NG-Tax, a highly accurate and validated pipeline for analysis of 16S rRNA amplicons from complex biomes [version 2; peer review: 2 approved, 1 approved with reservations, 1 not approved]

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Abstract

Background: Massive high-throughput sequencing of short, hypervariable segments of the 16S ribosomal RNA (rRNA) gene has transformed the methodological landscape describing microbial diversity within and across complex biomes. However, several studies have shown that the methodology rather than the biological variation is responsible for the observed sample composition and distribution. This compromises meta-analyses, although this fact is often disregarded.

Results: To facilitate true meta-analysis of microbiome studies, we developed NG-Tax, a pipeline for 16S rRNA gene amplicon sequence analysis that was validated with different mock communities and benchmarked against QIIME as a frequently used pipeline. The microbial composition of 49 independently amplified mock samples was characterized by sequencing two variable 16S rRNA gene regions, V4 and V5-V6, in three separate sequencing runs on Illumina's HiSeq2000 platform. This allowed for the evaluation of important causes of technical bias in taxonomic classification: 1) run-to-run sequencing variation, 2) PCR–error, and 3) region/primer specific amplification bias. Despite the short read length (~140 nt) and all technical biases, the average specificity of the taxonomic assignment for the phylotypes included in the mock communities was 97.78%. On average 99.95% and 88.43% of the reads could be assigned to at least family or genus level, respectively, while assignment to 'spurious genera' represented on average only 0.21% of the reads per sample.
Analysis of α- and β-diversity confirmed conclusions guided by biology rather than the aforementioned methodological aspects, which was not achieved with QIIME.

**Conclusions:** Different biological outcomes are commonly observed due to 16S rRNA region-specific performance. NG-Tax demonstrated high robustness against choice of region and other technical biases associated with 16S rRNA gene amplicon sequencing studies, diminishing their impact and providing accurate qualitative and quantitative representation of the true sample composition. This will improve comparability between studies and facilitate efforts towards standardization.

**Keywords**
16S rRNA amplicon analysis, microbial community analysis, microbial ecology, next-generation sequencing, bioinformatic pipeline
Background

Recent advances in massive high-throughput, short-amplicon sequencing are revolutionizing efforts to describe microbial diversity within and across complex biomes. Cultivation-independent whole metagenome sequencing has received increasing attention in the functional characterization of individual communities. These efforts, however, remain relatively expensive on a per sample basis, and the richer but much more unstructured information content requires complex data modelling and analysis procedures. Therefore targeted surveys for specific taxonomic marker genes, such as the 16S ribosomal RNA (rRNA) gene, remain essential in many microbial ecological studies. These surveys rely on sequencing of short, PCR amplified, hypervariable subregions rather than the full-length gene, mostly for reasons of throughput, sequence depth and cost-efficiency.

Despite great efforts to address the accuracy and reproducibility of scientific insights generated from 16S rRNA gene amplicon sequencing studies, methodology rather than biology has been shown to be the largest driver of variation in many microbiome studies. Replicability rather than reproducibility, by providing widely adopted defaults. However, there is a large distillation between the two. Drummond suggested that exact replication of an experiment (i.e., replicability) is less informative (although a necessary pre-requisite for any scientific endeavour) than the corroboration of findings by reproduction in different independent setups (i.e., reproducibility), because biological findings that are robust to independent methodologies are arguably more dependable than any single-track analysis. This distinction is highly relevant for the field of microbial ecology, where replicability is often confused with reproducibility, which is apparent from many often non-interchangeable methodologies.

Accuracy can typically be evaluated by the addition of positive controls. Generally these are synthetic or mock communities (MCs) consisting of phylotypes that, ideally, are representative of the ecosystem of interest. MCs allow researchers to answer two essential questions concerning accuracy. 1) Do I retrieve the number of species I put in, and if so are they correctly assigned? 2) How well does the PCR, sequencing and data analysis procedure reproduce species relative abundances? Reproducibility can be evaluated by comparing separate sequencing runs and different primer pairs that cover distinct 16S rRNA gene regions. Although replicability is often achieved, accuracy has been shown to be challenging especially at higher taxonomic resolution such as at genus level.

Central to all 16S rRNA gene amplicon studies are Operational Taxonomic Units (OTUs). These are often regarded as a synthetic proxy for microbial species and are typically clustered at 97% sequence similarity. However, the prokaryotic species definition remains a hotly debated topic without any satisfying solution so far. Moreover, the 97% sequence similarity threshold is based on the complete 16S rRNA gene (~1500 nt), and although sequence variability is not evenly distributed it is routinely applied to short reads of 100–500 nt. Different regions would therefore require their own species level cut-off. The combination of an ambiguous prokaryotic species definition and its application to short reads is the foundation for many complications regarding `correct’ OTU clustering. So far, there is little consensus on key experimental choices such as primers, targeted variable regions and OTU picking/clustering algorithms. Each of these technical aspects generate biases, and different methods produce clearly distinct results, leading to a situation where results of current studies cannot be easily compared or extrapolated to other study designs.

