Rarefy your data

Running title: Rarefy your data
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10 Abstract

Importance

Introduction

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- Motivation
 - Problem of uneven sampling effort
- What is rarefaction? History, reason for rarefaction
 - Repeated down sampling of datasets to a common number of observations to calculate the average value to ascertain the expected value of a metric for the metric under study; typically richness
 - Control for unneven sampling effort
 - Methods vary in their sensitivity to uneven sampling
 - Compositional data analysis
- · Reasons behind "rarefaction is inadmissable"
 - Weird simulation
- · Alternative approaches and claims
 - sampling invariance
- · Goal of this study

This analysis included 16S rRNA gene sequence data from from 12 studies that characterized the variation in bacterial communities from diverse environments (Table 1). The original studies generated the sequence data by pooling separate PCR products that were generated by amplifying the V4 region of the 16S rRNA gene from the bacterial DNA in multiple samples. Because pooling equimolar quantities of DNA is frought with difficulties, it was common to observe wide variation in the number of sequences in each sample (Table 1 and Figure S1).

Results

Without rarefaction, metrics of alpha diversity are sensitive to sampling effort. To test the sesitivity of various approaches of measuring alpha diversity to sampling effort, I generated null models for each study. Under a null model, each community from the same study would be expected to have the same alpha diversity regardless of the sampling effort. I measured the richness of the communities in each study without any correction, using scaled ranked subsampling (SRS) normalized OTU counts, with estimates based on

non-parametric and parametric approaches, and using rarefaction (e.g. Figure S2). For each study, all of
the approaches, except for rarefaction, showed a strong correlation between richness and the number of
sequences in the sample (Figure 1A). Next, I assessed diversity using the Shannon diversity index and
the inverse Simpson diversity index without any correction, using normalized OTU counts, and rarefaction; I
also used a non-parametric estimator of Shannon diversity. The correlation between sampling depth and the
diversity metric was not as strong as it was for richness and the inverse Simpson diversity values were less
sensitive than the Shannon diversity values; however, the correlation to the rarefied diversity metrics were
the lowest for all of the metrics and studies (Figure 1A). The rarefied alpha-diversity metrics consistently
demonstrated a lack of sensitivity to sampling depth.

Without rarefaction, metrics of beta diversity are sensitive to sampling effort. To test the sesitivity of various approaches of measuring beta diversity to sampling effort, I used the same null models used for studying the sensitivity of alpha diversity. Under a null model, the ecological distance between any pair of samples would be the same regardless of the difference in the number of sequences observed in each sample (e.g., Figure S3). First, I calculated the Jaccard distance coefficient between all pairs of communities within a study. The Jaccard distance coefficient is the fraction of OTUs that are unique to either community and does not account for the abundance of the OTUs. Jaccard distances were calcualted using the uncorrected OTU counts, with rarefaction, relative abundances, and following normalization using cumulative sum scaling (CSS) and SRS. Only the rarefied distances showed a lack of sensitivity to sampling effort (Figure 1B). Second, I analyzed the sensitivity of the Bray-Curtis distance coefficient, which is a popular metric that incorporates the abundance of each OTU. Similar to what I observed with the Jaccard coefficient, only the rarefied data showed a lack of sensitivity to sampling effort (Figure 1B). Third, I calcualted the Euclidean distance on raw OTU counts where the central log-ratio (CLR) was calculated (i.e., Aitchison distances) by ignoring OTUs in samples with zero counts (Robust CLR), adding a pseudocount of one to all OTU counts prior to calculating the CLR (One CLR), adding a pseudocount of one divided by the total number of sequences obtained for the community (Nudge CLR), and imputing the value of zero counts (Zero CLR). The Aitchison distances were all strongly sensitive to sampling effort (Figure 1B). Finally, I used the variance stabilization technique (VST) from DeSeq2 prior to calculating Euclidean distances. Again, there was a strong sensitivity to sampling effort (Figure 1B). Although Euclidean distances are not typically used on raw or rarefied count data in ecology, rarefied Euclidean distances were not sensitive to sampling effort. Across each of the beta diversity metrics and approaches used to account for uneven sampling effort and sparsity, rarefaction was the least sensitive approach to differences in sampling effort.

Rarefaction limits the detection of false positives when sampling effort and treatment group are con-

founded. Next, I investigated the impact of the various strategies and metrics on falsely detecting a significant difference using the the same communities generated from the null model in the analysis of alpha and beta diversity metrics. To test for differences in alpha and beta diversity I used the non-parametric Wilcoxon test and non-parametric permutation-based multivariate analysis of variance (PERMANOVA). First, within each study, I randomly assigned each sample to one of two treatment groups. My expectation was that approximately 5 of the 100 (5%) random tests for each comparison would yield a significant test result. Indeed, for each study and alpha and beta diversity metric and strategy for accounting for uneven sampling, approximately 5% of the tests yielded a significant result (Figure 2). Second, within each study, I assigned samples with more than the median number of sequences per sample to one treatment group and the rest to another treatment group. If there was no sensitivity to sampling effort, I would have expected that 5% of the tests would yield a significant result. In fact, only the rarefied data consistently resulted in a 5% false positive rate for alpha and beta diversity metrics (Figure 2). These results align with the observed sensitivity of alpha and beta diversity metrics to sampling effort and underscore the value of rarefaction.

