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A framework for assessing 16S rRNA marker-gene survey data analysis methods using mixtures.

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Olson et al. Page 2 of 31

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Abstract

Background: Analysis of 16S rRNA marker-gene surveys , used to characterize prokaryotic microbial communities, may be performed by numerous a variety of bioinformatic pipelines and downstream analysis methods. However, appropriate assessment datasets and statistics are needed as there is limited guidance on how to decide between methods, appropriate data sets and statistics for assessing these methods are needed. We developed a mixture dataset with real data complexity and an expected value for assessing available analysis methods. Mixtures of environmental samples are useful for assessment as they provide values calculated from measurements of the unmixed samples and the mixture design that can be compared to values recovered by each bioinformatic method. While experiments mixing complex samples have been used to assess other sequencing methods such as RNAseq, they have yet to be used to assess 16S rRNA sequencing.

Results: We developed an assessment framework for 16S rRNA bioinformatic pipelines and downstream analysis methods. We generate an assessment dataset using sequencing analysis methods based on a two-sample titration mixture design. The sequencing data were processed using multiple bioinformatic pipelines, dataset and metrics to evaluate OTU count table characteristics. Our qualitative assessment evaluates feature presence/absence exploiting features only present in unmixed samples or titrations by testing if random sampling can explain their observed relative abundance. Our quantitative assessment evaluates how well relative and differential abundance values agree with values expected from the mixture design. We evaluated count tables generated by three commonly used bioinformatic pipelines as demonstration: i) DADA2 a sequence inference method, ii) Mothur a de novo clustering method, and iii) QIIME with which uses open-reference clustering. The mixture dataset was used to qualitatively and quantitatively assess count tables generated using the pipelines.

The qualitative assessment was used to evalute features only present in unmixed samples and titrations. The abundance Qualitative assessment indicated that the majority of Mothur and QIIME features specific to unmixed samples and or titrations were explained by sampling alone. However, for random sampling alone but not DADA2 over a third of the unmixed sample and titration specific feature abundance could not be explained by sampling alone. The quantitative assessment evaluated pipeline performance by comparing observed to expected relative and differential abundance values. Overall the observed relative abundance features. When combined with assessments of count table sparsity, these results indicate that DADA2 has a higher false negative rate whereas Mothur and QIIME have higher false positive rates. Quantitative assessment indicated that, overall, observed relative abundance and differential abundance values were consistent with the expected values. Though outlier features were observed across all pipelines . expected values for all three pipelines. We also identified subsets of features measured with high error by all pipelines evaluated. We could not identify the source of bias in these poor performing features based on previously studied sources of bias, indicating that further analysis of potentially unknown and unaccounted for biases is warranted.

Conclusions: Using a novel mixture dataset and assessment methods we quantitatively and qualitatively. We developed a novel framework for assessing 16S rRNA marker-gene survey analysis methods based on mixture experiments. To demonstrate the assessment framework we evaluated count tables generated using three bioinformatic pipelines. The dataset and methods assessment framework developed for this study will serve as a valuable community resource for assessing 16S rRNA marker-gene survey bioinformatic methods.

Keywords: 16S rRNA gene; assessment; bioinformatic pipeline; normalization; differential abundance

Olson et al. Page 3 of 31

¹Background 2 Targeted sequencing of the 16S rRNA gene , commonly known as 16S rRNA 2 ³marker-gene-surveys, is a commonly used method for characterizing microbial³ ⁴communities, microbiomesis commonly used to characterize microbial communities. ⁴ ⁵The 16S rRNA marker-gene-survey measurement process includes molecular ⁵ ⁶(e.g. PCR and sequencing) and computational steps (e.g., sequence clustering)⁶ ⁷[1]. Molecular steps are used steps to selectively target and sequence the 16S rRNA⁷ ⁸gene from prokaryotic organisms within a sample . The computational steps and ⁸ ⁹computational steps [1] computational steps convert the raw sequence data into⁹ ¹⁰a matrix with feature (e.g., operational taxonomic units) count table of feature ¹⁰ ¹¹relative abundance values , feature abundance relative to all other features, for ¹¹ ¹²each sample [1]. Both molecular and computational measurement process steps ¹² ¹³processes contribute to the overall measurement bias and dispersion [2, 1, 3]. Proper ¹³ ¹⁴measurement method evaluation allows for The need for datasets characterizing ¹⁴ ¹⁵complex microbial communities with some degree "ground truth" has emerged in ¹⁵ ¹⁶ order to properly characterize the accuracy of the characterization of how individual 16 ¹⁷steps impact the measurement processes as a whole and determine where to focus ¹⁷ ¹⁸efforts for improving the measurement process. Appropriate datasets and methods ¹⁸ ¹⁹are needed to evaluate the 16S rRNA marker-gene-survey measurement process. ¹⁹ ²⁰A sample or dataset with "ground truth" is needed to characterize measurement ²⁰ ²¹process accuracy. marker-gene-survey measurement process. Diverse bioinformatic pipelines used to generate count tables produce data²² ²³with diverse characteristics. For example the commonly used QIIME, Mothur, ²³ ²⁴and DADA2 pipelines produce feature sets and count tables with different ²⁴ ²⁵characteristics. Mothur uses de novo clustering for feature inference [4, 5]. Pairwise ²⁵ ²⁶distances used in clustering are calculated from a multiple sequence alignment. ²⁶ ²⁷Quality filtered paired-end reads are merged into contigs, then aligned to a reference ²⁷ ²⁸multiple sequence alignment, followed by the removal of uninformative positions. ²⁸ ²⁹As a result the feature set representative sequences are shorter than the input²⁹ ³⁰ amplicons. For the QIIME open-reference clustering pipeline merged paired-end ³⁰ ³¹reads are first assigned to reference cluster centers [6, 7]. Next, unassigned reads³¹ ³²are clustered de novo. Unlike Mothur, the QIIME pipeline clustering method³² ³³uses pairwise sequence distances calculated from pairwise sequence alignments. ³³ ³⁴As a result, the QIIME pairwise distances are calculated using the full amplicon ³⁴ ³⁵ sequence, whereas Mothur pairwise distances are calculated using multiple sequence ³⁵ ³⁶ alignment with only informative positions. The DADA2 pipeline uses a probability ³⁶ ³⁷ model and maximization expectation algorithm for feature inference [8]. Unlike ³⁷ ³⁸distance-based clustering methods employed by the Mothur and QIIME pipelines, ³⁸ ³⁹DADA2 parameters determine if low abundance sequences are grouped with a³⁹ ⁴⁰higher abundance sequence. Numerous studies have evaluated quantitative and qualitative characteristics of 41 ⁴²the 16S rRNA measurement process using mock communities, simulated data, and ⁴² ⁴³environmental samples. To Mock communities are commonly used to assess the qualitative characteristics 44 ⁴⁵ of the 16S rRNA sequencing measurement process mock communities are commonly ⁴⁵ ⁴⁶used [9]. As the number of organisms in the mock community is known, the total ⁴⁶

Olson et al. Page 4 of 31

¹[9]. The use of mock communities in this fashion shows that surveys often result in ¹ ²number of features can be compared to the expected value. The number of observed ³features in a mock community is significantly higher than the expected number of ³ ⁴organism—that are significantly higher than the underlying features in the mock⁴ ⁵community [10]. The higher than expected number of features is often attributed ⁵ ⁶to sequencing and PCR artifacts as well as reagent contaminants [3, 11]. A notable ⁶ ⁷exception to this is mock community benchmarking studies evaluating sequencing ⁷ ⁸inference method is count tables generated using feature inference methods, such ⁸ ⁹as DADA2 [8]. Sequence inference methods which aim to reduce the number of ⁹ ¹⁰sequence artifact features. While mock communities have an expected number of ¹⁰ ¹¹features and composition features from sequence artifacts by using statistical models ¹¹ ¹²to group sequences by both similarity and abundance. Nonetheless, while mock ¹² ¹³communities are useful in this type of assessment, they lack the feature diversity and ¹³ ¹⁴ relative abundance diversity and dynamic range of feature present in real samples ¹⁴ The quantitative characteristics Quantitative assessment of 16S rRNA sequence 16 ¹⁷data are normally assessed using mock communities and simulated data . Mock ¹⁷ ¹⁸communities of equimolar and staggered concentration are used to assess relative ¹⁸ ¹⁹ abundance estimate quantitative accuracy [10] is informative but provides an ¹⁹ ²⁰incomplete characterization of the measurement process. Results from relative abun-²⁰ ²¹dance estimates using mock communities generated from mixtures of single organ-²¹ ²²ism's DNA have shown taxonomic specific effects where individual taxa are under ²² ²³or over represented in a sample. For example, Gram-negative bacteria have higher ²³ ²⁴extraction efficiency compared to Gram-positive bacteria, and are thus likely over ²⁴ ²⁵represented in count tables[12, 13]. Mismatches in the primer binding sites are ²⁵ ²⁶also responsible for taxonomic specific biases [3, 14, 15]. Additionally, taxon spe-²⁶ ²⁷cific biases due to sequence template properties such as GC content, secondary ²⁷ ²⁸structure, and gene flanking regions have been observed [16, 17, 15]. Simulated ²⁸ ²⁹count tables However, due to limited community complexity the applicability of ²⁹ ³⁰mock community assessment results to more complex environmental samples is ³⁰ ³¹unknown. Environmental sample complexity can be modeled using simulated and ³¹ ³²have been used to assess differential abundance method (fold change differences in ³²) ³³relative abundance) methods, where specific taxa are artificially overrepresented over ³³ ³⁴represented in one set of samples compared to another [18]. Using However, using ³⁴ ³⁵simulated data to assess log fold-change estimates only evaluates the computational³⁵ ³⁶steps of the measurement process. ³⁷ Quantitative and qualitative assessment can also be performed using sequence ³⁷ ³⁸data generated from mixtures of environmental samples. While simulated data and ³⁸ ³⁹mock communities are useful in evaluating and benchmarking new methods, one³⁹ ⁴⁰needs to consider that methods optimized for mock communities and simulated data ⁴⁰ ⁴¹ are not necessarily optimized for the sequencing error profile and feature diversity ⁴¹ ⁴²of real samples. Data from environmental samples, which are real samples, are real ⁴³environmental samples are often used to benchmark new molecular laboratory and ⁴³ 44 computational methods. However, without an expected value to compare to expected 44 ⁴⁵values for use in assessment, only measurement precision, or agreement with other ⁴⁵ ⁴⁶methods—can be evaluated. By mixing environmental samples, expected values ⁴⁶

Olson et al. Page 5 of 31

AssessmentFramework.pdf

Figure 1 Assessment Framework. A) Count tables evaluated by the assessment framework are generated from the assessment dataset using marker-gene survey bioinformatic pipelines. Count table rows are features identified by the bioinformatic pipeline and column are samples, four PCR replicates (labeled A-D) were sampled for PRE and POST and titrations, to simplify the diagram only three titrations are shown. B) Pictorial depiction of abundance values of the seven feature types observed and used in the assessment framework, C) Qualitative and quantitative assessment metrics used in the assessment framework. The artifactual feature proportion metric (AFP) is a qualitative assessment of feature presence/absence based on unmixed-specific or titration-specific artifactual features. Sparsity (SPAR) is a qualitative assessment of the proportion of observed features in each sample relative to the total observed features. Relative abundance metric (Rel) plot is a quantitative assessment of the relationship between the observed and expected relative abundance values. The difference is used to calculate the error rate (|Obs-Exp|/Exp) from which the bias metric (median(error)) and variance metric (RCOV) are calculated. The differential abundance (Diff) metric assesses the relationship between the expected log fold-change and estimated log fold-change is shown. Points represent the log fold-change between two titrations, point text indicates the titrations compared. A linear model is fit to the data. The model fit information is used for the differential abundance bias (1 - slope) and variance metrics (R^2) . Each feature type in (B) is labeled with the assessments shown in (C) in which they are employed.

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 18 $^{19}{\rm are}$ calculated using information from the unmixed samples and mixture design. 19

20Mixtures of environmental samples were previously used to evaluate gene expression 20 21 measurements [19, 20, 21].

In the present study, we developed a mixture dataset of extracted DNA from 22 23 human stool samples for assessing. Here we present a framework for assessing 23 24 computational methods used to analyze 16S rRNA sequencing. The mixture 24 25 datasets were processed using three bioinformatic pipelines. We developed metrics 25 26 for qualitative and quantitative assessment of the bioinformatic pipeline results 26 27 The quantitative results were similar across pipelines, but the qualitative results 27 28 varied by pipeline. We have made both marker-gene-survey data. The framework is 28 29 comprised of a 16S rRNA two-sample titration dataset, generated using mixtures 29 30 of human stool sample DNA extracts, along with metrics to assess the quantitative 30 31 and qualitative characteristics of count tables generated using marker-gene-survey 31 32 computational methods. To demonstrate usage of this assessment framework, we 32 33 evaluated three bioinformatic pipelines. Both the dataset and metrics developed in 33 4 this study publicly available for evaluating are publicly available and can be used 34 35 to evaluate and optimize new and existing bioinformatic pipelines.

³⁷Results

³⁸Two-Sample Titration DesignAssessment Framework

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³⁹ Samples collected at multiple timepoints during a Enterotoxigenic E. coli (ETEC)³⁹

⁴⁰vaccine trial [22] were used to generate a two-sample titration dataset for assessing ⁴⁰

⁴¹the-Our framework assesses the qualitative and quantitative characteristics of the ⁴¹

⁴²16S rRNA marker-gene survey measurement process . Samples from five trial ⁴²

⁴³participants were selected for our measurement process (Fig. 1). The framework ⁴³

44 evaluates count tables generated by bioinformatic pipelines from a dataset developed 44

⁴⁵specifically for use in this framework. The qualitative assessment provides insight ⁴⁵

⁴⁶into how much confidence a user can have in feature presence/absence. The⁴⁶

Olson et al. Page 6 of 31

experimentalDesign.pdf

Figure 2 Sample selection and experimental design for the two-sample titration 16S rRNA marker-gene-survey assessment dataset. A) Pre- and post-exposure (PRE and POST) samples from five vaccine trial participants were selected based on *Escherichia coli* abundance measured using qPCR and 454 16S rRNA sequencing (454-NGS), data from Pop et al. [23]. Counts represent normalized relative abundance values for 454-NGS and copies of the heat-labile toxin gene per μL , a marker gene for ETEC, for qPCR. PRE and POST samples are indicated with orange and green data points, respectively. Grey points are other samples from the vaccine trial time series. B) Proportion of DNA from PRE and POST samples in titration series samples. PRE samples were titrated into POST samples following a log_2 dilution series. The NA titration factor represents the unmixed PRE sample. C) PRE and POST samples from the five vaccine trial participants, subjects, were used to generate independent two-sample titration series. The result was a total of 45 samples, 7 titrations + 2 unmixed samples times 5 subjects. Four replicate PCRs were performed for each of the 45 samples resulting in 190 PCRs.