Historically, 16S rRNA gene sequences generated in a project were initially clustered de novo into OTUs at >97% sequence similarity using various clustering algorithms, mostly because available 16S rRNA gene reference databases were thought to provide insufficient coverage. Although new clustering algorithms that reduce the influence of clustering parameters, such as a hard cutoff for cluster similarity, have been specifically developed for amplicons, cluster generation is context-dependent, i.e. different datasets generate different clusters, and different algorithms may produce different end-results. Therefore, even though the same analysis framework is used, independent studies remain incomparable at OTU level. Consequently, reference-based OTU clustering has received increasing attention, due to the need for standardization, and because de-novo OTU clustering for very large datasets, such as those generated by HiSeq and MiSeq sequencers, has become computationally very intensive, unless greedy heuristics are

See referee reports
employed which suffer from the problems described above. With reference-based OTU clustering, sequences are mapped to pre-clustered reference sets of curated 16S rRNA gene sequences, provided by dedicated databases such as the Ribosomal Database Project (RDP), Greengenes and SILVA. The consequence of this approach is that the ‘quality’ of the clustering of the reference set propagates to reference-picked OTUs. Clustering has limited robustness and unbalances in databases due to over- or under-representation of certain species as well as error hotspots that are not necessarily matched to the variable regions, can potentially lead to a biased cluster formation, driven by non-biological factors. These effects have been previously ignored or underestimated in reference OTU picking protocols.

Another essential experimental choice concerns the selection of a targeted variable region of the 16S rRNA gene, because it should represent the sequence variability encountered with the full-length gene. Despite several studies comparing the performance of diverse regions, sequence lengths, sequencing platforms and taxon assignment methodologies, both within and across laboratories, there is still no complete consensus about the best variable regions of the 16S rRNA gene to assess, although some initiatives such as the Earth Microbiome Project are setting some standards that are increasingly being adopted by the field. There are several factors that can lead to the commonly observed highly region-specific differences across datasets: 1) PCR bias of varying degrees, 2) different regions are associated with different error profiles and different rates of chimera formation, and 3) the actual variation contained in the sequence is dissimilar (e.g. some regions are not variable enough to differentiate between genera, while others are), which in turn can affect clustering.

Apart from the use of a diverse range of primers and OTU picking protocols that can cause differences in results between studies and/or laboratories, sequencing error is a third important factor that defines data quality. Massive high throughput, short read length sequencing platforms have not been developed for amplicon sequencing but rather for whole genome sequencing, where sequence errors in individual reads is less important. However, in 16S rRNA gene amplicon sequencing every sequencing error could potentially lead to an incorrect OTU classification which may ultimately lead to the false discovery of a new phylotype. To avoid overestimation of microbial diversity, stringent quality filtering is therefore considered essential.

To address all of the aforementioned challenges associated with microbiota profiling, multiple standardized mock communities (MCs) were specifically designed. Those MCs were sequenced in multiple sequencing runs using a Illumina HiSeq2000 instrument (101nt paired end). Furthermore, two tandem variable 16S rRNA gene regions were sequenced in parallel (V4 and V5-V6). This led to the development of NG-Tax, a pipeline that accounts for biases associated with technical aspects associated with 16S rRNA gene amplicon sequencing. Therefore, NG-Tax will improve comparability by removing technical bias and facilitate efforts towards standardization, by focusing on reproducibility as well as accuracy. To assess the performance regarding key output parameters such as taxonomic classification, composition, richness and diversity measures we benchmarked the results obtained with NG-Tax with results obtained with QIIME, a common pipeline used for the analysis of this type of data.

**Results and discussion**

**NG-Tax layout**

NG-Tax consists of three core elements, namely barcode-primer filtering, OTU-picking and taxonomic assignment (**Figure 1**). Examples of use and details of each step of the pipeline can be found in the user manual in **Dataset 1**.

**Barcode-Primer filtering.** In a first step, paired end libraries are combined, and only read pairs with perfectly matching primers and barcodes are retained. To this end, both primers are barcoded to facilitate identification of chimeras produced during library generation after pooling of individual PCR products.

**OTU picking.** For each sample an OTU table is created with the most abundant sequences, using a minimum user defined relative abundance threshold. In this particular study we employed a threshold of 0.1% minimum relative abundance. Lowering the threshold will lead to the acceptance of low abundant OTUs, with an increased probability of these OTUs being artifacts due to sequencing and PCR errors. Abundance thresholds are commonly used to remove spurious OTUs generated by sequencing and PCR errors, but previous studies applied thresholds defined by the complete dataset, thereby ignoring sample size heterogeneity which may lead to under-representation of asymmetrically distributed OTUs.

Commonly employed quality filtering parameters based on Phred score, such as minimum average Phred score, maximum number of ambiguous positions, maximum bad run length, trimming and minimum read length after quality trimming, are not utilized in NG-Tax because quality scores from the Illumina base caller have been shown to be of limited use for the identification of actual sequence errors for 16S rRNA gene amplicon studies. Additionally, these quality scores only check for errors that occurred during sequencing, but do not account for other sources of error, such as PCR amplification, whereas quality filtering by abundance is sensitive to any source of error. Moreover, the application of global parameters (e.g. average Phred score) ignores that error is sequence-specific, and hence some sequences could be affected more than others. If a species specific amplicon is more prone to PCR or sequencing errors, the relative abundance of that particular species will be underestimated. To compensate for this potential bias, discarded reads are clustered to the OTUs with one mismatch.

Finally, all OTUs are subjected to non-reference based chimera checking according to the following principle: given three OTUs named A, B and C, C will be considered a chimera when the following conditions are satisfied: C and B 3' reads are identical, and C and B 5' reads are identical and both OTUs, A and B, are at least twice as abundant as OTU C. A complete overview of the number of sequences retained in both pipelines, *i.e.* NG-Tax
and QIIME, as well as the final number of OTUs, is provided in Dataset 1.

Taxonomic assignment. In the current version of NG-Tax, taxonomy is assigned to OTUs utilizing the USEARCH algorithm and the Silva 128 SSU Ref database, containing 1,922,223 unique full length 16S rRNA gene sequences. To ensure maximum resolution and avoid the risk of errors due to clustering-associated flaws (e.g., reference sequence error hotspots, over-representation of certain species and lack of robustness in cluster formation by clustering algorithms), we use a non-clustered database. To speed up the procedure by several orders of magnitude, 16S rRNA gene sequences from the reference database are trimmed to the amplified region using the primers as a guide. For each OTU, a taxonomic assignment is retrieved at six different identity thresholds levels (100%, 98%, 97%, 95%, 92% and 90%) and at two taxonomic levels (genus and family). The final taxonomic label is determined by the assignments that show concordance at the highest taxonomic resolution. Similar dynamic thresholds are used in rtax.