Rarefaction preserves the statistical power to detect differences between treatment groups. To assess the impact of different approaches to control for uneven sampling effort I performed two additional simulations. In the first simulation, for each study communities were randomly assigned to one of two treatment groups. Communities in the first treatment group were generated by sampling from the null distribution. Samples in the second treatment group were generated by perturbing the null distribution by increasing the relative abundance of 10% of the OTUs by 5%. These values were determined after empirically searching for conditions that resulted in a large fraction of the randomizations yieleding a significant result across most of the studies. The fraction of tests that yielded a significant test was a measure of the statistical power for the test. The power to detect differences in richness by all approaches was low (Figure 4A). This was likely because the approach for generating the perturbed community did not necessarily change the number of OTUs in each treatment group. Regardless, the simulations testing differencse in richness using the Rice and Stream datasets had the greatest power when the richness data were rarefied. To explore this further, in a second simulation the second treatment group was perturbed by removing 3% of the OTUs from the model. As suggested by the first simulation, the rarefied richness data had a higher statistical power than the other approaches when measuring richness (Figure 5). The simulations testing the power to detect differences in Shannon diversity also showed that rarefied data performed other methods (Figure 4A). When testing for differences in the Inverse Simpson diversity index the the difference between rarefaction and the other methods was negligible (Figure 4A). For tests of beta diversity I found that rarefaction was the most reliable approach to maintain statistical power to detect differences between

two communities. Among the tests using the Jaccard and Bray-Curtis metrics, raw count data and CSS normalized data had little power relative to rarefied, relative abundance, and SRS normalized data. The differences in power between rarefied, relative abundance, and SRS normalized data was small, but if there were differences, the power obtained using rarefied data was greater than the other methods. Among the tests using Euclidean distances, using raw counts and CLR and DeSeq2 transformed data had little power relative to the distances calcualted using rarefied and relative abundance data. This power-based analysis of the simulated communities using different methods of handling uneven sample sizes demonstrated the value of rarefaction for preserving the statistical power to detect differences between treatment groups for measures of alpha and beta diversity.

Increased rarefaction depth reduces intra-sample variation in alpha and beta diversity. Once concern with rarefying communities is the perceived loss of sequencing information when more a large fraction of data appears to be removed when the community with the greatest sequencing depth is rarefied to the size of the community with the least (e.g., 99% with the Bioethanol dataset). To assess the sensitivity of alpha and beta diversity metrics to rarefaction depth, I again used the dataset generated using the null models, but rarefied each community to varying sampling depths (Figure 6). The richness values increased with sampling effort as rare OTUs would continue be detected. In contrast, the Shannon diversity and Bray-Curtis values plateaued with increased sampling effort. This result was expected since increased sampling would lead to increased precision in the measured abundance of OTUs. Next, I measured the coefficient of variation (i.e., the mean divided by the standard deviation) between samples for richness, Shannon diversity, and Bray-Curtis distances. Although the richness values appeared to increased unbounded with smapling effort, the coefficient of variation for each dataset was relatively stable. In general, the coefficient of variation increased slightly with sampling depth only to decline once smaller samples were removed from the analysis at higher sampling depths. Interestingly, the coefficient of variation between Shannon diveristy values decreased towards zero with increased sampling effort and the coefficient of variation between Bray-Curtis distances tended to increased. Regardless, the coefficients of variation were relatively small.

Most studies have a high level of sequencing coverage. To explore the concern over loss of sequencing depth further, I calculated the Good's coverage for the observed data. The median coverage for each dataset ranged between 89.4 and 99.8% for the Seagrass and Human datasets, respectively (Figure 7). When I rarefied each dataset to the size of the smallest community in the dataset, with the exception of the Seagrass, Rice, and Stream datasets, the median coverage for the rarefied communities was still greater than 90%. These results suggest that most studies had a level of sequencing coverage that aligned with the diversity of the communities. Next, I used the null model for each dataset to ask how much sequencing

effort was required to obtain higher levels of coverage. To obtain 95 and 99% coverage, an average of 2.70 and 101.20-fold more sequence data was estimated to be required than was required to obtain 90% coverage, respectively (Figure 7). As suggested by the simulated coverages curve in Figure 7, these estimates are conservative. Regardless, the sequencing effort required to acheive higher sequencing depth would likely limit the number of samples that could be sequenced when controlling for costs. Although it may be disconcerting to rarefy to a sequencing depth that is considerably lower than that obtained for the best sequenced community in a study, sequencing coverage for many environments is probably adequate even at the lower sequencing depth. Of course, the results above have demonstrated that rarefaction is necessary to avoid problems with making inferences.