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  quantitative assessment evaluates the bias and variance of relative and differential 14
  abundance estimates.
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<sup>17</sup>Assessment Dataset - Mixture Design
<sup>18</sup>Using mixtures of environmental samples we generated a dataset with expected <sup>18</sup>
<sup>19</sup> values for use in our assessment framework. For mixture datasets, expected values
<sup>20</sup>can be obtained using information from unmixed samples and the mixture design.
<sup>21</sup>Our mixture dataset uses a two-sample titration dataset. Trial participants <sup>21</sup>
<sup>22</sup>(subjects) and sampling timepoints were selected based on E. coli abundance <sup>22</sup>
<sup>23</sup>data collected using qPCR and 16S rRNA sequencing from Pop et al. [23]. Only <sup>23</sup>
24 individuals with no E. coli detected in samples collected from trial participants prior 24
<sup>25</sup>to ETEC exposure (PRE) were used for our two-samples titrations. Post ETEC<sup>25</sup>
<sup>26</sup>exposure (POST) samples were identified as the timepoint titration mixture design, <sup>26</sup>
<sup>27</sup>where DNA collected from five vaccine trial participants before and after exposure to <sup>27</sup>
<sup>28</sup>ETEC with the highest-pathogenic E. Escherichia coliconcentration for each subject<sup>28</sup>
<sup>29</sup>(Fig. 2A). Due to limited sample availability, for E01JH0016 the timepoint with the <sup>29</sup>
30 second highest E. coli concentration was used as the POST sample. Independent 30
<sup>31</sup>titration series were generated for each subject, where POST samples were titrated <sup>31</sup>
<sup>32</sup>into PRE samples with POST proportions of 1/2, 1/4, 1/8, 1/16, 1/32, 1/1,024, <sup>32</sup>
<sup>33</sup> and 1/32,768 was mixed following a log<sub>2</sub> dilution series (Fig. 2B). Unmixed (PRE<sup>33</sup>
<sup>34</sup> and POST) sample DNA concentration was measured using NanoDrop ND-1000<sup>34</sup>
<sup>35</sup>(Thermo Fisher Scientific Inc. Waltham, MA USA). Unmixed samples were diluted <sup>35</sup>
^{36}to 12.5 nq/\mu L in tris-EDTA buffer before mixing.
37 ). Each sample was sequenced in quadruplicate. For our two-sample titration 37
<sup>38</sup>mixture design, the expected feature relative abundance can be is calculated using <sup>38</sup>
<sup>39</sup>equation (1), where \theta_i, is the proportion of POST DNA in titration i, q_{ij} is the <sup>39</sup>
<sup>40</sup>relative abundance of feature j in titration i, and the relative abundance of feature ^{40}
^{41}j in the unmixed PRE and POST samples is q_{pre,j} and q_{post,j}. Throughout the rest
<sup>42</sup>of the manuscript, samples collected prior to and after E. coli exposure are referred <sup>42</sup>
<sup>43</sup>to as PRE and POST respectively.
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                                                                                                     45
                                                                                                 (1)^{46}
        q_{ij} = \theta_i q_{post,j} + (1 - \theta_i) q_{pre,j}
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Olson et al. Page 7 of 31

Sample selection and experimental design for the two-sample titration 16S ¹
² rRNA marker-gene-survey assessment dataset. A) Pre- and post-exposure (PRE ²
³ and POST) samples from five vaccine trial participants were selected based on ³
⁴ Escherichia coli abundance measured using qPCR and 454 16S rRNA sequencing ⁴
⁵ (454-NGS), data from Pop et al. [23]. Counts represent normalized relative ⁵
⁶ abundance values for 454-NGS and copies of the heat-labile toxin gene per μL , ⁶
⁷ a marker gene for ETEC, for qPCR. PRE and POST samples are indicated with ⁷
⁸ orange and green data points, respectively. Grey points are other samples from the ⁸
⁹ vaccine trial time series. B) Proportion of DNA from PRE and POST samples in ⁹
¹⁰ titration series samples. PRE samples were titrated into POST samples following a ¹⁰
¹¹ log ₂ dilution series. The NA titration factor represents the unmixed PRE sample. ¹¹
¹² C) PRE and POST samples from the five vaccine trial participants, subjects, were ¹²
¹³ used to generate independent two-sample titration series. The result was a total of
¹⁴ 45 samples, 7 titrations + 2 unmixed samples times 5 subjects. Four replicate PCRs ¹⁴
¹⁵ were performed for each of the 45 samples resulting in 190 PCRs. 16
Quantutive Assessment
¹⁸ The qualitative assessment shows how well pipelines differentiate true biological ¹⁸
¹⁹ sequences from measurement process artifacts. Inadequate processing of artifacts.
²⁰ results in false positive and false negative features where false positives are features
²¹ in a count table that are not present in the sequenced sample and false negative ²¹
²² features are biological sequences in a sample not represented in the count table. ²²
²³ Our qualitative assessment methods characterize the artifactual feature proportion ²³
²⁴ (the frequency of artifactual features in a count table) by estimating the proportion ²⁴
²⁵ of titration- and unmixed-specific features (Fig. 1B) that cannot be explained by ²⁵
²⁶ sampling alone. We combine the artifactual feature proportion assessment results ²⁶
²⁷ with sparsity estimates to hypothesize whether the artifactual features are primarily ²⁷
28 false positives or negatives. Sparsity is defined as the fraction of 0 valued cells in 28
²⁹ the count table (Fig. 1C).
30
³¹ Dataset characteristics
$^{32}Quantitative_Assessment$
³³ To evaluate count table abundance values, our quantitative assessment uses ³³
³⁴ error, bias, and variance metrics (Fig. 1C). Error metrics measure agreement ³⁴
³⁵ between observed and expected abundance values. The bias and variance metrics ³⁵
³⁶ summarise feature-level performance. Bias metrics summarise the overall agreement ³⁶
³⁷ with expected values and the variance metric characterizes the distribution of ³⁷
38 the agreement. Overall, pipeline performance is evaluated by comparing count 38
³⁹ table metric distributions. Additionally, feature-level metrics are indicators of
feature-specific biases. 40
teature-specific biases. 41 41
⁴² Assessment Dataset Characterization and Validation
Assessment Dataset Characterization and Validation 43 To assure the mixture dataset is suitable for use in our assessment framework, 43
44 we first validated the titration series and raw sequence data. The mixture dataset 45
⁴⁵ had sufficient sample coverage, reads per sample, and read quality for use in our ⁴⁵
⁴⁶ assessment framework. The number of reads per sample and distribution of base ⁴⁶

Olson et al. Page 8 of 31

¹ quality scores by position was consistent across subjects (Fig. S5). There were
$^28.9548 \times 10^4 \ (152,267 - 3,195)$ sequences per sample, median and range. Average ²
³ base quality score was greater than 30 over the length of the amplicon when ³
⁴ considering both forward and reverse reads (Fig. S5B).
⁵ Additionally, we characterized subject specific differences to inform the interpretation
⁶ of our assessment results. No subject specific differences in base quality score ⁶
⁷ were observed (Fig. S5). However, average read depth was greater for E01JH004 ⁷
⁸ compared to the other individuals (Fig. S5). Community composition differences
⁹ between PRE and POST samples and individuals was characterized using alpha ⁹
¹⁰ and beta diversity (Fig. S6). Overall alpha diversity was higher for POST except ¹⁰
¹¹ for E01JH0011, though differences in diversity between PRE and POST varied ¹¹
¹² by individual. Based on the beta diversity the community composition within ¹³
¹³ individuals differed between the PRE and POST samples. Note that assessment ¹³
¹⁴ metrics defined above and results reported below are based on within subject ¹⁴ 15
comparisons.
10 validate the two-sample titration assessment dataset, we evaluated two
¹⁷ assumptions about the titrations: (1) The samples were mixed volumetrically in a ¹⁷
$^{18}log_2$ dilution series according to the mixture design. (2) The unmixed PRE and 18
¹⁹ POST samples have the same proportion of prokaryotic DNA. To validate the ¹⁹
²⁰ sample volumetric mixing exogenous DNA (ERCC plasmids) were spiked into the ²⁰
²¹ unmixed samples before mixing and quantified using qPCR (Fig. S1B). The stool ²¹
²² samples used to generate the mixtures have both eukaryotic (primarily human) ²²
²³ DNA and prokaryotic DNA. If the proportion of prokaryotic DNA differs between ²³
²⁴ the unmixed samples, then the amount of DNA from the unmixed samples in a ²⁴
²⁵ titration targeted by 16S rRNA gene sequencing is not consistent with the mixture ²⁵
²⁶ design. We quantified the proportion of prokaryotic DNA in the unmixed samples ²⁶
²⁷ using a qPCR assay targeting the 16S rRNA gene (Fig. S1C).
Our assessment dataset validation results indicated that the samples were 28
²⁹ volumetrically mixed according to the mixture design (Table S1) but prokaryotic ²⁹
³⁰ DNA proportion varied across the titration series (Fig. S2). To account for ³⁰
³¹ deviations from the mixture design due to differences in the proportion of ³¹
³² prokaryotic DNA in the unmixed samples, we estimated the proportion of ³²
³³ POST in each titration using the 16S rRNA sequencing data (Fig. S3) and the ³³
³⁴ estimated POST proportions were used in our assessment metric calculations. See ³⁴
³⁵ Supplemental Material for the assessment dataset validation methods and results. ³⁵
36 36
³⁷ Count Table Assessment Demonstration 37
³⁸ Next, we demonstrate the utility of our assessment framework on count tables
³⁹ generated using three different bioinformatic pipelines; DADA2, Mothur and ³⁹
⁴⁰ QHME. First, we provide high level summary statistics for initial insight into how ⁴¹ the count tables differ. Next, we compare the assessment framework results for the ⁴¹
⁴² three count tables. 43 44 43
⁴⁴ Count Table Characteristics Sequence dataset characteristics. (A) Distribution in ⁴⁵
⁴⁵ the number of reads per barcoded sample (Library Size) by individual. Boxplots ⁴⁵
⁴⁶ summarize data distribution with horizontal bar as median, boxes indicating ⁴⁶

Olson et al. Page 9 of 31

1 Table 1 Summary statistics for the different bioinformatic pipelines. DADA2 is a denoising sequence inference pipeline, QIIME is an open-reference clustering pipeline, and Mothur is a de-novo clustering 2 pipeline. No template controls were excluded from summary statistics. Sparsity is the proportion of 2 0's in the count table. Features is the total number of OTUs (QIIME and Mothur) or SVs (DADA2) in the count. Sample coverage is the median and range (minimum-maximum) per sample total 4 abundance. Drop-out rate is the proportion of reads removed while processing the sequencing data for 4 each bioinformatic pipeline.

Pipelines	Features	Sparsity	Total Abundance	Drop-out Rate
DADA2	3144	0.93	68649 (1661-112058)	0.24 (0.18-0.59)
Mothur	38358	0.98	53775 (1265-87806)	0.4 (0.35-0.62)
QIIME	11385	0.94	25254 (517-46897)	0.7 (0.62-0.97)

interquartile range, whiskers $\pm 1.5 \times IQR$, and black points outliers. The dashed 11 horizontal line indicates overall median library size. Excluding one PCR replicate 12 from subject E01JH0016 titration 5 that had only 3,195 reads. (B) Smoothing 13 spline of the base quality score (BQS) across the amplicon by subject. Vertical lines 14 indicate approximate overlap region between forward and reverse reads. Forward 15 reads go from position 0 to 300 and reverse reads from 464 to 164.

Relationship between the number of reads and features per sample by bioinformatic pipeline. (A) Scatter plot of observed features versus the number of reads per sample. (B) Observed feature distribution by pipeline and individual. Excluding one PCR replicate from subject E01JH0016 titration 5 with only 3,195 reads, and the Mothur E01JH0017 titration 4 (all four PCR replicates), with 1,777 observed features.

Comparison of dataset taxonomic composition across pipelines. Phylum (A) and Order (B) relative abundance by pipeline. Taxonomic groups with less than 1% total relative abundance were grouped together and indicated as other. Pipeline genus-level taxonomic assignment set overlap for the all features (C) and the upper quartile genera by relative abundance for each pipeline (D).

We first characterize the number of reads per sample and base quality score distribution. The number of reads per sample and distribution of base quality scores by position was consistent across subjects (Fig. ??). Two barcoded experimental samples had less than 35,000 reads. The rest of the samples with less than 35,000 reads were no template PCR controls (NTC). Excluding one failed reaction with 22,700 reads and NTCs, there were 8.9548 × 10⁴ (3195-152267) sequences per sample, 33 median and range. Forward reads had consistently higher base quality scores relative 4 to the reverse reads with a narrow overlap region with high base quality scores for 5 both forward and reverse reads (Fig. ??B).

The resulting The count tables generated using the four bioinformatic pipelines of twere characterized for three bioinformatic pipelines vary in pre-processing and seature inference methods. These differences are reflected in the count table number of features, sparsity, and filter total abundance, and drop-out rate (Table 1, Figs. sparsity, sparsity, and filter total abundance, and drop-out rate (Table 1, Figs. sparsity, sparsity, and filter total abundance, and drop-out rate (Table 1, Figs. sparsity). The pipelines evaluated employ different approaches for handling sparsity reads resulting in large differences in the drop-out rate and the pipeline sparsity fraction of raw sequences not included in the count table (Table 1). QIIME pipeline sparsity has the highest drop-out rate and number of features per sample but fewer total sparsity fractions than Mothur. The targeted amplicon region has a relatively small overspansity fraction of the pipelines sparsity fraction of the count table (Table 1). The features than Mothur features than Mothur features and number of features per sample but fewer total sparsity fractions are reflected in the count table (Table 1). The features than Mothur features per sample but fewer total sparsity fractions are reflected in the count table (Table 1).