Validation Datasets
Our main objective was to develop a pipeline that accurately reproduces the composition of the synthetic MCs and also reduces the impact of experimental choices. To achieve this goal, four synthetic communities of varying complexity were created, consisting of full length 16S rRNA gene amplicons of phylogenotypes (PTs) associated with the human GI-tract (Dataset 1). This specific setup limited the likelihood of overfitting to a particular OTU composition or distribution and allowed us to assess (1) the quantification potential, (2) noise floor and (3) the effect of richness and diversity on quality filtering parameters, thus ensuring a higher fidelity with biological samples than by...
using a single MC. As a reference, to assess the quality of the taxonomic classifications, full length sequences for all PTs were obtained through Sanger sequencing. Expected MCs were created in silico by trimming the full length sequences to the sequenced region. MC1 and MC2 consisted of equimolar amounts of 17 and 55 PTs, respectively. MC3 contained 55 PTs in staggered concentrations typical for the human GI-tract, and MC4 included 50 PTs with relative abundances ranging between 0.001 and 2.49%. To account for pipetting errors, each of the four MCs was produced in triplicate. These 12 MC templates were used to sequence the MCs with different conditions that cover most of the technical bias associated with 16S rRNA gene amplicon studies reported in literature. To this end, we 1) targeted either region V4 or region V5-V6, 2) used four PCR protocols differing in the number of PCR cycles and reaction volumes 3) PCR products were analysed in three different sequencing runs and in seven different libraries, and 4) two different library preparation protocols (with and without an extra amplification of 10 cycles) were applied (Dataset 1). In addition the sequencing depth ranged from 1911 to 334613 reads per sample (Dataset 1).

**NG-Tax classification of short reads versus full length classification**

To evaluate the accuracy and reproducibility of taxonomic classification using a low information content of ~140 nt compared to a maximum information content of ~1500 nt, we compared the NG-Tax classification of all 55 reference sequences trimmed to V4 and V5-V6, with a classification of the corresponding full length reference sequences using the Silva Incremental Aligner (SINA) with SILVA taxonomy\(^40\) (Figure 2). At family level, all three classifications (i.e. full length, V4 and V5-V6)
were in complete concordance for all phylotypes. Correspondingly, the consistency at genus level was very high. Only five phylotypes for V4 that belong to the poorly classified family Enterobacteriaceae, attained higher resolution using the full length sequences. In turn, for *Intestinibacter* (PT39, V5-V6) and *Klebsiella* (PT46, V5-V6), a higher resolution was attained with short reads due to the high specificity of the hypervariable region, which can be overshadowed when using the full length sequence. Lastly, only two assignment at genus level, both Enterobacteriaceae (PT52, V4 and PT45, V5-V6) were incongruent between classification of the short and full length sequences. Overall, the V5-V6 amplicons outperformed the V4 amplicons because this region allowed for differentiation between Enterobacteriaceae and even attained a higher resolution than full length sequences for some sequences. The average taxonomic specificity (percentage of hits with an identical taxonomic label) for all reference phylotypes was 97.78% for both regions with an average of 4837 and 1688 hits for regions V4 and V5-V6, respectively. The high specificity and high number of hits at very high identity thresholds, combined with the fact that the vast majority of V4 and V5-V6 based assignments matched to each other as well as to the full length classification, testifies for the reliability and quality of the assignments.

**Observed versus expected microbial profiles**

To assess the ability to reproduce the expected composition of the MCs we benchmarked NG-Tax with QIIME, a common 16S rRNA gene amplicon analysis pipeline. Table 1 shows the comparison between NG-Tax and QIIME per region and taxonomic rank with the percentage of classified reads, the amount of spurious taxa and the total percentage of misclassified reads. The number of classified sequences without considering their accuracy is higher for NG-Tax at each taxonomic rank, with relatively small differences with QIIME. However, the number and percentage of spurious reads is considerably higher for QIIME with some regions generating an average of 18.65% incorrectly assigned reads at the genus level, compared to 0.3% for NG-Tax. Consequently, NG-Tax ensured excellent reproduction of the expected profiles (Figure 3), while the QIIME profiles suffered from high a high fraction of poorly classified and spurious OTUs (Table 1, Figure 4).

**Observed versus expected diversity**

To quantify the distances to the expected profiles, the sum of weighted differences were calculated. Given two taxonomical profiles x and y, for each taxon i, we defined the difference in abundance as $d(x,y) = (x_i - y_i)$ and a weighing factor $w_i$ as $w(x,y) = (x_i - y_i)/\text{avg}(x_i + y_i)$. The weighted difference was obtained by multiplying the difference in abundance by its weighing factor. This weighing factor is used to take the relative change as well the absolute change into account, because a 1% absolute change becomes a 200% or 20% relative change depending on whether the expected abundance is 0.5% or 5%, respectively. Distances to the expected profile were significantly lower for NG-Tax ($p<1e-4$) compared to QIIME using a two-tailed t-test (Figure 5 and Dataset 1).

One template, PT17 (*Parabacteroides*), triggered so much sequencing error in the V4 region that it was rendered undetectable although it was amplified by the primers (Supplementary Figure 1). Therefore, to test both pipelines without this sequencing anomaly, it was removed from the analysis.

