Microbiome Projects. The Empirical power is calculated using Equation (11). Differences between any two data sets $X_1$ and $X_2$ is assessed using alpha diversity metrics using a Kruskal-Wallis test. Empirical power is calculated as a function of the sample size $n_1$ of group 1 (with $n_2 = n_3$ and total sample size $n = n_1 + n_2$) over $k=2000$ tests per each sample size.
Figure 8

Different beta diversity metrics to estimate different sample sizes (n=100 repetitions). A. Chicken data set, B. data set B, C data set C.
The distribution of the sample size of the data sets that were analysed, using the Chao1 diversity measure (among others) in 28 of the 100 papers considered in the literature review.
Overview of data set characteristics of the different data sets. The mean alpha and beta diversity, the standard deviation between brackets (A,B). The mean of the beta diversities is also calculated as the mean distance of groups members to the group centroid (C). \( n = \) sample size. Feed 1, is the intervention without same medium chain fatty acid. PD is the phylogenetic diversity.

### A

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### C

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Table 2
Overview of data set characteristics of the simulated data sets. The mean alpha and beta diversity, the standard deviation between brackets (A,B). n = sample size, PD is the phylogenetic diversity, BC is Bray-Curtis dissimilarity.

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The frequency of the different alpha and beta metrics in published papers with microbiome or microbiota in the title and published between 01/2020 and 02/2020 (n=100, multiple metrics per paper were often used)

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Supplementary figures

Figure S1
Principal coordinate plots (PCoA) based on (a) Bray-Curtis and (b) Jaccard and (c) unweighted UniFrac and (d) Weighted UniFrac distances between the different human sample types. Different colors indicate different sample types.
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