Olson et al. Page 10 of 31

¹ ends of the reads especially the reverse reads resulting in a high proportion of un- ¹
² successfully merged reads pairs (Fig. ??B). Furthermore, increasing the drop-out ²
³ S5B). Further increasing the filter rate, QIIME excludes singletons — (features only ³
⁴ observed once in the dataset , to remove potential sequencing artifacts from the ⁴
⁵ dataset. QHME and DADA2 pipelines were similarly sparse (the fraction of zero ⁵
⁶ values in count tables)despite differences in the number of features and drop-out ⁶
⁷ rate. The expectation is that this mixture dataset will be less sparse relative to ⁷
⁸ other datasets. This is due to the redundant nature of the samples where the 35 ⁸
⁹ titration samples are derived directly from the 10 unmixed samples, along with ⁹
¹⁰ four PCR replicates for each sample. With sparsity greater than 0.9 for the three ¹⁰
¹¹ pipelines it is unlikely that any of the pipelines successfully filtered out a majority ¹¹
¹² of the sequencing artifacts.
¹³ Dataset taxonomic assignments Feature taxonomic composition also varied by ¹³
¹⁴ pipeline (Fig. ??). S8). The three pipelines generated unique feature sets in terms ¹⁴
¹⁵ of sequence length and amplicon position (see pipeline description). Therefore, ¹⁵
¹⁶ we used feature taxonomic assignments for cross-pipeline community composition ¹⁶
¹⁷ comparison. Phylum and order relative abundance is similar across pipelines (Fig. ¹⁷
¹⁸ ??A S8A & B). The observed differences are attributed to different taxonomic ¹⁸
¹⁹ classification methods and databases used by the pipelines. DADA2 and QHME ¹⁹
²⁰ pipelines differed from Mothur and QHME for Proteobacteria and Bacteriodetes. ²⁰
²¹ Regardless of the relative abundance threshold, for genus sets most genera were 21
²² unique to individual pipelines (Fig. ???C_S8C & D). Sets —(shared taxa between ²²
²³ pipelines, with QIIME had the fewest genera, excluding the DADA2-QIIME set. ²³
²⁴ QIIME was the only pipeline to use open-reference clustering and the Greengenes ²⁴
²⁵ database. Mothur and DADA2 both used the SILVA dataset. The Mothur and ²⁵
²⁶ DADA2 pipeline use different implementations implementations of the RDP naïve ²⁶
²⁷ Bayesian classifier, which may be partially responsible for the Mothur, unclustered, ²⁷
²⁸ and DADA2 differences.
29
³⁰ Titration Series Validation
³¹ Qualitative Assessment
³² To validate the two-sample titration dataset for use in abundance assessment ³²
³³ we evaluated two assumptions about the titrations: 1. The samples were mixed ³³
34 volumetrically in a log_2 dilution series according to the mixture design. 2. The 34
35 unmixed PRE and POST samples have the same proportion of prokaryotic DNA.35
³⁶ The stool samples used to generate the mixtures have both eukaryotic (primarily ³⁶
³⁷ human) DNA and prokaryotic DNA. If the proportion of prokaryotic DNA differs ³⁷
³⁸ between the unmixed samples, then the amount of DNA from the unmixed samples ³⁸
³⁹ in a titration targeted by 16S rRNA gene sequencing is not consistent with the ³⁹
⁴⁰ mixture design. To validate the sample volumetric mixing exogenous DNA was ⁴⁰
⁴¹ spiked into the unmixed samples before mixing and quantified using qPCR . To ⁴¹
⁴² evaluate if the PRE and POST samples had the same proportion of prokaryotic ⁴²
⁴³ DNA total prokaryotic DNA in the titrations samples was quantified using a qPCR ⁴³
⁴⁴ assay targeting the 16S rRNA gene.
45 45
⁴⁶ Spike-in qPCR results 46
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Olson et al. Page 11 of 31

qualPlot-1.pdf

Figure 3 Prokaryotic DNA concentration Distribution of (ng/ulA) across titrations measured using a 16S rRNA qPCR assay. Separate linear models, Prokaryotic DNA concentration versus θ were fit observed count values for each individual, titration-specific (TS) features and R^2 (B) expected count values for unmixed-specific (US) features by pipeline and p-values were reported individual. Red lines indicate negative slope estimates and blue lines positive slope estimates. p-value The orange horizontal dashed line indicates significant difference from the expected slope a count value of 0. The grey regions indicate the linear model 95% confidence interval 1. (C) Artifactual feature proportion (Art. Multiple test correction was performed using the Benjamini-Hochberg method Feat. One of the E01JH0004 PCR replicates Prop.) for titration 3 titration-specific and ($\theta = 0.125$ D) was identified as an outlier, unmixed-specific features with a concentration of 0.003, and was excluded from an adjusted p-value 0.05 for the linear model Bayesian hypothesis test and binomial test respectively. The linear model slope was still significantly different from 0. We failed to accept the null hypothesis when the outlier was included p-value 0.05, indicating that the discrepancy between the feature only being observed in the titrations or unmixed samples cannot be explained by sampling alone.

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Fitration series volumetric mixing was validated by quantifying ERCC plasmids 16 ₁₇spiked into the POST samples using qPCR. The qPCR assay standard curves had a $_{\rm 18}{\rm high\ level\ of\ precision\ with\ }R^2$ values close to 1 and amplification efficiencies between $_{\rm 18}$ 190.84 and 0.9 for all standard curves indicating the assays were suitable for validating the titration series volumetric mixing (Table ??). For our log₂ two-sample-titration 21 mixture design the expected slope of the regression line between titration factor and 22 Ct is 1, corresponding to a doubling in template DNA every PCR cycle. The qPCR 22 assays targeting the ERCCs spiked into the POST samples had R² values and slope estimates close to 1 (Table ??). Slope estimates less than one were attributed to 24 assay standard curve efficiency less than 1 (Table ??). ERCCs spiked into PRE samples were not used to validate volumetric mixing as PRE sample proportion₂₆ differences were too small for qPCR quantification. The expected C_t difference for the entire range of PRE concentrations in only 1. When considering the quantitative 28 imitations of the qPCR assay these results confirm that the unmixed samples were volumetrically mixed according to the two-sample titration mixture design. ERCC Spike-in qPCR assay information and summary statistics. ERCC is the ERCC identifier for the ERCC spike-in, Assay is TaqMan assay, and Length and GC are the size and GC content of the qPCR amplicon. The Std. R^2 and Efficiency (E) statistics were computed for the standard curves. R^2 and slope for titration qPCR results for the titration series. Subject ERCC Assay Length Std. R^2 E R^2 SlopeE01JH0004 012 Ac03459877-a1 77 0.9996 86.19 0.98 0.92E01JH0011 157 Ac03459958-a1 71 0.9995 87.46 0.95 0.90E01JH0016 108 Ac03460028-a1 74 0.9991 87.33 0.95 0.84E01JH0017 002 Ac03459872-a1 69 0.9968 85.80 0.89 0.93E01JH0038 38 035 Ac03459892-a1 65 0.9984 86.69 0.95 0.94 39 $^{40}Prokaryotic\ DNA\ Concentration$ Observed changes in prokaryotic DNA concentration across titrations indicate 41 ⁴²the proportion of prokaryotic DNA from the unmixed PRE and POST samples in a ⁴³titration is inconsistent with the mixture design (Fig. ??). A qPCR assay targeting ⁴³ 44 the 16S rRNA gene was used to quantify the concentration of prokaryotic DNA in 44 ⁴⁵the titrations. An in-house standard curve with concentrations of 20 ng/ul, 2ng/ul, ⁴⁵ ⁴⁶and 0.2 ng/ul was used, with efficiency 91.49, and R² 0.999. If the proportion of 46 Olson et al. Page 12 of 31

¹ prokaryotic DNA is the same between PRE and POST samples the slope of the ¹
² concentration estimates across the two-sample titration would be 0. For subjects ²
³ where the proportion of prokaryotic DNA is higher in the PRE samples, the slope ³
⁴ will be negative, and positive when the proportion is higher for POST samples. ⁴
⁵ The slope estimates are significantly different from 0 for all subjects excluding ⁵
⁶ E01JH0011 (Fig. ??). These results indicate that the proportion of prokaryotic ⁶
⁷ DNA is lower in POST when compared to the PRE samples for E01JH0004 and ⁷
⁸ E01JH0017 and higher for E01JH0016 and E01JH0038.
9
¹⁰ Theta Estimates
¹¹ Human stool sample DNA extracts vary in the proportion of eukaryotic (primarily ¹¹
¹² human) and prokaryotic DNA in the sample. To account for differences in the ¹²
¹³ proportion of prokaryotic DNA in PRE and POST samples (Fig. ??) we inferred ¹³
¹⁴ the proportion of POST sample prokaryotic DNA in a titration, θ , using the 16S ¹⁴
¹⁵ rRNA sequencing data (Fig. ??). Overall the relationship between the inferred and ¹⁵
¹⁶ mixture design θ values were consistent across pipelines but not subject whereas ¹⁶
¹⁷ the θ estimate 95% CI varied by both subject and pipeline. For study subjects ¹⁷
¹⁸ E01JH0004, E01JH0011, and E01JH0016 the inferred and mixture design θ values ¹⁸
¹⁹ were in agreement, in contrast to study subjects E01JH0017 and E01JH0038. For ¹⁹
²⁰ E01JH0017 the inferred values were consistently less than the mixture design values. ²⁰
²¹ Whereas for E01JH0038 the inferred values were consistently greater than the ²¹
²² mixture design values. These results were consistent with the qPCR prokaryotic ²²
²³ DNA concentration results with significantly positive slopes for E01JH0004 and ²³
²⁴ E01JH0016 and significantly negative slope for E01JH0038 (Fig. ??).
Theta estimates by titration, biological replicate, and bioinformatic pipeline. The 25
²⁶ points indicates mean estimate of 1000 bootstrap theta estimates and errorbars 95% ²⁶
points indicates mean estimate of 1000 bootstrap theta estimates and errorbars 93% ²⁷ confidence interval. The black bar indicate expected theta values. Theta estimates ²⁷
²⁸ below the expected theta indicate that the titrations contain less than expected
²⁹ bacterial DNA from the POST sample. Theta estimates greater than the expected
30 theta indicate the titration contains more bacterial DNA from the PRE sample 30 31.1
³¹ than expected. 32
Weasurement Assessment
³⁴ Next, we assessed the qualitative and quantitative nature of 16S rRNA measurement ³⁴
³⁵ process using our two-sample titration dataset. For the qualitative assessment, we ³⁵
³⁶ analyzed the relative abundance of features only observed in To evaluate feature
³⁷ presence-absence, the framework's qualitative assessment measures artifactual ³⁷
³⁸ feature proportion and count table sparsity. Low abundance features present only in ³⁸
³⁹ unmixed samples or titrations. These features are not expected given the titration ³⁹
40 experimental design. The quantitative assessment evaluated relative and differential 41
abundance estimates.
42 42
⁴³ Qualitative Assessment
⁴⁴ Distribution of (A) observed count values for titration-specific features and (B) ⁴⁴
⁴⁵ expected count values for unmixed-specific features by pipeline and individual. ⁴⁵
⁴⁶ The grange horizontal dashed line indicates a count value of 1. (C) Proportion of

Olson et al. Page 13 of 31

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<sup>1</sup>unmix-specific features and (D) titration-specific features with an adjusted p-value <sup>1</sup>
<sup>2</sup>< 0.05 for the Bayesian hypothesis test and binomial test respectively. We failed to <sup>2</sup>
<sup>3</sup>accept the null hypothesis when the p-value < 0.05, indicating that the discrepancy
<sup>4</sup>between the feature only being observed in the titrations or unmixed samples cannot <sup>4</sup>
<sup>5</sup>be explained by sampling alone.
<sup>6</sup> Unmixed- and titration-specific titration samples are expected due to random<sup>6</sup>
<sup>7</sup>sampling. Unmixed- and titration-specific features were observed for all pipelines<sup>7</sup>
*(titration-specific titration-specific: Fig. 3A, unmixed-specific unmixed-specific: Fig. 3A)
<sup>93</sup>B). For mixture datasets low abundance features present only in unmixed samples<sup>9</sup>
<sup>10</sup>and mixtures are expected due to random sampling Overall, the DADA2 count<sup>10</sup>
11table had the largest number of artifactual features (Table S3). A summary of the 11
12 titration-specific artifactual features is provided in the supplementary material.
13 We next assessed the proportion of these artifactual features that could be 13
14explained by sampling effects alone. For our two-sample titration dataset there were 14
15unmixed-specific, there were unmixed-specific features with expected counts not 15
16which could not be explained by sampling alone for all individuals and bioinformatic 16
17pipelines (Fig. 3C). However, the proportion of unmixed-specific unmixed-specific 17
18 features that could not be explained by sampling alone varied by bioinformatic 18
19pipeline. DADA2 had the highest proportion of unmixed-specific features not 19
20 explained by sampling unmixed-specific artifactual features whereas QIIME had 20
21the lowest proportion . Consistent which is consistent with the distribution of 21
220bserved counts for titration-specific features more of the DADA2 features could not 22
23be explained by sampling alone compared to the other pipelines titration-specific 23
<sub>24</sub>feature observed counts (Fig. 3D). Overall, the
   We expected this mixture dataset to be less sparse relative to other datasets due<sub>25</sub>
26 to the redundant nature of the samples where the 35 titration samples are derived 26
ardirectly from the 10 unmixed samples, along with four PCR replicates for each
sample. We observed overall sparsity of 0.93 and 0.94 for DADA2 count table had
29 the largest number of observed features inconsistent with the titration experiment,
30 design, while the same phenomenon is significantly reduced in the other pipelines 30
and QIIME respectively, and a higher value of 0.98 for Mothur 1).
   To account for differences in microbial community composition across the five
33 individuals we also measured sparsity at the individual level (Table S2). Sparsity
at the individual-level is lower than overall sparsity for all three pipelines. In this
case, average sparsity across individuals for 0.70 and 0.76 for DADA2 and Mothur,
 while QIIME had a lower average sparsity across individuals of 0.56. Differences in _{36}
 alpha and beta diversity for the five individual unmixed samples are consistent with
  individual level sparsity and therefore reflects differences in individual microbial
 community composition.
   Based on the artifactual feature proportions and count table sparsity, \mathrm{DADA2}^{39}
 artifactual features are likely due to false negative features, whereas the Mothur and 40
<sup>41</sup>QIIME high sparsity values were attributed to false positive features. Based on the
<sup>42</sup> observed sparsity levels it is unlikely that any of the pipelines successfully filtered <sup>42</sup>
out a majority of the sequencing artifacts. Both unmixed- and titration-specific 43
44 features that can and cannot be explained by sampling alone contribute to sparsity
<sup>45</sup> and the differences in the artifactual feature proportion and sparsity provide insight <sup>45</sup>
<sup>46</sup>into how the pipelines treat sequencing artifacts.
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Olson et al. Page 14 of 31

relAbuError-1.pdf

Figure 4 Relative abundance assessment. (A) A linear model of the relationship between the expected and observed relative abundance. The dashed grey line indicates expected 1-to-1 relationship. The plot is split by individual and color is used to indicate the different bioinformatic pipelinespipeline indicated by line color. A negative binomial model was used to calculate an average relative abundance estimate across the four PCR replicates. Points To highlight quantitative performance for higher abundance features, points with observed and expected relative abundance values less than 1/medianlibrary size (total abundance) were excluded from the data used to fit the linear modelplot. (B) Relative abundance error rate (|expected - observed|/expected) distribution by individual and pipeline.

relAbuErrorMetrics-1.pdf

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Figure 5 Comparison of pipeline relative abundance assessment feature-level error metrics. Distribution of feature-level relative abundance (A) bias metric - median error rate and (B) variance - robust coefficient of variation (RCOV = (IQR)/|median|RCOV = IQR/|medianerrorrate|) by individual and pipeline. For both the bias and variance metrics lower values are better. Boxplot outliers, $1.5 \times IQR$ from the median were excluded from the figure to prevent extreme metric values from obscuring metric value visual comparisons.

18 19 Quantitative Assessment 19

Maximum feature-level error rate bias (median error rate) and variance (robust²⁰
 21COV) by pipeline and individual.

22 For the relative abundance assessment

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²⁴Relative Abundance Assessment To assess count table feature relative abundance²⁴ ²⁵values, we evaluated the consistency of the observed and expected relative abun-²⁵ ²⁶dance estimates for a feature and titration as well as feature-level bias and vari-²⁶ ²⁷ance. The PRE and POST estimated relative abundance and inferred θ values were²⁷ ²⁸used to calculate titration and relative abundance error rates. Relative abundance²⁸ ²⁹error rate is defined as |exp - obs|/exp, where exp and obs is the expected and ²⁹ ³⁰observed relative abundance. To control for biases in feature inference, the three³⁰ ³¹pipelines were compared to an unclustered dataset. The unclustered count table³¹ ³²was generated using the 40,000 most abundant features from Mothur's initial³² ³³preprocessing (see Methods for details). Unclustered pipeline θ estimates were used ³³ ³⁴to calculate the error rates for all pipelines to prevent over-fitting. Only features ob-³⁴ ³⁵served in all PRE and POST PCR replicates and PRE and POST specific features³⁵ ³⁶were included in the analysis (Table ??). PRE and POST specific features were ³⁶ ³⁷defined as present in all four of the PRE or POST PCR replicates, respectively, but ³⁷ ³⁸none of the PCR replicates for the other unmixed samples. There is lower confidence³⁸ ³⁹in PRE or POST feature relative abundance when the feature is not observed all³⁹ ⁴⁰4 PCR replicates, therefore these features were not included in the analysis, S3). ⁴⁰ ⁴¹Overall, agreement between inferred and observed relative abundance was high for ⁴¹ ⁴²all individuals and bioinformatic pipelines (Fig. 4A). The error rate distribution ⁴² ⁴³was similarly consistent across pipelines, including long tails (Fig. 4B). To assess quantitative accuracy across pipelines, we compared the feature-level ⁴⁵relative abundance error rate bias (median error rate, Fig. 5A) and variance 45

 $^{46}(RCOV = (IQR)/|median| \text{ Fig. 5B})$ across pipelines and individuals and variance 46

Olson et al. Page 15 of 31

logFCerror-1.pdf

Figure 6 Differential abundance quantitative assessment. (A) Linear model or of the relationship between log fold-change estimates estimated and expected values log fold-change relative abundance between titrations for PRE-specific and PRE-dominant features by pipeline and individual, line color indicates pipelines. Dashed grey line indicates expected 1-to-1 relationship between the estimated and expected log fold-change. (B) Log fold-change error (|exp-est|) distribution by pipeline and individual.

logFcErrorMetrics-1.pdf

Figure 7 Feature-level $\frac{\log - \operatorname{fold}}{\operatorname{differential}}$ abundance assessment. Log-fold change error bias (A) and variance (B) metric distribution by subject and pipeline. The bias (1-slope) and variance (R^2) metrics are derived from the linear model fit to the estimated and expected log fold-change values for individual features. Boxplot outliers, $1.5 \times IQR$ from the median were excluded from the figure to prevent extreme metric values from obscuring metric value visual comparisons.