Richness and diversity measures are important for understanding community complexity and dynamics. Among these measures, $\alpha$-diversity is defined as the diversity within a sample, which is often estimated based on the abundance distribution

| Family          | Genus          | V4                              |
|-----------------|----------------|---------------------------------|
| NG-Tax          | 100            | 100.00                          |
| Spurious taxa   |                 | Spurious reads (%)              |
| (%)             | 1              | 4                               |
| (%)             | 0.19           | 6.40                            |
| (%)             | 15.05          | 5.47                            |
| QIIME F & R     | 100            | 100.00                          |
| NG-Tax          | 0.19           | 6.40                            |
| Spurious taxa   | 5              | 53                              |
| (%)             | 4              | 4                               |
| (%)             | 15.05          | 5.47                            |
| Spurious reads  | 0.19           | 5.47                            |

**Table 1. Performance of NG-Tax and QIIME at different taxonomic levels for region V4 and V5-V6.** Classified reads are defined as reads mapped to a sequence for which a genus, family or order level classification is given, without considering accuracy. The percentage represents the average over all samples. Spurious taxa are taxonomic classes not included in the MCs. The percentage of spurious reads is the percentage of total reads in the misclassified classes. F: forward read, R: reverse read.
Figure 3. Observed composition of all MCs compared with the expected ones (EXP) for both regions obtained with NG-Tax.
Figure 4. Observed composition of all MCs compared with the expected ones (EXP) for both regions and each read separately obtained with QIIME.
(evenness) and number (richness) of species, whereas \( \beta \)-diversity is defined as the partitioning of diversity among communities. The ability of researchers to quantify richness and diversity hinges on an accurate assessment of the composition of these communities\(^{41} \). For microbial communities, this has been particularly challenging, as none of the existing molecular microbial ecology methods normally captures more than a small proportion of the estimated total richness in most microbial communities\(^{42} \).

For deep sequencing based approaches, filtering strategies that remove low-abundance reads make it impossible to apply richness estimation metrics such as the Chao1 index and the ACE coverage estimator, because low-abundance read counts are included in their calculations. Conversely, richness estimates based on unfiltered datasets are unlikely to be accurate, if many of the reads actually represent PCR and/or sequencing errors\(^ {16} \). In contrast to purely OTU-based methods, divergence-based methods account for the fact that not all species within a sample are equally related to each other, considering two communities to be similar if they harbour the same phylogenetic lineages, even if the species representing those lineages in each of the communities are different. Consequently, these methods are more powerful than purely OTU-based methods, because similarity in 16S rRNA gene sequence often correlates with phenotypic similarity in key features such as metabolic capabilities. An added benefit is that small errors that are likely due to unfiltered sequencing errors, are punished less severely because OTUs that are only a few nt distant from each other due to error are still closely related using divergence based indices\(^ {15} \). Therefore, these indices probably provide a better estimate of the true diversity for data generated by high throughput next generation technology sequencers.

Because the aim of NG-Tax is to enhance the biological signal as much as possible by minimizing the impact of any technical aspect, divergence-based \( \alpha \)-diversity (Phylogenetic Diversity (PD))\(^ {44} \) and \( \beta \)-diversity (Unifrac\(^ {41} \)) metrics were used to visualize the diversity within and between MCs (Figure 6). The results obtained with QIIME suffered from all of the previously described technological artifacts. The MCs clustered by primer pair instead of MC, and within each cluster the structure, \textit{i.e.} the position of MCs relative to each other, was different. More importantly, the true biological variation depicted by the expected composition was reproduced by neither primer pair (Figure 6C). Based on these results not only the Principle Coordinates Analysis (PCoA) based conclusions would have been different for both primer pairs, but also the differences in taxonomic classification could lead to significant changes in identified biomarkers, in line with what has previously been observed by He and co-workers\(^ {30} \) as well as Edgar\(^ {43} \). Here we show that replicability within a variable region was attained. The more important reproducibility, however, \textit{i.e.} the corroboration of findings by reproduction in different independent setups that use \textit{e.g.} different primers, was not. This is an important observation because biological findings should be insensitive to independent methodologies\(^ {11} \). In line with the above, also the observed \( \alpha \)-diversity (PD) was found highly inflated and the biological order was not reproduced (Figure 6D). In contrast, NG-Tax provided a clear separation of samples by MC type and their representative expected samples regardless of variable region, PCR protocol, sequencing run, library and sequencing depth. These results are remarkable, given the biases associated with each of these categories and the difference in resolution between the two regions (Figure 6A). Moreover, MC2, MC3 and MC4 were very similar, sharing the same genera and largely the same phylotypes, only differing in relative distribution (Dataset 1). Correspondingly, rarefaction curves for \( \alpha \)-diversity (Figure 6B) showed excellent reproduction of the true diversity. A perfect overlap cannot be achieved since the expected MCs do not account for sequencing or PCR errors, and these factors cannot be completely removed from real sequencing data. Results for \( \alpha \)-diversity and \( \beta \)-diversity using different metrics can be found in Dataset 1.

**Figure 5. Distances to expected taxonomical profiles.** NG-Tax results are depicted in blue and QIIME in red.
Small distances to expected MCs show the accuracy of NG-Tax, reproducibility on the other hand can be evaluated by the within MCs distances and also by the dispersion of the between MCs distances (Figure 7). Distances to the expected MCs, within MC distances and dispersion of the between MCs distances were significantly (p<1e-10) lower for NG-Tax (Dataset 1).