Discussion

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- Rarefy your data
- Problems with recommended methods...
 - Many recommended methods are borrowed from gened expression analysis
 - Meaning of zeroes in data structural vs. below limit of detection
- · Factors that determine what number of sequences to rarefy to
- Need better methods of pooling libraries that result in more even distribution of sequences across samples
- · Rarefy your data

Materials and Methods

Choice of datasets. The specific studies were selected because their data was publicly accessible through the Sequence Read Archive, the original investigators sequenced the V4 region of the 16S rRNA gene using paired 250 nt reads, and my previous familiarity with the data. The use of paired 250 nt reads to sequence the V4 region resulted in a near complete two-fold overap of the V4 region resulting in high quality contigs with a low sequencing error rate (1). These data were processed through the standard sequence curation pipeline to generate operational taxonomic units (OTUs) using the mothur software package (1, 2).

Null community model. Null models were generated by randomly assigning each sequence in the study to an OTU and sample while keeping constant the number of sequences per sample and the total number

of sequences in each OTU. Because the construction of the null models was a stochastic process, 100 replicates were geneated for each study.

10%/5% model

65 Remove model

Null treatment model

Size-based treatment model

Data availability.

Reproducible data analysis.

doi: 10.1038/nmeth.2658. Should CSS be used in alpha diversity analysis? No. The richness and shannon diversity values were no different from those obtained with the raw abundances. From the paper, "The relative proportion of the features is unaffected by the normalization". CSS paper refers to relative abundance approach as total-sum normalization (TSN)

Robust CLR - remove zero counts and calculate CLR One CLR - add a pseudocount of 1 to all observations Nudge CLR - add a pseudocount of 1/total number of sequences in sample Zero CLR - Impute the value of zeroes using zCompositions package

https://edepot.wur.nl/ 547087. "Log- ratio PCA is designed to give results that are library size- independent. However, as we demonstrated mathematically and with examples based on simulated and real data, log- ratio PCA becomes library size- dependent, if there are many infrequent taxa (many zeroes) and library sizes differ largely. In this situation, the row centring used in log- ratio PCA brings an effect of r (the row mean of the log- transformed counts) in the clr- transformed matrix. Note that this effect is irrespective of whether or not these infrequent taxa are genuine or due to sequencing noise or allocation error. This library size dependence is unexpected in the sense that, after applying the clr, the transformed matrix is free of the effect of the row totals for strictly positive data (yij>0 for all i and j). We additionally demonstrate that library size variability causes a loss in power in detecting an effect of x with log- ratio RDA. If there is additionally a correlation between treatment and the library size, the type 1 error for detecting the effect of x can be seriously inflated."

https://www.nature.com/articles/s41522-020-00160-w. "An important characteristic of a feature table is that it is typically sparse, sometimes as many as ~90% are zero entries21, which creates a challenge for analyzing rare taxa. A quick and simple strategy to deal with excess zeros is to add a small positive constant (e.g. 1) called pseudo-count14,22 to each cell of the feature table. The addition of a pseudo-

count becomes necessary when using methods of analysis that require log transformation of the observed counts. Even though adding a pseudo-count is simple and widely used, the choice of the pseudo-count is ad hoc. Studies have shown that differential abundance or clustering results could be sensitive to the choice of pseudo count23,24. Although different values of pseudo counts have been discussed in the literature23,24,25,26, to the best of our knowledge, there is no consensus on how to choose the optimal value. Other strategies involve modeling zero counts by some probability models21,27."

https://academic.oup.com/bioinformatics/article/34/16/2870/4956011 https://www.ncbi.nlm.nih.gov/pmc/articles/PMC6755255/

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Table 1. Summary of studies used in the analysis. For all studies, the number of sequences used from each study was rarefied to the smallest sample size. A graphical representation of the distribution of sample sizes for each study and the samples that were removed from each study are provided in Figure S1.

Study (Ref)	Samples	Total	Median	Range of	SRA study
		sequences	sequences	sequences	accession
Bioethanol (3)	95	3,970,972	16,014	3,690-356,027	SRP055545
Human (4)	490	20,828,275	32,452	10,439-422,904	SRP062005
Lake (5)	52	3,145,486	69,205	15,135-110,993	SRP050963
Marine (6)	7	1,484,068	213,091	132,895-256,758	SRP068101
Mice (1)	348	2,785,641	6,426	1,804-30,311	SRP192323
Peromyscus (7)	111	1,545,288	12,393	4,454-33,502	SRP044050
Rainforest (8)	69	936,666	11,464	4,880-37,403	ERP023747
Rice (9)	490	22,623,937	43,399	2,777-192,200	SRP044745
Seagrass (10)	286	4,135,440	13,538	1,830-45,076	SRP092441
Sediment (11)	58	1,151,389	17,606	7,686-67,763	SRP097192
Soil (12)	18	932,563	50,487	46,622-58,935	ERP012016
Stream (13)	201	21,017,610	90,621	8,931-394,419	SRP075852

Figure 1.

Figure S1.