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¹⁵using mixed effects models. To control for subject specific differences, subject was ¹⁵ ¹⁶included in the model as a random effect. Large bias and variance metric values were ¹⁶ ¹⁷observed for all pipelines (Table ??). Features with large bias and variance metrics, ¹⁷ $^{18}1.5 \times IQR$ from the median, were deemed outliers. To prevent these outlier features 18 19 from biasing the comparison they were not used to fit the mixed effects model. 19 20 Multiple comparisons test (Tukey) was used to test for significant differences in 20 21 feature-level bias and variance between pipelines. A one-sided alternative hypothesis 21 22was used to determine which pipelines had smaller feature-level error rateS3).22 23 Feature-level relative abundance error rate bias (median error rate, Fig. 5A) was 23 24significantly different between pipeline, but no statistically significant differences 24 25were observed for the variance metric, (RCOV = (IQR)/|median|, Fig. 5B) across 25 26pipeline. The Mothur, DADA2, and QIIME feature-level bias-biases were all sig-26 ₂₇nificantly different from each other $(p < 1 \times 10^{-8})$. DADA2 had the lowest mean₂₇ ₂₈feature-level bias (0.2), followed by Mothur (0.28), with QIIME having the high-₂₈ 29est bias (0.33) (5B). Large variance metric values were observed for all individuals 29 30 and pipelines (Table ?? S3). The feature-level variance was not significantly differ-30 ₃₁ent between pipelines; Mothur = 0.83, QIIME = 0.71 and DADA2 = 1 (Fig.₃₁ ₃₂5B). We evaluated whether poor feature-level relative abundance metrics can be₃₂ ₃₃attributed to specific taxonomic groups or phylogenetic clades. While a significant 24 overall phylogenetic signal was detected for both the bias and variance metric, no 35 specific taxonomic groups or phylogenetic clades were identified with exceptionally 35 36 poor performance in our assessment.

³⁷ Differential Abundance Assessment The agreement between log-fold change esti³⁸ mates and expected values were individual specific and consistent across pipelines
³⁹ (Fig. 6A). The individual specific effect was attributed to the fact that unlike rela⁴⁰ tive abundance assessment, the inferred θ values were not used to calculate expected
⁴¹ values. Inferred θ values were not used to calculate the expected values because all
⁴² of the titrations and the θ estimates for the higher titrations were included and they
⁴³ were not monotonically decreasing and therefore. Using the inferred θ resulted in
⁴⁴ unrealistic expected log fold-change values, e.g., negative log-fold changes for PRE
⁴⁵ specific features. The log-fold change estimates and expected values were consis⁴⁶ tent across pipelines with one notable exception. For: for subject E01JH0011, the ⁴⁶

Olson et al. Page 16 of 31

¹ Mothur log fold-change estimates were more consistent with expected values than ¹
² the other pipelines. However, as θ was not corrected for differences in the proportion ²
³ of prokaryotic DNA between the unmixed PRE and POST samples, it cannot be ³
⁴ said whether Mothur's performance was better than the other pipelines.
⁵ The log fold-change error distribution was consistent across pipelines (Fig. 6B). ⁵
⁶ There was a long tail of high error features in the error distribution for all pipelines ⁶
⁷ and individuals. The log fold-change estimates responsible for the long tail could not ⁷
⁸ be attributed to specific titration comparisons. Additionally, we compared log-fold ⁸
⁹ change error distributions for log-fold change estimates using different normalization ⁹
10 methods. Error rate distributions, including the long tails, were consistent across 10
¹¹ normalization methods. Furthermore, Seeing as the long tail was observed for the ¹¹
¹² unclustered data as well, the log-fold change estimates contributing to the long tail ¹²
¹³ are likely due to a bias associated with the molecular laboratory portion of the ¹³
¹⁴ measurement process and not the bioinformatic pipelines computational portion. ¹⁴
¹⁵ Exploratory analysis of the relationship between the log fold-change estimates and ¹⁵
¹⁶ expected values for individual features indicated that the long tails were attributed ¹⁶
¹⁷ to feature specific performance.
¹⁸ Feature-level log fold-change bias and variance metrics were used to compare ¹⁸
¹⁹ pipeline performance (Fig. 6). Similar to relative abundance, feature-level bias and ¹⁹
²⁰ variance metrics are defined as the $1-slope$ and R^2 for linear models of the esti- ²⁰
²¹ mated and expected log fold-change for individual features and all titration com- ²¹
²² parisons. For the bias metric, $1 - slope$, the desired value is 0 (i.e., log fold-change ²²
²³ estimate = log fold-change expected), with negative values indicating the log-fold ²³
²⁴ change was consistently underestimated and positive values consistently overesti- ²⁴
²⁵ mated. The linear model R^2 value was used to characterize the feature-level \log^{25}
²⁶ fold-change variance as it indicates consistency between log fold-change estimates ²⁶
²⁷ and expected values across titration comparisons. To compare bias and variance ²⁷
²⁸ metrics across pipelines, mixed-effects models were used. The log fold-change bias ²⁸
²⁹ and variance metrics were not significantly different between pipelines (Bias: $F = {}^{29}$
30 0, 2.51, p = 0.99, 0.08, 6B, Variance: F = 47.39, 0.23, p = 0, 0.8, Fig. 6C). 30 0
³¹ also evaluated whether poor feature-level metrics could be attributed to specifie ³¹
³² clades for taxonomic groups. Similar to the relative abundance estimate, while a ³²
³³ phylogenetic signal was detected for both the bias and variance metrics, no specific ³³
³⁴ taxonomic groups or phylogenetic clades that performed poorly were identified.
35
³⁶ Discussion
37 Mixtures of environmental samples have been used to assess RNAseq and microarray 37
³⁸ gene expression measurements [19, 20, 21]. However, this is the first time mixtures ³⁸
³⁹ have been used to assess microbiome measurement methods. We developed a novel ³⁹
⁴⁰ assessment framework utilizing a mixture dataset for evaluating marker-gene-survey ⁴⁰
⁴¹ computational methods (Fig. 1).
We assessed the quantitative and qualitative characteristics of count tables 42
⁴³ generated using different bioinformatic pipelines and 16S rRNA marker-gene survey ⁴³
⁴⁴ mixture dataset. The mixture dataset followed Using mixtures of environmental 44
⁴⁵ samples, expected values for use in assessment can be obtained using information ⁴⁵
⁴⁶ from unmixed samples and how the samples were mixed. Our assessment dataset ⁴⁶

Olson et al. Page 17 of 31

¹follows a two-sample titration mixture design, where DNA collected from five ¹ ²vaccine trial participants before and after exposure to pathogenic Escherichia coli² ³from five vaccine trial participants (subjects) were was mixed following a log_2 ³ ⁴dilution series (Fig. 2). Qualitative count table Count table qualitative character-⁴ ⁵istics were assessed using relative abundance information for features observed⁵ ⁶only in titrations (titration-specific) and unmixed samples . We quantitatively ⁶ ⁷assed count tables by comparing feature relative and differential abundance⁷ ⁸(unmixed-specific) (Fig. 1B). Statistical tests were used to determine if the absence⁸ 9 of unmixed-specific features from titrations or absence of titration-specific features 9 ¹⁰from unmixed samples could be explained by random sampling. Count tables¹⁰ ¹¹were quantitatively assessed by comparing observed feature relative abundance¹¹ ¹²and feature differential abundance estimates to expected values. Quantitative ¹² 13 performance was characterized using error rate, along with feature-level bias 13 14 variance metrics we developed (Fig. 1C). 14 15 ₁₆Count Table Assessment Demonstration 16 ₁₇We demonstrated our novel assessment approach by evaluating assessment framework ₁₈on count tables generated using different by three commonly used bioinformatic₁₈ ₁₉pipelines, QIIME, Mothur, and DADA2. The Mothur pipeline uses de novo₁₉ 20 clustering for feature inference [4, 5]. Pairwise distances used in clustering are 20 21 calculated using a multiple sequence alignment. The quality filtered paired end reads 21 22 are merged into contigs. The pipeline then aligns contigs to a reference multiple 22 23 sequence alignment and removes uninformative positions in the multiple sequence 2d alignment. The QHME pipeline uses open-reference clustering where merged 2d ₂₅paired-end reads are first assigned to reference cluster centers [6, 7]. Next QHME₂₅ 26 clusters unassigned reads de novo. Unlike Mothur, the QHME clustering method 26 ₂₇uses pairwise sequence distances calculated from pairwise sequence alignments. As a ₂₈ result, the QHME pairwise distances are calculated using the full ~436 bp sequences ²⁹ whereas Mothur pairwise distances were calculated using a 270 bp multiple sequence 30 alignment. The objective of any pipeline is to differentiate true biological sequences from measurement process artifacts along with accurate abundance estimates. Our qualitative assessment results, when combined with sparsity information 32 provides a new method for evaluating how well bioinformatic pipelines account for sequencing artifacts without loss of true biological sequences. Additionally, our 34 quantitative assessment results identified previously unknown feature specific biases $^{35}_{35}$ in abundance estimates. The qualitative assessment evaluates if titration- and unmixed-specific features can be explained by random sampling alone (Fig. 1B). Titration- and unmixed-specific features not explained by sampling are artifacts of the measurement process. These ³⁹ artifacts can be viewed as false-positives, not representative of actual sequences in a sample, or false-negatives, actual sequences in a sample not represented by ⁴¹count table features. Artifacts can be PCR errors such as chimeras, reads with ⁴²high sequencing error rates, or cross sample contamination [26, 27, 28]. Count table ⁴³sparsity information (the proportion of zero-valued cells) provides additional insight ⁴³ ⁴⁴into the qualitative assessment results. A high false negative rate provides an explanation for DADA2pipeline uses a 45 ⁴⁶probability model and maximization expectation algorithm for feature inference 46

Olson et al. Page 18 of 31

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<sup>1</sup>[8]. Unlike distance-based clustering methods employed by the Mothur and QHME<sup>1</sup>
 <sup>2</sup>pipelines, 's high proportion of artifact titration, and unmixed-specific features<sup>2</sup>
 <sup>3</sup>and count table having comparable sparsity to the other pipelines despite having
 <sup>4</sup>significantly fewer features (Fig. S5 and Table 1). The DADA2 parameters determine <sup>4</sup>
 <sup>5</sup>if low abundance feature inference algorithm may be aggressively grouping lower <sup>5</sup>
 <sup>6</sup>abundance true sequences with higher abundance sequences are grouped with a <sup>6</sup>
 <sup>7</sup>higher abundance sequence. As a control, we compared our quantitative assessment<sup>7</sup>
 <sup>8</sup>results for the three pipelines to a count table of unclustered features. The <sup>8</sup>
 <sup>9</sup>unclustered features were generated using the Mothur pipeline preprocessing <sup>9</sup>
<sup>10</sup>methods result, the low abundance sequences are not present in samples leading <sup>10</sup>
<sup>11</sup>to increased sparsity and high abundance unmixed- and titration-specific features. <sup>11</sup>
<sup>12</sup>This aggressive grouping of sequences is a design choice made by the algorithm <sup>12</sup>
<sup>13</sup>developers. The DADA2 documentation states that the default setting for OMEGA_A<sup>13</sup>
<sup>14</sup> is conservative to prevent false positives at the cost of increasing false negatives <sup>14</sup>
<sup>15</sup>[8]. Using the qualitative assessment methods described here, a user can adjust the <sup>15</sup>
<sup>16</sup>OMEGA_A parameter to obtain a false-negative rate appropriate for their study.
^{18}Quantitative Assessment While the relative abundance bias metric was sig-^{18}
<sup>19</sup> nificantly different between pipelines, overall, pipeline choice had minimal im-<sup>19</sup>
<sup>20</sup>pact on the quantitative assessment results when accounting for subject-specific<sup>20</sup>
<sup>21</sup>effects deviations in the proportion of prokaryotic DNA from PRE and POST<sup>21</sup>
<sup>22</sup>samples in a titration from the mixture design. Outlier features — (those with extreme<sup>22</sup>
<sup>23</sup> quantitative analysis bias and variance metrics.) were observed for all pipelines and <sup>23</sup>
<sup>24</sup>both relative and differential abundance assessments. Outlier features are not likely <sup>24</sup>
<sup>25</sup>a pipeline artifact
Outlier features could not be attributed to bioinformatic pipelines and are likely<sup>26</sup>
<sup>27</sup>due to biases in the molecular biology part of the measurement process. Outlier <sup>27</sup>
<sup>28</sup>features are unlikely pipeline artifacts as they were observed in count tables gener-<sup>28</sup>
<sup>29</sup> ated using the unclustered pipeline as well as standard bioinformatic pipelines. We<sup>29</sup>
<sup>30</sup>Additionally, we were unable to attribute outlier features to relative abundance val-<sup>30</sup>
<sup>31</sup>ues, log fold-change between unmixed samples, and sequence GC content. Features <sup>31</sup>
<sup>32</sup>Furthermore, features with extreme metric values were not limited to any specific<sup>32</sup>
<sup>33</sup>taxonomic group or phylogenetic clade. Outlier features could not be attributed to <sup>33</sup>
<sup>34</sup>bioinformatic pipelines and are likely due to biases in the molecular biology part<sup>34</sup>
<sup>35</sup> of the measurement process. PCR amplification is PCR amplification bias (a well-<sup>35</sup>
<sup>36</sup>known source of bias in the molecular biology part of the measurement process) <sup>36</sup>
<sup>37</sup> is one possible explanation for the outlier features [29]. Mismatches in the primer <sup>37</sup>
<sup>38</sup>binding regions impact PCR efficiency and are a potential cause for poor feature-<sup>38</sup>
<sup>39</sup>specific performance [30]. Additional research is needed before outlier features are <sup>39</sup>
<sup>40</sup>can be attributed to mismatches in the primer binding regions.
<sup>42</sup>Qualitative Assessment The qualitative assessment evaluated whether features <sup>42</sup>
<sup>43</sup>only observed in unmixed samples or titrations could be explained by sampling <sup>43</sup>
<sup>44</sup>alone. Features present only in titrations or unmixed samples not due to random <sup>44</sup>
<sup>45</sup>sampling are bioinformatic pipeline artifacts. These artifacts can be categorized <sup>45</sup>
<sup>46</sup>as false negative or false positive features. A false negative occurs when a lower <sup>46</sup>
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Olson et al. Page 19 of 31