Conclusions

An increasing number of studies have shown that the chosen methodology rather than the natural variance is responsible for the greatest variance in microbiome studies. Some authors raised their concern when comparing data generated using different strategies, which basically suggests that true reproducibility (i.e. using different approaches and drawing the same biological conclusions) is unattainable. This is an alarming observation since studies are often used to identify biomarker organisms, associated with certain host phenotypes (often comparing a diseased state to a healthy state), yet the use of different primers might show different biomarkers. So far, neither currently available pipelines nor taxonomic classifiers have been able to efficiently reduce the noise in this type of data. Nevertheless, in properly de-noised datasets, taxonomical profiles, richness and diversity should be close to the expected values and the abundance of unassigned and poorly assigned reads should be low except when dealing with largely unexplored environments that are not sufficiently covered yet by the reference databases. At lower noise levels different variable regions should yield similar conclusions with small variations due to region

Dataset 1. Raw data of NG-Tax pipeline for analysis of 16S rRNA amplicons from complex biome

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Figure 6. PCoA using Weighted Unifrac of all sequenced and expected MCs as obtained after processing of data using NG-Tax (A) and QIIME (C). Darker colored triangles represent the expected composition while lighter colored circles represent sequenced samples. B/D. Rarefaction curves of PD for all MCs and their expected counterparts for NG-Tax (B) and QIIME (D). Dashed lines represent the expected composition while solid lines represent sequenced samples.
specific resolution, and minor changes in the experiment should still deliver the same biological conclusions. Here we presented NG-Tax, an improved pipeline for 16S rRNA gene amplicon sequencing data, which continues to be a backbone in the analysis of microbial ecosystems. Several novel steps ensure much needed improved robustness against errors associated with technical aspects of these studies, such as PCR protocols, choice of 16S rRNA gene variable region and variable rates of sequencing error\textsuperscript{5,6,12}. The commonly reported problems such as many un- or poorly classified OTUs, inflated richness and diversity, taxonomic profiles that do not match the expected ones, region dependent taxonomic classification and results being highly dependent on minor changes in the experimental setup have been tackled with NG-Tax. Despite the short read length (~140 nt) and all technical biases, the average taxonomic assignment specificity for the phylotypes included in the MCs was 97.78%. In addition, 89.43% of the reads could be assigned to a genus level and 99.95% to at least family. Spurious genera represented only 0.21% of the reads per sample. More importantly, rarefaction curves and PCoA plots confirmed improved performance of NG-Tax with respect to clustering based on biology rather than technical aspects, such as sequencing run, library or choice of 16S rRNA gene region. Therefore, NG-Tax represents a method for 16S rRNA gene amplicon analysis with improved qualitative and quantitative representation of the true sample composition. Additionally, the high robustness against technical bias associated with 16S rRNA gene amplicon studies will improve comparability between studies and facilitate efforts towards standardization.

**Methods**

**Primers**

Primer pairs 515F (5’-GTGCCAGCMCCGCGGTAA) - 806R (5’-GGACTACHVGGGTWTCTAAT) and BSF784F (5’-RGGATTAGATAACC) - 1064R (5’-CGACRRCCATGCANACACT) have been previously reported for amplification of the V4\textsuperscript{17} and V5- V6\textsuperscript{6} regions of the bacterial 16S rRNA gene, respectively. They were selected based on 1) experimental validation, 2) taxonomic coverage of the relevant ecosystem (Supplementary Figure 4) and 4) adherence to specific rules associated with the sequencing platform, such as a maximum amplicon size of <500 nt. Unless noted otherwise all primers were ordered at Biolegio (Nijmegen, Netherlands).

**Barcoding strategy**

At the time of sequencing Illumina’s Hiseq2000 allowed for multiplexing of up to 12 samples per lane using an index or barcode read provided by Illumina. To achieve optimal sample throughput and phylogenetic depth, 70 primers containing a custom designed 8nt barcode were developed to combine with the Illumina barcodes to reach a maximum throughput of 12×70 samples per lane. Each set of 70 barcoded samples are referred to as “library”. Low diversity samples, such as 16S rRNA gene amplicons, can lead to problems with base calling due to overexposure of fluorescent labels. Therefore, the set of 70 barcodes was specifically designed to possess an equal base distribution over their complete length. Additionally, to avoid differential amplification, a two-base “linker” sequence that is not complementary to any 16S rRNA sequence at the corresponding position, from a database that contains 1132 phylotypes associated with the Human GI tract\textsuperscript{46}, was inserted between the primer and barcode. The resulting set of 70 barcoded primers was checked for avoidance of secondary structure formation within or between primers (i.e., primer-dimers) or between barcodes and primers, using PrimerProspector\textsuperscript{97}.

**Mock communities**

All MCs were mixed in triplicate to account for pipetting error. These MCs ranged from 17–55 species in both equimolar and staggered compositions. One MC contained members at very
low abundances of 0.1, 0.01 and 0.001% (Dataset 1). Amplicons were generated either from cloned 16S rRNA gene amplicons, isolates available in the local culture collection of the Laboratory of Microbiology, Wageningen University, or strains ordered from DSMZ and cultured according to DSMZ recommendations, after which genomic DNA was isolated using the Genejet genomic DNA isolation kit (Thermo Fisher scientific AG, Reimach, Switzerland). A 16S rRNA gene specific PCR was performed using the universal primers 27F (5’-GTTTGAATCCTGTGCTCAAG) and reverse (806R) primers (5’-GGTACCTTGTACGACTT) to obtain full length amplicons of which size and concentration were checked on a 1% agarose gel and which were column purified and quantified with the Qubit 2.0 fluorometer, and dsDNA BR assay kit (Invitrogen, Eugene, USA). Amplicons were mixed in the MCs to obtain the specified relative abundances. High quality full length reference sequences of all MC members were obtained by Sanger sequencing at GATC Biotech AG (Constance, Germany) with sequencing primers 27F - 1492R in order to confirm their identity. The MCs were diluted 10^3-fold and subsequently used as templates in PCRs for the generation of barcoded PCR products.

Barcoded PCR

Unless noted otherwise, each sample was amplified in triplicate using Phusion hot start II high fidelity polymerase (Thermo fisher scientific AG), checked for correct size and concentration on a 1% agarose gel and subsequently combined and column-purified with the High pure PCR cleanup micro kit (Roche diagnostics, Mannheim, Germany). Forty μl PCR reactions contained 28.4 μL nucleotide free water (Promega, Madison, USA), 0.4 μL of 2 U/μl polymerase, 8 μL of 5× HF buffer, 0.8 μl of 10 μM stock solutions of each of the forward (515F) and reverse (806R) primers, 0.8 μL 10mM dNTPs (Promega) and 0.8 μL template DNA (10^1 × diluted 200 ng/μl stock). Reactions were held at 98°C for 30 s and amplification proceeding for 25 cycles at 98°C for 10 s, 50°C for 10 s, 72°C for 10 s and a final extension of 7 min at 72°C. Purified amplicons were quantified using Qubit. For primer pair BSF784F-1064R the thermal cycling conditions were identical to those described above except that the annealing temperature was 42°C. To quantify noise generated by the PCR protocol, several reactions were performed with 30 or 35 cycles and 1× 100μl reaction instead of pooling 40μl in triplicate (Dataset 1).

A composite sample for sequencing was created by combining equimolar amounts of amplicons from the individual samples, followed by gel purification and ethanol precipitation to remove any remaining contaminants. The resulting libraries were sent to GATC Biotech AG for sequencing on an Illumina Hiseq2000 instrument.

Supplementary material

Supplementary Figure 1. A) Nucleotide distribution of PT17 (Parabacteroides) for each of the four primers. Positions under the black segment are fixed and specific for PT17 preventing the inclusion of sequences belonging to a different PT. B) Percentage of 10 most abundant sequences for PT17 obtained with each of the primers. PT17 (Parabacteroides) presented a sequencing anomaly in the reverse V4 region (primer R806) (Supplementary Figure 1A). From positions 50 to 67 this region had higher error rate than the other three regions. The noise generated from this anomaly masked the biological

Sequence analysis with QIIME

We have used QIIME to benchmark NG-Tax. Illumina fastq files were de-multiplexed, quality filtered and analyzed using QIIME (v. 1.9) with closed reference OTU picking, using default settings and quality parameters as previously reported

NG-tax pipeline and user manual

The NG-tax pipeline, user manual and files and code to reproduce the presented results, are available for download at http://github.com/JavierRamiroGarcia/NG-Tax.

Abbreviations

rRNA: ribosomal RNA; MC: Mock Community; OTU: Operational Taxonomic Unit; PT: Phylotype; RDP: Ribosomal Database Project; RDPc: RDP classifier; PD: Phylogenetic Diversity; PCoA: Principle Coordinates Analysis

Data availability

Supplementary material

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signal rendering PT17 undetectable. In fact the most abundant sequence represented less than 0.45% of the total reads, while for the other three regions the most abundant sequence represented more than 80% (Supplementary Figure 1B). We decided to remove the sequences belonging to PT17 from V5-V6 samples to avoid region clustering due to the presence of PT17. Our intention in this study was to test region performance under conditions in which sequencing anomalies like the one showed in Supplementary Figure 1 are not present.

Click here to access the data.

**Supplementary Figure 2. K-means cluster prediction for QIIME results.**

The number of clusters is chosen using the “elbow criterion”. When the marginal gain of variance explained drops the line bends indicating the number of clusters.

Click here to access the data.

**Supplementary Figure 3. K-means cluster prediction for NG-Tax results.**

The number of clusters is chosen using the “elbow criterion”. When the marginal gain of variance explained drops the line bends indicating the number of clusters.

Click here to access the data.

**Supplementary Figure 4. Taxonomic coverage of primers.**

Forward (left bars) and reverse (right bars) primer coverage of the major bacterial phyla associated with the human GI tract using RDP’s probematch program with one mismatch allowed.

Click here to access the data.

**Supplementary Figure 5. Beta-diversity measures for NG-Tax results.**

Click here to access the data.

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Open Peer Review

Current Peer Review Status: ?? ? ✔ ✔

Version 2

Reviewer Report 13 March 2019

https://doi.org/10.5256/f1000research.18667.r43331

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George Watts
The University of Arizona Cancer Center and Department of Pharmacology, The University of Arizona, Tucson, AZ, USA

Bonnie Hurwitz
Department of Biosystems Engineering, The University of Arizona, Tucson, AZ, USA

The manuscript reports a new tool, NG-Tax, for analysis of 16S data which has been tested and benchmarked utilizing several mock communities. The manuscript is well written and clear. In particular, the introduction demonstrates the authors' expertise and understanding of the issues and hurdles in analyzing 16S data. The data presented depicts the performance of NG-tax as compared to QIIME using default settings for both tools. As it stands, the manuscript is ready for indexing following the relatively minor comments below. There is one caveat, however, to our recommendation for indexing, stemming from the use of a now deprecated version of QIIME (version 1.X) for benchmarking. Since the initial submission of the manuscript in July 2016, a major revision of QIIME has been released (version 2.X) and therefore a more appropriate benchmark would be to compare against the latest version of QIIME. Importantly, some of the changes made in QIIME 2.X were to address the very problem that motivated the development of NG-Tax. A similar concern was raised by a previous reviewer (J. Tremblay), who criticized the use of QIIME with default settings given that these settings are known to be sub-optimal. The issues raised by default settings in QIIME have been examined by the QIIME team and optimal settings analyzed1. Nonetheless, the authors' response to Tremblay's criticism applies to ours and therefore we don't feel it is a requirement for recommending indexing. The real test of NG-tax will be when it is utilized by disparate researchers on real datasets over time, and thus dwelling on which is the most appropriate benchmark is beyond the scope of the current paper. Minor edits required/recommended before indexing are below:

1. The authors state that RDP was replaced with SILVA SINA to classify sequences, however, the Figure 2 column heading reads SINA, while the Figure legend still lists RDP.