¹abundance sequence representing an organism within the sample is clustered with a ¹ ²higher abundance sequence from a different organism. False positives are sequencing ³or PCR artifacts not appropriately filtered or assigned to an appropriate feature by ³ ⁴the bioinformatic pipeline. ⁵ Count table sparsity, the proportion of zero-valued cells, provides additional⁵ ⁶insight into the qualitative assessmentresults. A high rate of false negative ⁶ ⁷features is a potential explanation for Based on our assessment results, we⁷ ⁸suggest using DADA2 count table's poor performance in the qualitative assessment⁸ ⁹and comparable sparsity for feature-level abundance analysis, e.g. differential ¹⁰abundance testing. While DADA2 performed poorly in our qualitative assessment, ¹⁰ ¹¹the pipeline performed better in the quantitative assessment compared to the ¹¹ ¹²other pipelinesdespite having significantly fewer features (Fig. 3 and Table 1). ¹² ¹³The. Additionally, the DADA2 feature inference algorithm may be aggressively ¹³ ¹⁴grouping lower abundance true sequences with higher abundance sequences. As a ¹⁴ ¹⁵result, the low abundance sequences are not present in samples leading to increased ¹⁵ ¹⁶ sparsity and higher abundance unmixed- and titration-specific features. Adjusting ¹⁶ ¹⁷the-poor qualitative assessment results due to false-negative features are unlikely ¹⁷ ¹⁸to negatively impact feature-level abundance analysis. When determining which ¹⁸ ¹⁹pipeline to use for a study, users should consider whether minimizing false positives ¹⁹ ²⁰(DADA2parameters, specifically the OMEGA A parameter in setDada0pt, Along these²⁰ ²¹lines, the DADA2 documentation states that the default setting for OMEGA is ²¹ ²²conservative to prevent false positives at the cost of increasing false negatives [8].)²² ²³or false negatives (Mothur) is more appropriate for their study objectives. Based on ²³ ²⁴our findings we find that users of DADA2 can be more confident that an observed ²⁴ ²⁵feature represents a member of the microbial community and not a measurement ²⁵ ²⁶ artifact, but careful examination of sequences assigned to features of interest should ²⁶ ²⁷still be performed. ²⁸ False positive features provide an explanation for Mothur and OHME pipelines²⁸ ²⁹having lower proportion of unmixed- and titration-specific features not explained²⁹ ³⁰by sampling but high sparsity (Fig. 3 and Table 1). The statistical tests used ³⁰ ³¹to determine if the specific features could be explained by sampling alone only ³¹ ³²considers feature abundance. Therefore, the statistical test is not able to distinguish ³² 33 between true low abundance unmixed- and titration-specific features and low 33 ³⁴abundance sequence artifacts. Mothurand QHME count tables have ten times and ³⁴ 35 three times more features compared to DADA2, respectively (Table 1). While 35 ³⁶microbial abundance distributions are known to have long tails, it is likely that the ³⁶ ³⁷observed sparsity is an artifact of the 16S rRNA sequencing measurement process. ³⁷ ³⁸Similarly, significantly more features than expected are commonly observed for mock ³⁸ ³⁹community benchmarking studies evaluating the QHME and Mothur pipelines [24]. ³⁹ False positive features can be reduced, but not eliminated, using smaller amplicon⁴¹ ⁴²and prevalence filtering. The 16S rRNA region sequenced in the study is larger than ⁴² ⁴³the region the de-novo, and open clustering pipelines were developed for, potentially ⁴³ 44 explaining the higher than expected sparsity [24]. Kozich et al. [24] reduced the 44 45 sequence error rate from 0.29% to 0.06% by using paired-end reads that completely 45 ⁴⁶overlap. The larger region used in this study has a smaller overlap between the ⁴⁶

Olson et al. Page 20 of 31

² not allow for sequence error correction that occurs when a smaller amplicon is ²
³ used. However, even when targeting smaller regions of the 16S rRNA gene both the ³
⁴ de-novo (Mothur) and open-reference clustering (QHME) pipelines produced count
⁵ tables with significantly more features than expected in evaluation studies using ⁵
⁶ mock communities. Prevalence filtering is used to exclude low abundance features, ⁶
⁷ predominantly measurement artifacts [31]. For example, a study exploring the ⁷
⁸ microbial ecology of the Red-necked stint <i>Calidris ruficollis</i> , a migratory shorebird, ⁸
⁹ used a hard filter to validate their study conclusions are not biases by false positive ⁹
¹⁰ features . The study authors compared results with and without prevalence filter ¹⁰
¹¹ ensuring that the study conclusions were not biased by using the arbitrary filter or ¹¹
¹² including the low abundant features [32].
13
¹⁴ Using Mixtures to Assess 16S rRNA Sequencing
¹⁵ Mixtures of environmental samples have previously been used to assess RNAseq ¹⁵
¹⁶ and microarray gene expression measurements. However, this is the first time ¹⁶
¹⁷ mixtures have been used to assess microbiome measurement methods. Using our ¹⁷
¹⁸ mixture dataset we developed novel methods for assessing marker-gene-survey ¹⁸
¹⁹ computational methods. Our quantitative assessment allowed for the characterization ¹⁹
²⁰ of relative abundance values using a dataset with a larger number of features and ²⁰
²¹ dynamic range compared to mock community assessments. As a result, we identified ²¹
²² previously unknown feature specific biases. Based on our subject-specific results ²²
²³ observation, we recommend that studies using stool samples seeking inferences in a ²³
²⁴ longitudinal series of multiple subjects carefully estimate bacterial DNA proportions ²⁴
²⁵ and adjust inferences accordingly. Additionally, our qualitative assessment results, ²⁵
²⁶ when combined with sparsity information provide a new method for evaluating how ²⁶
²⁷ well bioinformatic pipelines account for sequencing artifacts without loss of true ²⁷
well biblinormatic bibelines account for sequencing artifacts without loss of true
²⁸ biological sequences. ²⁸
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28 biological sequences. 29 There were also 30 30
 28 biological sequences. 29 There were also 30 30 31 Using Mixtures to Assess 16S rRNA Sequencing - Lessons Learned 31
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Olson et al. Page 21 of 31

¹the proportion of prokaryotic DNA in the unmixed samples using qPCR data. Us-¹ ²ing the 16S sequencing data rRNA sequencing data, we inferred the proportion of ² ³prokaryotic DNA from the POST sample in each titration. However, the uncertainty ³ ⁴and accuracy of the inference method are not known, resulting in an unaccounted ⁴ ⁵for error source source of error. ⁶ A better method for quantifying sample prokaryotic DNA proportion or using ⁶ ⁷samples with consistent proportions would increase confidence in the expected value⁷ ⁸and, in-turn, error metric accuracy. Limitations in the prokaryotic DNA qPCR as-⁸ ⁹say's concentration precision limits the assay's suitability s suitability for use in ⁹ ¹⁰mixture studies. Digital PCR provides a more precise alternative to qPCR and ¹⁰ ¹¹is, therefore, a more appropriate method. Alternatively using samples where the ¹¹ ¹²majority of the DNA is prokaryotic would minimize this issue. Mixtures of environ-¹² ¹³mental samples can also be used to assess shotgun metagenomic methods as well. ¹³ ¹⁴As shotgun metagenomics is not a targeted approach, differences in the proportion ¹⁴ ¹⁵ of prokaryotic DNA in a sample would not impact the assessment results in the ¹⁵ ¹⁶same way as 16S rRNA marker-gene-surveys. ¹⁷ Using samples from a vaccine trial allowed for the use of a specific marker with an ¹⁷ ¹⁸expected response, E. coli, during methods development. However, the high level ¹⁸ ¹⁹ of similarity between the PRE and POST unmixed samples resulted in a limited ¹⁹ ²⁰number of features that could be used in the quantitative assessment results. Using ²⁰ ²¹more diverse samples to generate mixtures would address this issue. Alternatively, ²¹ ²²instead of mixing PRE and POST samples from the same individual, mixing PRE²² ²³and POST samples from different individuals would have resulted in additional ²³ ²⁴features for use in our quantitative assessment. While unmixed sample similarity ²⁴ ²⁵impacts the number of features that can be used in the quantitative assessment, ²⁵ ²⁶the qualitative assessment is not impacted by unmixed sample similarity. Finally, ²⁶ ²⁷a symmetric mixture design, for example one with unmixed PRE and POST ratios²⁷ ²⁸ of 1:4, 1:2, 1:1, 2:1, and 4:1, would provide a larger dynamic range of abundance ²⁸ ²⁹values for assessing both PRE and POST specific features. 30 31 31 Conclusions ³²Our two-sample-titration dataset and assessment methods Our assessment framework ³³can be used to evaluate and characterize bioinformatic pipelines and clustering ³³ 34 methods. The sequence dataset presented in this study can be processed with $16S^{34}$ ³⁵rRNA marker-gene survey analysis methods, in particular count tables produced by ³⁵ ³⁶any 16S rRNA bioinformatic pipeline. Our quantitative and qualitative assessment ³⁶ ³⁷can then be performed on the count table and the results compared to those obtained ³⁷ ³⁸using the pipelines presented here. The three pipelines we evaluated produced sets³⁸ ³⁹of features varying in total feature abundance, number of featuresper samples, ³⁹ ⁴⁰and total features. The objective of any pipeline is to differentiate true biological ⁴⁰ ⁴¹sequences from measurement process artifacts. In general, based on our evaluation ⁴¹ ⁴²results we suggest using We demonstrated our assessment framework with three ⁴² ⁴³commonly used bioinformatic pipelines. Our qualitative assessment results indicated ⁴³ 44 that the QIIME and Mothur pipelines produced count table with more false-positive 44 ⁴⁵features whereas the DADA2 for count table had more false-negative features. ⁴⁵ ⁴⁶Overall the three pipelines performed well in our quantitative assessment. However, ⁴⁶