2. Figure 4. The figure obfuscates the point by presenting too much material. Since the figure's point is to show the poorer estimates of prevalence and mis-identifications in QIIME compared to
NG-tax, it would be easier to see this point if there were fewer barcharts presented. Since any set of charts would suffice to make the point, we would recommend that the Reverse read barcharts (QIIME) be moved to supplemental data to simplify the figure. Similarly, move the V4 data to supplemental and only present V5-V6 for both NG-Tax and QIIME. Further, perhaps discuss an exemplary disparity between expected and observed in QIIME versus NG-Tax - especially one of mis-identification. Lastly, the figure legends do not provide enough information so that the figures stand alone without the manuscript text.

3. Figure 6. It is difficult to see that triangles are “darker”. We propose you omit the word “darker” in the legend and only call attention to the “circles” and “triangles” that distinguish the samples.

4. Correct typo in the word “assess” in the following sentence in the introduction “still is no complete consensus about the best variable regions of the 16S rRNA gene to assess”.

5. In the “Barcoded PCR” methods, it is stated that 30 or 35 cycles of PCR were tested to quantify noise generated by the PCR protocol. The results are alluded to in the text as being presented in figure 6, however, it is unclear where they are shown as the figure only lists the four mock communities (MC1-4) and expected outcomes.

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Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** metagenomics, algorithms, tool development, genomics, analytics, viral ecology
We confirm that we have read this submission and believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 30 November 2018

https://doi.org/10.5256/f1000research.18667.r41304

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Fiona Fouhy
Teagasc Food Research Centre, Fermoy, Ireland

I wish to approve the manuscript for acceptance as all of my previous comments have now been addressed.

Is the work clearly and accurately presented and does it cite the current literature?  
Yes

Is the study design appropriate and is the work technically sound?  
Yes

Are sufficient details of methods and analysis provided to allow replication by others?  
Yes

If applicable, is the statistical analysis and its interpretation appropriate?  
Yes

Are all the source data underlying the results available to ensure full reproducibility?  
Yes

Are the conclusions drawn adequately supported by the results?  
Yes

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 1

Reviewer Report 02 August 2016

https://doi.org/10.5256/f1000research.9931.r15389
Fiona Fouhy
Teagasc Food Research Centre, Fermoy, Ireland

This is a novel and important piece of research. Extensive research is being conducted using next generation sequencing but researchers are becoming increasingly aware that many factors such as PCR bias, region of the 16S rRNA gene targeted etc. can impact on the results achieved. This has a negative impact on the ability to compare results across studies. This manuscript sets about to address this with their new analysis pipeline NG-Tax.

The title of the manuscript is good.

The abstract accurately summarises the research but the results section should have less methods and more results.

Figure 1 is vague and fails to show the unique aspects of how NG-Tax differs from e.g. QIIME. More details would make this figure useful.

I think greater details on the filtering and the classification used by this approach would benefit the reader. Perhaps a table showing the differences between this approach and e.g. RDP, QIIME etc. would improve the readers ability to interpret the novelty of the work.

This work was done only using HiSeq data. Do the authors feel that the approach would be equally successful on approaches e.g. Ion, MiSeq etc where longer reads are achieved? It would also be nice to test the approach with a real life data set and not a mock community and see how the results compare to those achieved using traditional analysis approaches.

Figures 3 and 4 are difficult to interpret, perhaps remake as tables.

Is the work clearly and accurately presented and does it cite the current literature?  
Yes

Is the study design appropriate and is the work technically sound?  
Yes

Are sufficient details of methods and analysis provided to allow replication by others?  
Yes

If applicable, is the statistical analysis and its interpretation appropriate?  
Yes

Are all the source data underlying the results available to ensure full reproducibility?  
Yes
Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

**Author Response 14 Nov 2018**

**Javier Ramiro-Garcia,** TI Food and Nutrition (TIFN), Wageningen, The Netherlands

This is a novel and important piece of research. Extensive research is being conducted using next generation sequencing but researchers are becoming increasingly aware that many factors such as PCR bias, region of the 16S rRNA gene targeted etc. can impact on the results achieved. This has a negative impact on the ability to compare results across studies. This manuscript sets about to address this with their new analysis pipeline NG-Tax.

The title of the manuscript is good.

Thank you.

The abstract accurately summarizes the research but the results section should have less methods and more results.

We tried to find a proper balance between results and methods but taking into account that the paper describes a tool, a description of it should be included because the pipeline is both method and results at the same time. But we tried to include those results needed to prove that NG-Tax is suitable for 16S amplicon analysis:

1) Taxonomy assignment using short reads should be comparable with the assignment using the complete 16S rRNA gene.
2) Composition profiles based on sequencing data should resemble the real composition of the biological sample.
3) α and β diversity should match the expected α and β diversity.
4) Results should be reproducible and therefore robust against biological variation (different sample compositions) and technical (PCR and sequencing settings) biases.

We consider that these requirements were met by NG-Tax and hope that they will convince readers of the actual improvements that were made, regarding robustness against methodological aspects as well as a more accurate reproduction of the MC compositions. In the new version of the manuscript we included more statistical tests to measure accuracy and reproducibility of NG-Tax.

**Figure 1 is vague and fails to show the unique aspects of how NG-Tax differs from e.g.**
QIIME. More details would make this figure useful.

As suggested by the reviewer Figure 1 now includes those unique aspects of NG-Tax.