Olson et al. Page 22 of 31

¹ feature-level abundance analysis, e.g. differential abundance testing. While DADA	42^{1}
² performed poorly in our qualitative assessment, the pipeline performed better	in ²
³ the quantitative assessment compared to the other pipelines. Additionally, analysis	sis^3
⁴ identified poorly performing features and the DADA2 poor qualitative assessme	nt^4
⁵ results due to false-negative features are unlikely to negatively impact sources.	of^5
⁶ bias responsible for this poor feature-level abundance analysis . When determining	~ ~
⁷ which pipeline to use for a study, users should consider whether minimizing fal	
⁸ positives (DADA2) or false negatives (Mothur) is more appropriate for their study	
⁹ objectives. When a sequencing dataset is processed using DADA2, the user can	
¹⁰ more confident that an observed feature represents a member of the microb	
¹¹ community and not a measurement artifact. Pipeline parameter optimization	
¹² could address DADA2 false-negative issue. For the Mothur and QHME pipeline	
¹³ prevalence filtering will reduce the number of false-positive features. Feature-lev	
¹⁴ quantitative performance are unknown. Therefore, feature-level results for any 10	
¹⁵ rRNA marker-gene survey should be interpreted with care, as the biases responsible	
¹⁶ for poor quantitative assessment are unknown. Addressing both of these issues 1	
¹⁷ quires advances in both the molecular biology and computational components	01
¹⁸ the measurement process.	19
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²⁰ Methods	21
²¹ Titration Validation	
²² qPCR was used to validate volumetric mixing and check for differences in t	
²³ proportion of prokaryotic DNA across titrations. To ensure the	23
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²⁵ Assessment Framework	25
²⁵ Assessment Framework ²⁶ To assess the qualitative and quantitative performance of marker-gene surv	25 ey ²⁶
²⁵ Assessment Framework	25 ey ²⁶
²⁵ Assessment Framework ²⁶ To assess the qualitative and quantitative performance of marker-gene surv	ey ²⁶
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²⁵ Assessment Framework ²⁶ To assess the qualitative and quantitative performance of marker-gene surv ²⁷ analysis methods we developed a framework utilizing our two-sample titrations we ²⁸ volumetrically mixed according to the mixture design, independent ERCC plasmi ²⁹ were spiked into the unmixed PRE and POST samples [33] (NIST SRM SRM 237) ³⁰ (Table ??). The ERCC plasmids were resuspended in 100 µL tris-EDTA buffer as	25 ere ²⁷ 6ds ²⁸ 74) ²⁹
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²⁵ Assessment Framework ²⁶ To assess the qualitative and quantitative performance of marker-gene surverally analysis methods we developed a framework utilizing our two-sample titrations we volumetrically mixed according to the mixture design, independent ERCC plasmid were spiked into the unmixed PRE and POST samples [33] (NIST SRM SRM 237) ³⁰ (Table ??). The ERCC plasmids were resuspended in 100 μL tris-EDTA buffer and μL of resuspended plasmids was spiked into the appropriate unmixed samples after unmixed sample concentration of the spiked into the appropriate unmixed sample and μL are spiked into unmixed samples after unmixed sample concentration of the spiked into the appropriate unmixed sample samples after unmixed sample concentration of the spiked into the appropriate unmixed samples after unmixed sample concentration of the spiked into the appropriate unmixed samples after unmixed sample concentration of the spiked into the appropriate unmixed samples after unmixed sample concentration of the spiked into the appropriate unmixed samples after unmixed sample concentration of the spiked into the appropriate unmixed samples after unmixed sample concentration of the spiked into the appropriate unmixed samples after unmixed sample concentration of the spiked into the appropriate unmixed samples after unmixe	25 ey ²⁶ ere ²⁷ eds ²⁸ eds ²⁸ nd ³⁰ de. ³¹ en ³² en ³² en ³³
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²⁵ Assessment Framework ²⁶ To assess the qualitative and quantitative performance of marker-gene surverally analysis methods we developed a framework utilizing our two-sample titrations we volumetrically mixed according to the mixture design, independent ERCC plasmid were spiked into the unmixed PRE and POST samples [33] (NIST SRM SRM 237) ³⁰ (Table ??). The ERCC plasmids were resuspended in 100 µL tris-EDTA buffer as 2 µL of resuspended plasmids was spiked into the appropriate unmixed sample samples after unmixed sample concentration was normalized to 12.5 ng/µL. POST sample ERCC plasmid abundance were adapted using TaqMan gene expression assays (FAM-MGB, Catalog # 444889). ThermoFisher) specific to each ERCC plasmid and TaqMan Universal Master Most HarmoFisher was also as a series of the sample and taqMan Universal Master Most HarmoFisher was also as a series of the sample and taqMan Universal Master Most HarmoFisher was also as a series of the sample and taqMan Universal Master Most HarmoFisher was a series of the sample and taqMan Universal Master Most HarmoFisher was a series of the sample and taqMan Universal Master Most HarmoFisher was a series of the sample and taqMan Universal Master Most HarmoFisher was a series of the sample and tagMan Universal Master Most HarmoFisher was a series of the sample and tagMan Universal Master Most HarmoFisher was a series of the sample and tagMan Universal Master Most HarmoFisher was a series of the sample and tagMan Universal Master Most HarmoFisher was a series of the sample and tagMan Universal Master Most HarmoFisher was a series of the sample and tagMan Universal Master Most HarmoFisher was a series of the sample and tagMan Universal Master Most HarmoFisher was a series of the sample and tagMan Universal Master Most HarmoFisher was a series of the sample and tagMan Universal Master Most HarmoFisher was a series of the sample and tagMan Universal Master Most HarmoFisher was a series of the sample and tagMan Universal Master Most HarmoFisher Was HarmoFisher Was Ha	28 ey 26 con 27 con 28 con 32
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²⁵ Assessment Framework ²⁶ To assess the qualitative and quantitative performance of marker-gene surversulation assess the qualitative and quantitative performance of marker-gene surversulation assess the qualitative and quantitative performance of marker-gene surversulation assessment evaluates the relative and differential abundance estimates. ²⁶ Assessment Framework To assess the qualitative and quantitative performance of marker-gene surversulation assessment evaluates the quantitative performance of marker-gene surversulation assessment evaluates the relative and differential abundance estimates.	25 25 26 26 26 26 26 26 26 26 26 26 26 26 26
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²⁵ Assessment Framework ²⁶ To assess the qualitative and quantitative performance of marker-gene surversular analysis methods we developed a framework utilizing our two-sample titrations we volumetrically mixed according to the mixture design, independent ERCC plasmic were spiked into the unmixed PRE and POST samples [33] (NIST SRM SRM 237 and (Table ??). The ERCC plasmids were resuspended in 100 µL tris-EDTA buffer at 12 µL of resuspended plasmids was spiked into the appropriate unmixed sample and Plasmids were spiked into unmixed samples after unmixed sample concentration was normalized to 12.5 ng/µL. POST sample ERCC plasmid abundance were quantified using TaqMan gene expression assays (FAM-MGB, Catalog # 444885 and TaqMan Universal Master Mas	2! ev 26 26 26 26 26 26 26 26 26 26 26 26 26
²⁵ Assessment Framework ²⁶ To assess the qualitative and quantitative performance of marker-gene surverally mixed according to the mixture design, independent ERCC plasmic volumetrically mixed according to the mixture design, independent ERCC plasmic were spiked into the unmixed PRE and POST samples [33] (NIST SRM SRM 237) (Table ??). The ERCC plasmids were resuspended in 100 µL tris-EDTA buffer at 12 µL of resuspended plasmids was spiked into the appropriate unmixed samples after unmixed sample concentration was normalized to 12.5 ng/µL. POST sample ERCC plasmid abundance were quantified using TaqMan gene expression assays (FAM-MGB, Catalog # 444885) (ThermoFisher) specific to each ERCC plasmid and TaqMan Universal Master Most II (Catalog # 4440040, ThermoFisher Waltham, MA USA). titration dataset (Fassessment evaluates the relative and differential abundance estimates. To check for differences in the proportion of bacterial DNA in the 40 and 41 Assessment Dataset - Mixture Design 42 To provide a dataset with real-world complexity and expected values for qualitatical and 12 provide a dataset with real-world complexity and expected values for qualitatical and 25 provide a dataset with real-world complexity and expected values for qualitatical and 26 provide a dataset with real-world complexity and expected values for qualitatical and 27 provide a dataset with real-world complexity and expected values for qualitatical and 27 provide a dataset with real-world complexity and expected values for qualitatical 48 provides a dataset with real-world complexity and expected values for qualitatical 49 provides a dataset with real-world complexity and expected values for qualitatical 40 provides and 40 provides a dataset with real-world complexity and expected values for qualitatical 40 provides and 40 provides an	2! ey 26 26 26 26 26 26 26 26 26 26 26 26 26
²⁵ Assessment Framework ²⁶ To assess the qualitative and quantitative performance of marker-gene surversular analysis methods we developed a framework utilizing our two-sample titrations were resulted into the unmixed according to the mixture design, independent ERCC plasmic were spiked into the unmixed PRE and POST samples [33] (NIST SRM SRM 237) (Table ??). The ERCC plasmids were resuspended in 100 µL tris-EDTA buffer at 12 µL of resuspended plasmids was spiked into the appropriate unmixed sample supplies applies were spiked into unmixed samples after unmixed sample concentration was normalized to 12.5 ng/µL. POST sample ERCC plasmid abundance were quantified using TaqMan gene expression assays (FAM MGB, Catalog # 444885). ThermoFisher) specific to each ERCC plasmid and TaqMan Universal Master Maste	2! ev 26 de 20 26 de
Assessment Framework 26 To assess the qualitative and quantitative performance of marker-gene surver analysis methods we developed a framework utilizing our two-sample titrations we volumetrically mixed according to the mixture design, independent ERCC plasmid were spiked into the unmixed PRE and POST samples [33] (NIST SRM SRM 237) (Table ??). The ERCC plasmids were resuspended in 100 µL tris EDTA buffer at 2 µL of resuspended plasmids was spiked into the appropriate unmixed sample Plasmids were spiked into unmixed samples after unmixed sample concentrations was normalized to 12.5 ng/µL. POST sample ERCC plasmid abundance were quantified using TaqMan gene expression assays (FAM-MGB, Catalog # 444880) (ThermoFisher) specific to each ERCC plasmid and TaqMan Universal Master M	2! 2! 2! 2! 2! 2! 2! 2! 2! 2! 2! 2! 2! 2
²⁵ Assessment Framework ²⁶ To assess the qualitative and quantitative performance of marker-gene surversular analysis methods we developed a framework utilizing our two-sample titrations were resulted into the unmixed according to the mixture design, independent ERCC plasmic were spiked into the unmixed PRE and POST samples [33] (NIST SRM SRM 237) (Table ??). The ERCC plasmids were resuspended in 100 µL tris-EDTA buffer at 12 µL of resuspended plasmids was spiked into the appropriate unmixed sample supplies applies were spiked into unmixed samples after unmixed sample concentration was normalized to 12.5 ng/µL. POST sample ERCC plasmid abundance were quantified using TaqMan gene expression assays (FAM MGB, Catalog # 444885). ThermoFisher) specific to each ERCC plasmid and TaqMan Universal Master Maste	25

Olson et al. Page 23 of 31

¹Trial participants (subjects) and sampling timepoints were selected based on E. ¹ ²coli abundance data collected using qPCR and 16S rRNA sequencing from Pop² ³et al. [23]. Only individuals with no E. coli detected in samples collected from ⁴trial participants prior to ETEC exposure (PRE) were used for our two-samples ⁴ ⁵titrations. Post ETEC exposure (POST) samples were identified as the timepoint ⁵ ⁶after exposure to ETEC with the highest *E. coli* concentration for each subject of ⁷(Fig. 2A). Due to limited sample availability, for E01JH0016 the timepoint with the second highest E. coli concentration was used as the POST sample. Independent titration series were generated for each subject. POST samples were titrated into ¹⁰PRE samples with POST proportions of 1/2, 1/4, 1/8, 1/16, 1/32, 1/1,024, and ¹⁰ 11/32,768 (Fig. 2B). Unmixed (PRE and POSTsamples, titration bacterial) sample DNA concentration was quantified using the Femto Bacterial DNA quantification kit (Zymo Research, Irvine CA). All samples were run in triplicate along with an in-house E. coli DNA log₁₀ dilution standard curve, qPCR assays were performed using the QuantStudio Real-Time qPCR (ThermoFisher). Amplification data and Ct values were exported as tsv files using QuantStudio TM Design and Analysis Software v1.4.1. Statistical analysis was performed on the exported data using 18 custom scripts in R [34]. The qPCR data and scripts used to analyze the data are available at . 20 21 22 Sequencing 23measured using NanoDrop ND-1000 (Thermo Fisher Scientific Inc. Waltham, MA23 ₂₄USA). Unmixed samples were diluted to 12.5 $ng/\mu L$ in tris-EDTA buffer before₂₄ ₂₅mixing. The resulting titration series was composed of 45 samples, seven titrations₂₅ 26and two unmixed samples for each of five subjects) the five subjects. 27 The 45 samples were processed using the Illumina 16S library protocol (16S27 28Metagenomic Sequencing Library Preparation, posted date 11/27/2013, down-28 29loaded from https://support.illumina.com). This protocol specifies an initial 29 30PCR of the 16S rRNA genePCR, followed by a sample indexing PCR, sample 30 31 concentration normalization, and sequencing. ³² A total of 192 16S rRNA PCR assays were run including four sequenced across³² 33two 96-well plates including four PCR replicates per sample and 12 no-template33 ³⁴controls, using Kapa HiFi HotStart ReadyMix reagents (KAPA Biosystems, Inc. ³⁴ ³⁵Wilmington, MA)... The initial PCR assay targeted the V3-V5 region of the 16S³⁵ 36 rRNA gene, Bakt_ 341 F and Bakt_ 806 R [14]. The V3-V5 region is 464 base pairs 36 ³⁷(bp) long, with forward and reverse reads overlapping by 136 bp, using 2 X 300 bp³⁷ ³⁸paired-end sequencing [35] (http://probebase.csb.univie.ac.at). Primer se-³⁸ ³⁹quences include overhang adapter sequences for library preparation (forward primer³⁹ ⁴⁰5'- TCG TCG GCA GCG TCA GAT GTG TAT AAG AGA CAG CCT ACG GGN⁴⁰ ⁴¹GGC WGC AG - 3' and reverse primer 5'- GTC TCG TGG GCT CGG AGA⁴¹ ⁴²TGT GTA TAA GAG ACA GGA CTA CHV GGG TAT CTA ATC C - 3'). For ⁴² ⁴³quality control, the PCR product Kapa HiFi HotStart ReadyMix reagents (KAPA⁴³ ⁴⁴Biosystems, Inc. Wilmington, MA) was used to PCR the 16S rRNA gene. The ⁴⁴ ⁴⁵PCR product amplicon size was verified using agarose gel electrophoresisto cheek ⁴⁵ ⁴⁶ amplicon size. Concentration measurements were made after the initial 16S rRNA ⁴⁶

Olson et al. Page 24 of 31

¹PCR, the indexing PCR, and normalization steps. DNA concentration was mea-²sured using the QuantIT Picogreen dsDNA Kit (Cat # P7589, ThermoFisher Sci-² ³entific) and fluorescent measurements were made with a Synergy2 Multi-Detection³ ⁴MicroPlate Reader (BioTek Instruments, Inc, Winooski, VT). ⁵ Initial PCR products were purified using 0.8X AMPure XP beads (Beckman Coul-⁵ ⁶ter Genomics, Danvers, MA) following the manufacturer's protocol. After purifica-⁶ ⁷tion, the 192 samples were indexed using the Illumina Nextera XT index kits A⁷ ⁸and D (Illumina Inc., San Diego CA) and then purified using 1.12X AMPure XP⁸ ⁹beads. Prior to pooling purified sample concentration was normalized using Sequal-⁹ ¹⁰Prep Normalization Plate Kit (Catalog n. A10510-01, Invitrogen Corp., Carlsbad, ¹⁰ ¹¹CA), according to the manufacturer's protocol. Pooled library concentration was ¹¹ ¹²checked using the Qubit dsDNA HS Assay Kit (Part# Q32851, Lot# 1735902, ¹² ¹³ThermoFisher, Waltham, MA USA). Due to the low pooled amplicon library DNA¹³ ¹⁴concentration, a modified protocol for low concentration libraries was used. The li-¹⁴ ¹⁵brary was run on an Illumina MiSeq, and base calls were made using Illumina Real¹⁵ ¹⁶Time Analysis Software version 1.18.54. The sequence data was deposited in the ¹⁶ ¹⁷NCBI SRA archive under Bioproject PRJNA480312. Individual SRA run accession ¹⁷ ¹⁸numbers and metadata in Supplemental Table. Sequencing data quality control met-¹⁸ ¹⁹rics for the 384 fastg sequence files (192 samples with forward and reverse reads) ¹⁹ ²⁰were computed using the Bioconductor Rqc package [36, 37]. The sequence data was ²⁰ ²¹deposited in the NCBI SRA archive under Bioproject PRJNA480312. Individual²¹ ²²SRA run accession numbers and metadata in Supplemental Table. 23 ²⁴Sequence Processing</sup> ²⁵Sequence data were processed using four bioinformatic pipelines: a de-novo clus-²⁵ ²⁶tering method - Mothur [5], an open-reference clustering method - QIIME [7], ²⁶ 27 and a sequence inference method - DADA2 [8], and unclustered sequences as a^{27} ²⁸control. The code used to run the bioinformatic pipelines is available at https: ²⁸ 29//github.com/nate-d-olson/mgtst pipelines. The Mothur pipeline follows the developer's MiSeq SOP [5, 24]. The pipeline was³⁰ ³¹run using Mothur version 1.37 (http://www.mothur.org/). We sequenced a larger³¹ ³²16S rRNA region, with smaller overlap between the forward and reverse reads, ³² ³³than the 16S rRNA region the SOP was designed. Pipeline parameters modified to ³³ ³⁴account for difference in overlap are noted for individual steps below. The Make-³⁴ ³⁵file and scripts used to run the Mothur pipeline are available https://github.³⁵ 36 com/nate-d-olson/mgtst_pipelines/blob/master/code/mothur. The Mothur 36 ³⁷pipeline includes an initial preprocessing step where the forward and reverse reads³⁷ ³⁸are trimmed and filtered using base quality scores and were merged into sin-³⁸ ³⁹gle contigs for each read pair. The following parameters were used for the ini-³⁹ ⁴⁰tial contig filtering, no ambiguous bases, max contig length of 500 bp, and max ⁴⁰ ⁴¹homopolymer length of 8 bases. For the initial read filtering and merging step, ⁴¹ ⁴²low-quality reads were identified and filtered from the dataset based on the pres-⁴² ⁴³ence of ambiguous bases, failure to align to the SILVA reference database (V119, ⁴³ 44 https://www.arb-silva.de/) [38], and identification as chimeras. Prior to align-44 ⁴⁵ment, the SILVA reference multiple sequence alignment was trimmed to the V3-V5⁴⁵ ⁴⁶region, positions 6,388 and 25,316. Chimera filtering was performed using UChime ⁴⁶

Olson et al. Page 25 of 31

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<sup>1</sup>(version v4.2.40) without a reference database [26]. OTU clustering was performed <sup>1</sup>
 <sup>2</sup>using the OptiClust algorithm with a clustering threshold of 0.97 [4]. The RDP<sup>2</sup>
 <sup>3</sup>classifier implemented in Mothur was used for taxonomic classification against the <sup>3</sup>
 <sup>4</sup>Mothur provided version of the RDP v9 training set [39].
 <sup>5</sup> The QIIME open-reference clustering pipeline for paired-end Illumina data was<sup>5</sup>
 <sup>6</sup>performed according to the online tutorial (Illumina Overview Tutorial (an IPython<sup>6</sup>
 <sup>7</sup>Notebook): open reference OTU picking and core diversity analyses, http://qime.<sup>7</sup>
<sup>8</sup>org/tutorials/) using QIIME version 1.9.1 [7]. Briefly, the QIIME pipeline uses<sup>8</sup>
 <sup>9</sup>fastq-join (version 1.3.1) to merge paired-end reads [40] and the Usearch algorithm<sup>9</sup>
<sup>10</sup>[41] with Greengenes database version 13.8 with a 97% similarity threshold [42] was <sup>10</sup>
<sup>11</sup>used for open-reference clustering.
<sup>12</sup> DADA2, an R native pipeline was also used to process the sequencing data<sup>12</sup>
<sup>13</sup>[8]. The pipeline includes a sequence inference step and taxonomic classifica-<sup>13</sup>
<sup>14</sup>tion using the DADA2 implementation of the RDP naïve Bayesian classifier [39]<sup>14</sup>
<sup>15</sup>and the SILVA database V123 provided by the DADA2 developers [38, https: <sup>15</sup>
16//benjjneb.github.io/dada2/training.html].
<sup>17</sup> The unclustered pipeline was based on the Mothur de-novo clustering pipeline, <sup>17</sup>
<sup>18</sup>where the paired-end reads were merged, filtered, and then dereplicated. Reads were <sup>18</sup>
<sup>19</sup> aligned to the reference Silva alignment (V119, https://www.arb-silva.de/), and <sup>19</sup>
<sup>20</sup>reads failing alignment were excluded from the dataset. Taxonomic classification<sup>20</sup>
<sup>21</sup>of the unclustered sequences was performed using the same RDP classifier imple-<sup>21</sup>
^{22}mented in Mothur used for the de-novo pipeline. To limit the size of the dataset ^{22}
^{23}the most abundant 40,000 OTUs (comparable to the Mothur dataset), across all^{23}
<sup>24</sup>samples, were used as the unclustered dataset.
25
                                                                                                       25
<sup>26</sup> Titration Proportion Estimates
<sup>27</sup>The following linear model was used to infer the proportion of prokaryotic DNA, <sup>27</sup>
<sup>28\theta</sup>, in each titration. Where \mathbf{Q}_i is a vector of titration i feature relative abundance <sup>28</sup>
<sup>29</sup>estimates and \mathbf{Q}_{pre} and \mathbf{Q}_{post} are vectors of feature relative abundance estimates<sup>29</sup>
30 for the unmixed PRE and POST samples. Feature relative abundance estimates 30
31 were calculated using a negative binomial model.
32
                                                                                                       32
33
                                                                                                       33
         \mathbf{Q}_i = \theta_i(\mathbf{Q}_{post} - \mathbf{Q}_{pre}) + \mathbf{Q}_{pre}
34
                                                                                                       34
35
   To fit the model and prevent uninformative and low abundance features from 36
<sup>37</sup>biasing \theta estimates, only features meeting the following criteria were used. Features
<sup>38</sup>included in the model were observed in at least 14 of the 28 total titration PCR<sup>38</sup>
<sup>39</sup>replicates (4 replicates per 7 titrations), demonstrated greater than 2-fold difference <sup>39</sup>
40 in relative abundance between the PRE and POST samples, and were present in 40
<sup>41</sup>either all four or none of the PRE and POST PCR replicates.
   16S rRNA sequencing count data is known to have a non-normal mean-variance 42
<sup>43</sup>relationship resulting in poor model fit for standard linear regression [18]. <sup>43</sup>
<sup>44</sup>Generalized linear models provide an alternative to standard least-squares regression. <sup>44</sup>
<sup>45</sup>The above model is additive and therefore \theta_i cannot be directly inferred in log-space. <sup>45</sup>
<sup>46</sup>To address this issue, we fit the model using a standard least-squares regression <sup>46</sup>
```

Olson et al. Page 26 of 31

¹ then obtained non-parametric 95 % confidence intervals for the θ estimates by ¹
² bootstrapping with 1000 replicates.
3
⁴ Qualitative Assessment
⁵ Our qualitative measurement
6
⁷ Artifactual Feature Proportion Our qualitative assessment evaluated features only ⁷
⁸ observed in unmixed samples (PRE or POST) , unmixed-specific, or titrations, ⁸
⁹ titration-specific. Unmixed- or only in titrations. The former we will refer to as ⁹
¹⁰ unmixed-specific features and the latter we will refer to as titration-specific features ¹
¹¹ (Fig. 1B). <i>Unmixed</i> - and <i>titration-specific</i> features are can arise from errors in ¹
¹² the PCR/sequencing, feature inference processes, or due to differences in sampling ¹³
¹³ depth(number of sequences)between the unmixed samples and titrations, artifacts ¹
¹⁴ of the feature inference process, or PCR/sequencing artifacts. Measurement process ¹
¹⁵ artifacts should be considered false positives or negatives. To provide context for ¹
¹⁶ the artifactual feature proportion results count table sparsity was used (Fig. 1C). ¹⁰
¹⁷ Sparsity is defined as the proportion of 0 valued cells in a matrix.
¹⁸ Hypothesis tests were used to determine if differences in sampling depth random
¹⁹ sampling alone, here sequencing depth, could account for <i>unmixed-specific unmixed</i> - ¹⁹
²⁰ and <i>titration-specific</i> features. p-values were adjusted for multiple comparisons us- ²¹
²¹ ing the Benjamini & Hochberg method [43]. For <i>unmixed-specific</i> features, the a ²
binomial test was used to evaluate if true feature relative abundance is less than 2:
²³ the expected relative abundance. A binomial test could not be used to evaluate The 24 in a way in facility for title for title and the country and the bound having a solid by 22 in a way in facility for title for title for the country and the bound have a solid by 22 in a way in facility for title for title for the country and the bound have a solid by 22 in a way in facility for the country and the country
²⁴ binomial test was infeasible for <i>titration-specific</i> features, as the hypothesis would be ²
²⁵ formulated as such. Given observed counts and the titration total feature abundance ²⁶
²⁶ , the true feature relative abundance. Because the count table abundance values ²¹
²⁷ for these features was 0 in the unmixed samples, their estimated probability of ²
28 occurrence π_{min} is equal to 0. As non-zero counts were observed the true feature
²⁹ proportion is non-zero, and the test always 0, and thus, the binomial test fails. ²¹
³⁰ Therefore, we formulated a Bayesian hypothesis test for $titration$ -specific features $-$
A Bayesian hypothesis test detailed by equation (2). This Bayesian approach 3
32 was used to evaluate if the true feature proportion is less than the minimum de- 32
³³ tected proportion. The Bayesian hypothesis test was formulated using equation ³³
Which Note that when assuming equal priors, $P(\pi < \pi_{min}) = P(\pi \ge \pi_{min})$,
$P(\pi < \pi_{min}) = P(\pi > \pi_{min})$ (2) reduces to (3). For equations and π is We define
$\frac{36}{\pi}$ as the true feature proportion, $\frac{\pi_{min}}{\pi_{min}}$ is π_{min} the minimum detected proportion,
${}^{37}C$ is C the expected feature counts, and C_{obs} is C_{obs} the observed feature counts.
³⁸ Simulation was used to generate possible values of C , assuming C has a binomial 3
³⁹ distribution given the observed sample total feature abundance, and a uniform ³¹
40 probability distribution for π between 0 and 1. π_{min} Count values for C were
⁴¹ simulated using a beta prior (with varying alpha and beta values) for $\pi > \pi_{min}$
⁴² and a uniform distribution for $\pi < \pi_{min}$. Higher values of alpha and beta will skew ⁴
⁴³ the prior right and left respectively. Our Bayesian hypothesis tests (Eg. (3)) results ⁴
⁴⁴ were largely unaffected by beta distribution parameterization (Fig. S4). π_{min} was ⁴
⁴⁵ calculated using the mixture equation (1) where $q_{pre,j}$ and $q_{post,j}$ are $min(\mathbf{Q}_{pre})^{4}$
⁴⁶ and $min(\mathbf{Q}_{post})$ $q_{pre,j}$ and $q_{post,j}$ are $min(\mathbf{Q}_{pre})$ and $min(\mathbf{Q}_{post})$ across all features

Olson et al. Page 27 of 31

```
<sup>1</sup>for a subject and pipeline. Our assumption is that \frac{\pi}{\pi} is less than \frac{\pi_{min}}{\pi_{min}} for \frac{\pi_{min}}{\pi_{min}}
 <sup>2</sup>features not observed in unmixed samples<del>due to random sampling.</del> Artifacts not <sup>2</sup>
 <sup>3</sup>explained by sequencing alone are likely errors in the sequence measurement and
 <sup>4</sup>inference processes, and thus, false positives or negatives.
                                                                                                          6
 <sup>7</sup> p = P(\pi < \pi_{min} | C \ge C_{obs})
    = \frac{P(C \ge C_{obs} | \pi < \pi_{min}) P(\pi < \pi_{min})}{P(C \ge C_{obs} | \pi < \pi_{exp}) P(\pi < \pi_{min}) + P(C \ge C_{obs} | \pi \ge \pi_{min}) P(\pi \ge \pi_{min})}
                                                                                                      (2)^{8}
                                                                                                          10
11
                                                                                                          11
                                                                                                          12
12
        p = \frac{P(C \ge C_{obs} | \pi < \pi_{min})}{P(C > C_{obs})}
                                                                                                      (3)_{13}
13
14
                                                                                                          14
15 Quantitative Assessment
                                                                                                          15
16For quantitative assessment, we compared observed relative abundance and log16
17 fold-changes to expected values derived from the titration experimental design. 17
18Feature average relative abundance across PCR replicates was calculated using a18
19 negative binomial model, and used as observed relative abundance values (obs) for 19
20the relative abundance assessment. Average relative abundance values were used20
21to reduce PCR replicate outliers from biasing the assessment results. Equation (1)21
<sup>22</sup>and inferred \theta values were used to calculate the expected relative abundance values<sup>22</sup>
<sup>23</sup>(exp). Relative abundance error rate is defined as |exp - obs|/exp.
<sup>24</sup> We developed bias and variance metrics to assess feature performance. The<sup>24</sup>
<sup>25</sup>feature-level bias and variance metrics were defined as the median error rate and <sup>25</sup>
<sup>26</sup>robust coefficient of variation (RCOV = IQR/median) respectively. Mixed-effects<sup>26</sup>
<sup>27</sup>models were used to compare feature-level error rate bias and variance metrics across<sup>27</sup>
<sup>28</sup>pipelines with subject as a random effect. Extreme feature-level error rate bias and <sup>28</sup>
<sup>29</sup>variance metric outliers were observed, these outliers were excluded from the mixed<sup>29</sup>
<sup>30</sup>effects model to minimize biases due to poor model fit and were characterized <sup>30</sup>
<sup>31</sup>independently.
32 Log fold-change between samples in the titration series including PRE and POST<sup>32</sup>
<sup>33</sup>were compared to the expected log fold-change values to assess differential abun-<sup>33</sup>
<sup>34</sup>dance log fold-change estimates. Log fold-change estimates were calculated using<sup>34</sup>
^{35}EdgeR [44, 45]. Expected log fold-change for feature j between titrations l and m^{35}
<sup>36</sup>is calculated using equation (4), where \theta is the proportion of POST bacterial DNA<sup>36</sup>
<sup>37</sup>in a titration, and q is feature relative abundance. For features only present in PRE<sup>37</sup>
<sup>38</sup>samples, the expected log fold-change is independent of the observed counts for the <sup>38</sup>
<sup>39</sup>unmixed samples and is calculated using (5). Features only observed in POST sam-<sup>39</sup>
<sup>40</sup>ples, POST-specific, expected log fold-change values can be calculated in a similar <sup>40</sup>
<sup>41</sup>manner. However, POST-specific features were rarely observed in more than one<sup>41</sup>
<sup>42</sup>titration and therefore were not suitable for use in our assessment. Due to a limited <sup>42</sup>
<sup>43</sup>number of PRE-specific features, both PRE-specific and PRE-dominant features<sup>43</sup>
<sup>44</sup>were used in the differential abundance assessment. PRE-specific features were de-<sup>44</sup>
<sup>45</sup>fined as features observed in all four PRE PCR replicates and not observed in any <sup>45</sup>
<sup>46</sup> of the POST PCR replicates and PRE-dominant features were also observed in all <sup>46</sup>
```

Olson et al. Page 28 of 31

¹four PRE PCR replicates and observed in one or more of the POST PCR replicates ²with a log fold-change between PRE and POST samples greater than 5. 3 4 $logFC_{lm,j} = log_2 \left(\frac{\theta_l q_{post,j} + (1 - \theta_l) q_{pre,i}}{\theta_m q_{post,j} + (1 - \theta_m) q_{pre,j}} \right)$ 5 $(4)^5$ 8 8 $logFC_{lm,i} = log_2\left(\frac{1-\theta_l}{1-\theta_m}\right)$ $(5)^9$ 10 10 Count Table Assessment Demonstration Demonstrate framework by comparing the qualitative and quantitative assessment results across the three pipelines. We first characterized overall differences in the count tables produced by the three pipelines. This characterization included calculating the number of features, total abundance by sample, dropout-rate, and 15 taxonomic composition. 17 18 ¹⁸Qualitative Assessment ¹⁹For the qualitative assessment we compare the proportion of artifactual features. The artifactual feature proportion was defined as the proportion of *unmixed*- and ²⁰ ²¹titration-specific features with abundance values that could not be explained by ²¹ 22 sampling alone. These are PCR replicates with p-values less than 0.05 after multiple 22 ²³hypothesis test correction for the binomial and bayesian hypothesis tests described²³ ²⁴in the assessment framework methods section. We additionally used the count²⁴ ²⁵table sparsity values to draw conclusions regarding the mechanism responsible for ²⁵ ²⁶different artifactual feature proportions. 27 ²⁸Quantitative Assessment ²⁹Mixed-effects models were used to compare feature-level error rate bias and variance²⁹ ³⁰metrics across pipelines with subject as a random effect. Extreme feature-level³⁰ ³¹error rate bias and variance metric outliers were excluded from this analysis to³¹ ³²minimize biases due to poor model fit. Features with large bias and variance metrics. ³² $^{33}1.5 \times IQR$ from the median, were deemed outliers. These outlier features were 33 ³⁴characterized independently in a separate analysis. We fit the following mixed effect model to test for differences in measurement bias³⁵ ³⁶across pipelines 37 37 38 38 $e_{ijk} = b + b_i + z_j + \epsilon_{ijk}$ 39 where e_{ijk} is the observed error across features and tritations k for pipeline i^{40} on individual j. b_i is a fixed term modeling the pipeline effect, z_j is a random ⁴²effect (normally distributed with mean 0) capturing overall bias differences across ⁴² ⁴³individuals. We fit a similar model for differences in error variance across pipelines. We used estimated terms \hat{b}_i from the mixed effects model to test for pair-wise 45 ⁴⁶differences across pipelines. These multiple comparisons were performed with ⁴⁶

Olson et al. Page 29 of 31

¹ Tukey's HSD test. A one-sided alternative hypothesis was used to determine which	h ¹
² pipelines had smaller feature-level error rate.	2
3	3
4 Declarations	4
5Ethics approval and consent to participate	5
6 ^{Not} applicable.	6
7	7
7 Consent for publication 8Not applicable.	8
9	9
Availability of data and material	10
¹⁰ Sequence data was deposited in the NCBI SRA archive under Bioproject PRJNA480312. Individual SRA run 11accession numbers and metadata in Supplemental Table. The code used to run the bioinformatic pipelines is	11
available at https://github.com/nate-d-olson/mgtst_pipelines. Scripts used to analyze the data are available	9
12at https://github.com/nate-d-olson/mgtst_pub.	12
13 ₁₄ Competing interests	13
The authors declare that they have no competing interests.	14
15	15
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18 Authors' contributions	18
19NDO, HCB, OCS, MS, and WT designed the experiment, SL and SH performed the laboratory work. NDO,	19
HCBand MS, MS, and DJB analyzed the data. NDO, DJB, and HCB wrote the manuscript. All authors provided feedback on manuscript drafts and approved the final manuscript.	20
21	21
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24Provided helpful insight during the development of the project. Opinions expressed in this paper are the authors and	d ₂₄
do not necessarily reflect the policies and views of NIST, or affiliated venues. Certain commercial equipment, 25instruments, or materials are identified in this paper in order to specify the experimental procedure adequately. Sucl	
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35References	35
1. Goodrich, J.K., Di Rienzi, S.C., Poole, A.C., Koren, O., Walters, W.A., Caporaso, J.G., Knight, R., Ley, R.E.: Conducting a microbiome study. Cell 158 (2), 250–262 (2014)	36
37 2. D'Amore, R., Ijaz, U.Z., Schirmer, M., Kenny, J.G., Gregory, R., Darby, A.C., Quince, C., Hall, N.: A	37
comprehensive benchmarking study of protocols and sequencing platforms for 16S rRNA community profiling. BMC Genomics 17, 1–40 (2016). doi:10.1186/s12864-015-2194-9	38
39 3. Brooks, J.P., Edwards, D.J., Harwich, M.D., Rivera, M.C., Fettweis, J.M., Serrano, M.G., Reris, R.A., Sheth,	39
N.U., Huang, B., Girerd, P., <i>et al.</i> : The truth about metagenomics: quantifying and counteracting bias in 16s rrna studies. BMC microbiology 15 (1), 66 (2015)	40
4. Westcott, S.L., Schloss, P.D.: Opticlust, an improved method for assigning amplicon-based sequence data to	41
operational taxonomic units. mSphere 2 (2) (2017) 42 5. Schloss, P.D., Westcott, S.L., Ryabin, T., Hall, J.R., Hartmann, M., Hollister, E.B., Lesniewski, R.A., Oakley,	41
B.B., Parks, D.H., Robinson, C.J., et al.: Introducing mothur: open-source, platform-independent,	
community-supported software for describing and comparing microbial communities. Applied and environmenta microbiology 75 (23), 7537–7541 (2009)	
6. Rideout, J.R., He, Y., Navas-Molina, J.A., Walters, W.A., Ursell, L.K., Gibbons, S.M., Chase, J., McDonald,	44
D., Gonzalez, A., Robbins-Pianka, A., Clemente, J.C., Gilbert, J.A., Huse, S.M., Zhou, HW., Knight, R., Caporaso, J.G.: Subsampled open-reference clustering creates consistent, comprehensive OTU definitions and	45
scales to billions of sequences. Peer J 2, 545 (2014)	46

Olson et al. Page 30 of 31

```
7. Caporaso, J.G., Kuczynski, J., Stombaugh, J., Bittinger, K., Bushman, F.D., Costello, E.K., Fierer, N., Peña,
       A.G., Goodrich, J.K., Gordon, J.I., Huttley, G.A., Kelley, S.T., Knights, D., Koenig, J.E., Ley, R.E., Lozupone, 2
       C.A., McDonald, D., Muegge, B.D., Pirrung, M., Reeder, J., Sevinsky, J.R., Turnbaugh, P.J., Walters, W.A.,
       Widmann, J., Yatsunenko, T., Zaneveld, J., Knight, R.: Qiime allows analysis of high-throughput community
       sequencing data. Nature Methods 7, 335 (2010). Correspondence
      Callahan, B.J., McMurdie, P.J., Rosen, M.J., Han, A.W., Johnson, A.J.A., Holmes, S.P.: Dada2:
                                                                                                                        5
 5
       High-resolution sample inference from illumina amplicon data. Nature Methods 13, 581-583 (2016).
       doi:10.1038/nmeth.3869
       Bokulich, N.A., Rideout, J.R., Mercurio, W.G., Shiffer, A., Wolfe, B., Maurice, C.F., Dutton, R.J., Turnbaugh,
       P.J., Knight, R., Caporaso, J.G.: mockrobiota: a public resource for microbiome bioinformatics benchmarking.
       mSystems 1(5), 00062-16 (2016)
       Kopylova, E., Navas-molina, J.A., Mercier, C., Xu, Z.: Open-Source Sequence Clustering Methods Improve the
       State Of the Art. mSystems 1(1), 1-16 (2014). doi:10.1128/mSystems.00003-15.Editor
      Huse, S.M., Welch, D.M., Morrison, H.G., Sogin, M.L.: Ironing out the wrinkles in the rare biosphere through
  11.
10
       improved OTU clustering. Environmental microbiology 12(7), 1889–98 (2010).
       doi:10.1111/j.1462-2920.2010.02193.x
       Costea, P.I., Zeller, G., Sunagawa, S., Pelletier, E., Alberti, A., Levenez, F., Tramontano, M., Driessen, M.,
       \mathsf{Hercog},\,\mathsf{R.},\,\mathsf{Jung},\,\mathsf{F.-E.},\,\mathsf{Kultima},\,\mathsf{J.R.},\,\mathsf{Hayward},\,\mathsf{M.R.},\,\mathsf{Coelho},\,\mathsf{L.P.},\,\mathsf{Allen-Vercoe},\,\mathsf{E.},\,\mathsf{Bertrand},\,\mathsf{L.},\,\mathsf{Blaut},\,\mathsf{M.},\,\mathsf{12}
       Brown, J.R.M., Carton, T., Cools-Portier, S., Daigneault, M., Derrien, M., Druesne, A., de Vos, W.M., Finlay,
13
       B.B., Flint, H.J., Guarner, F., Hattori, M., Heilig, H., Luna, R.A., van Hylckama Vlieg, J., Junick, J., Klymiuk,
       I., Langella, P., Le Chatelier, E., Mai, V., Manichanh, C., Martin, J.C., Mery, C., Morita, H., O'Toole, P.W.,
       Orvain, C., Patil, K.R., Penders, J., Persson, S., Pons, N., Popova, M., Salonen, A., Saulnier, D., Scott, K.P.,
15
       Singh, B., Slezak, K., Veiga, P., Versalovic, J., Zhao, L., Zoetendal, E.G., Ehrlich, S.D., Dore, J., Bork, P.:
       Towards standards for human fecal sample processing in metagenomic studies. Nat. Biotechnol. 35, 1069
16
                                                                                                                        16
^{17}13. Olson, N.D., Morrow, J.B.: DNA extract characterization process for microbial detection methods development ^{17}
       and validation. BMC Res. Notes 5, 668 (2012)
                                                                                                                        18
      Klindworth, A., Pruesse, E., Schweer, T., Peplies, J., Quast, C., Horn, M., Glöckner, F.O.: Evaluation of
19
       general 16s ribosomal rna gene pcr primers for classical and next-generation sequencing-based diversity studies. 19
       Nucleic acids research, 808 (2012)
      Gohl, D.M., Vangay, P., Garbe, J., MacLean, A., Hauge, A., Becker, A., Gould, T.J., Clayton, J.B., Johnson,
       T.J., Hunter, R., Knights, D., Beckman, K.B.: Systematic improvement of amplicon marker gene methods for 21
       increased accuracy in microbiome studies. Nat. Biotechnol. (2016)
<sup>22</sup>16.
                                                                                                                        22
      Pinto, A.J., Raskin, L.: PCR biases distort bacterial and archaeal community structure in pyrosequencing
       datasets, PLoS One 7(8), 43093 (2012)
                                                                                                                        23
      Hansen, M.C., Tolker-Nielsen, T., Givskov, M., Molin, S.: Biased 16S rDNA PCR amplification caused by
                                                                                                                        24
       interference from DNA flanking the template region. FEMS Microbiol. Ecol. 26(2), 141-149 (1998)
2518. McMurdie, P.J., Holmes, S.: Waste not, want not: why rarefying microbiome data is inadmissible. PLoS
                                                                                                                        25
       Comput. Biol. 10(4), 1003531 (2014)
2619
       Parsons, J., Munro, S., Pine, P.S., McDaniel, J., Mehaffey, M., Salit, M.: Using mixtures of biological samples 26
       as process controls for rna-sequencing experiments. BMC genomics 16(1), 708 (2015)
      Pine, P.S., Rosenzweig, B.A., Thompson, K.L.: An adaptable method using human mixed tissue ratiometric
       controls for benchmarking performance on gene expression microarrays in clinical laboratories. BMC
                                                                                                                        28
       biotechnology 11(1), 38 (2011)
<sup>29</sup>21.
      Thompson, K.L., Rosenzweig, B.A., Pine, P.S., Retief, J., Turpaz, Y., Afshari, C.A., Hamadeh, H.K., Damore, <sup>29</sup>
       M.A., Boedigheimer, M., Blomme, E., et al.: Use of a mixed tissue rna design for performance assessments on 30
       multiple microarray formats. Nucleic acids research 33(22), 187-187 (2005)
      Harro, C., Chakraborty, S., Feller, A., DeNearing, B., Cage, A., Ram, M., Lundgren, A., Svennerholm, A.-M.,
       Bourgeois, A.L., Walker, R.I., et al.: Refinement of a human challenge model for evaluation of enterotoxigenic 32
32
       escherichia coli vaccines. Clinical and Vaccine Immunology 18(10), 1719-1727 (2011)
3323.
       Pop, M., Paulson, J.N., Chakraborty, S., Astrovskaya, I., Lindsay, B.R., Li, S., Bravo, H.C., Harro, C., Parkhill, 33
       J., Walker, A.W., et al.: Individual-specific changes in the human gut microbiota after challenge with
       enterotoxigenic escherichia coli and subsequent ciprofloxacin treatment. BMC genomics f 17(1),\ 1\ (2016)
       Kozich, J.J., Westcott, S.L., Baxter, N.T., Highlander, S.K., Schloss, P.D.: Development of a dual-index
                                                                                                                        35
       sequencing strategy and curation pipeline for analyzing amplicon sequence data on the miseg illumina
                                                                                                                        36
       sequencing platform. Applied and environmental microbiology 79(17), 5112-5120 (2013)
       Walters, W., Hyde, E.R., Berg-Lyons, D., Ackermann, G., Humphrey, G., Parada, A., Gilbert, J.A., Jansson,
       J.K., Caporaso, J.G., Fuhrman, J.A., Apprill, A., Knight, R.: Improved bacterial 16S rRNA gene (v4 and v4-5)
38
       and fungal internal transcribed spacer marker gene primers for microbial community surveys. mSystems \mathbf{1}(1)
                                                                                                                        38
26. Edgar, R.C., Haas, B.J., Clemente, J.C., Quince, C., Knight, R.: Uchime improves sensitivity and speed of
                                                                                                                        39
       chimera detection. Bioinformatics 27(16), 2194-2200 (2011)
                                                                                                                        40
  27. Edgar, R.C.: UNCROSS2: identification of cross-talk in 16S rRNA OTU tables (2018)
<sup>41</sup>28.
                                                                                                                        41
      D'Amore, R., Ijaz, U.Z., Schirmer, M., Kenny, J.G., Gregory, R., Darby, A.C., Shakya, M., Podar, M., Quince,
       C., Hall, N.: A comprehensive benchmarking study of protocols and sequencing platforms for 16S rRNA
       community profiling. BMC Genomics 17, 55 (2016)
<sup>43</sup>29.
                                                                                                                        43
       Sze, M.A., Schloss, P.D.: The impact of dna polymerase and number of rounds of amplification in pcr on 16s
       rrna gene sequence data. bioRxiv (2019). doi:10.1101/565598
                                                                                                                        44
       https://www.biorxiv.org/content/early/2019/03/04/565598.full.pdf
      Wright, E.S., Yilmaz, L.S., Ram, S., Gasser, J.M., Harrington, G.W., Noguera, D.R.: Exploiting extension bias 45
       in polymerase chain reaction to improve primer specificity in ensembles of nearly identical dna templates
       Environmental microbiology 16(5), 1354-1365 (2014)
```

Olson et al. Page 31 of 31

¹ 31.	Callahan, B., Sankaran, K., Fukuyama, J., McMurdie, P., Holmes, S.: Bioconductor workflow for microbiome	1
2	data analysis: from raw reads to community analyses [version 2; referees: 3 approved]. F1000Research 5(1492) (2016). doi:10.12688/f1000research.8986.2	2
³ 32.	Risely, A., Waite, D., Ujvari, B., Klaassen, M., Hoye, B.: Gut microbiota of a long-distance migrant	3
4 33	demonstrates resistance against environmental microbe incursions. Molecular ecology (2017) Baker, S.C., Bauer, S.R., Bever, R.P., Brenton, J.D., Bromley, B., Burrill, J., Causton, H., Conley, M.P.,	4
5	Elespuru, R., Fero, M., et al.: The external rna controls consortium: a progress report. Nature methods 2(10),	5
6 34	731–734 (2005) R Core Team: R: A Language and Environment for Statistical Computing. R Foundation for Statistical	6
7	Computing, Vienna, Austria (2018), R Foundation for Statistical Computing, https://www.R-project.org/	7
8 ^{35.}	Yang, B., Wang, Y., Qian, PY.: Sensitivity and correlation of hypervariable regions in 16s rrna genes in phylogenetic analysis. BMC bioinformatics 17(1), 1 (2016)	8
936.	Souza, W., Carvalho, B.: Rqc: Quality Control Tool for High-Throughput Sequencing Data. (2017). R package	e 9
¹⁰ 37.	version 1.10.2. https://github.com/labbcb/Rqc Huber, W., Carey, J., V., Gentleman, R., Anders, S., Carlson, M., Carvalho, S., B., Bravo, C., H., Davis, S.,	10
11	Gatto, L., Girke, T., Gottardo, R., Hahne, F., Hansen, D., K., Irizarry, A., R., Lawrence, M., Love, I., M., MacDonald, J., Obenchain, V., Ole's, K., A., Pag'es, H., Reyes, A., Shannon, P., Smyth, K., G., Tenenbaum,	11
12	D., Waldron, L., Morgan, M.: Orchestrating high-throughput genomic analysis with Bioconductor. Nature	12
13	Methods 12(2), 115–121 (2015) Quast, C., Pruesse, E., Yilmaz, P., Gerken, J., Schweer, T., Yarza, P., Peplies, J., Glöckner, F.O.: The silva	13
14	ribosomal rna gene database project: improved data processing and web-based tools. Nucleic acids research	14
¹⁵ 39.	41(D1), 590–596 (2012) Wang, Q., Garrity, G.M., Tiedje, J.M., Cole, J.R.: Naive bayesian classifier for rapid assignment of rrna	15
16	sequences into the new bacterial taxonomy. Applied and environmental microbiology 73(16), 5261–5267 (2007)) 16
17	Aronesty, E.: ea-utils: Command-line tools for processing biological sequencing data. Expression Analysis, Durham, NC (2011)	17
18 ⁴¹ .	Edgar, R.C.: Search and clustering orders of magnitude faster than blast. Bioinformatics 26 (19), 2460–2461	18
1942.	(2010) DeSantis, T.Z., Hugenholtz, P., Larsen, N., Rojas, M., Brodie, E.L., Keller, K., Huber, T., Dalevi, D., Hu, P.,	19
20	Andersen, G.L.: Greengenes, a chimera-checked 16s rrna gene database and workbench compatible with arb. Applied and environmental microbiology 72 (7), 5069–5072 (2006)	20
2143.	Benjamini, Y., Hochberg, Y.: Controlling the false discovery rate: a practical and powerful approach to multiple	e21
2244	testing. Journal of the royal statistical society. Series B (Methodological), 289–300 (1995) Robinson, M.D., McCarthy, D.J., Smyth, G.K.: edger: a bioconductor package for differential expression	22
23	analysis of digital gene expression data. Bioinformatics ${f 26}(1),~139$ –140 (2010)	23
45. 24	McCarthy, D.J., Chen, Y., Smyth, G.K.: Differential expression analysis of multifactor RNA-Seq experiments with respect to biological variation. Nucleic Acids Res. 40 (10), 4288–4297 (2012)	24
25		25
26		26
27		27
28		28
29		29
30		30
31		31
32		32
33		33
34		34
35		35
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