I think greater details on the filtering and the classification used by this approach would benefit the reader. Perhaps a table showing the differences between this approach and e.g. RDP, QIIME etc. would improve the reader's ability to interpret the novelty of the work.

As suggested by the reviewer we detailed the filtering and the classification process in the user manual for those readers that want to dive into the technical details.

“This script demultiplexes the raw data into samples using the information contained in the mapping file. It also generates an OTU table per sample after removing chimeras and assigns taxonomy to the OTUs. NG-Tax is designed for short reads, 70 nucleotides is the recommended read length. Reads can be trimmed to this length by the script. Longer length can be selected by the user but comparison with 70 nucleotide analysis is advisable.

OTU picking: For each sample reads are ranked by abundance and OTUs are added to an OTU table starting from the most abundant sequence until the read abundance is lower than a percentage defined by the user (recommended is at is 0.1%). Subsequently, the discarded reads are clustered to the OTU table allowing one mismatch.

Chimera removal: OTUs are subjected to non-reference based chimera checking according to the following principle: given three OTUs named A, B and C, C will be considered a chimera when the following conditions are satisfied: C and A 5’ reads are identical, C and B 3’ reads are identical and both OTUs, A and B, are at least twice as abundant as OTU C.

Taxonomic assignment: For each OTU, usearch is used to retrieve hits for the forward and reverse reads against their respective trimmed reference database. Hits that are in common between both reads are divided in 6 identity thresholds 100, 98, 97, 95, 92, 90.

A hit belongs to a certain level, for example 97, when both reads have at least a 97 percentage identity with that hit.

Using the highest available identity threshold, NG-Tax assigns the consensus taxonomy to the OTU if this taxonomy is supported for at least half of the hits. Genus, Family or Order remains unassigned if the maximum identity percentage level is lower or equal to 97%, 95% and 92% respectively.”

Now we also included the main characteristics of NG-Tax in Figure 1. The OTU generation is the main difference between NG-Tax and QIIME. NG-Tax uses a de novo generation approach without clustering. This increases the resolution and allows for the distinction between OTUs with one nucleotide distance. In addition, NG-Tax generates OTUs independently for each sample, which avoids problems associated to sample size heterogeneity. Another important feature of NG-Tax is the use of non-fixed thresholds for the taxonomic assignment, which results in more accurate classifications. To highlight those points we added the text “No clustering → max resolution” to Figure 1 and indicated in the
workflow that OTUs are generated per sample. We also clearly state that NG-Tax does not use any clustering and explain that the taxonomic classification uses different identity levels. The authors of the RDP classifier stated that it does not perform well for short sequences, i.e. a length of 200 nt would give accurate family level classification but shorter reads will not, likely due to insufficient features \(^1\). In contrast, NG-Tax has been specifically designed for short reads.

This work was done only using HiSeq data. Do the authors feel that the approach would be equally successful on approaches e.g. Ion, MiSeq etc where longer reads are achieved? It would also be nice to test the approach with a real life data set and not a mock community and see how the results compare to those achieved using traditional analysis approaches.

NG-Tax has been the reference method for 16S rRNA gene amplicon analysis in our lab for more than two years now, and has been used in more than 30 manuscripts that have been submitted or are in preparation. One of these manuscripts \(^1\) was published before this manuscript. Since then another fifteen studies using NG-Tax have been published \(^2-16\). These studies contain biological samples that belong to very different and specific environments and were sequenced both on MiSeq and HiSeq instruments. These will contribute to the assessment of NG-Tax's performance, however these were not included in the current manuscript since the data is accessible in the aforementioned publications.

Figures 3 and 4 are difficult to interpret, perhaps remake as tables.

With Figure 3 and 4 we intended to visualize how close the observed sample composition resembled the expected composition at a glance and how many different taxa are found in the data. In order to further improve interpretation, we have now also added Table 1 to provide detailed information as to the number of misclassified reads at different taxonomic levels and Figure 5, which shows boxplots of the distances to the expected composition. We also performed statistical tests to quantitatively compare the performance of NG-Tax and QIIME. As suggested by the reviewer we included the tables with the taxonomical profiles as supplementary data, which can be used for evaluation of the results.

**Competing Interests:** No competing interests were disclosed.

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Reviewer Report 02 August 2016

https://doi.org/10.5256/f1000research.9931.r15175

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Julien Tremblay
Biomonitoring, National Research Council Canada, Montreal, QC, Canada

This paper describes a pipeline for processing 16S rRNA amplicon data. They implemented an experimental design in which they used data coming from three different HiSeq2000 runs using two variable regions (V4 and V5-V6). It is however not clear if their data has been generated in-house or if their data was actually coming from public databases. This should be explicitly stated somewhere (unless I missed it). Using this data as input, the authors developed a pipeline labeled NG-Tax, which according to them: 1) better accounts (compared to what?) for errors associated with a range of technical aspects of 16S rRNA amplicon sequencing and 2) improves comparability by removing technical bias and facilitating efforts towards standardization. In my view, the problem is that why their pipeline does 1) and 2) is not addressed in depth. The description of the technical aspects of their pipeline in the first part of the result section only very summarily describes the general workflow of the pipeline, but nowhere do they describe how exactly OTU picking is done (see comment below). How exactly Chimeras are detected? With an open-source package? In-house script? Taxonomic assignment methodology is unclear as well. The authors state that they are using uclust for taxonomic assignment, while uclust is a sequence clustering software (also see comments below).

Then the authors compares their pipeline results with the ones generate by Qiime with default parameters. Qiime with its default parameters is already known to not perform optimally (See UPARSE paper, Edgar, 2013). I think that comparing with Qiime for validation is okay, but do not spend too much time dissecting the results. What the authors should focus on is, I think, on improving substantially on the technical description of their pipeline -- describe each step in details. If open source packages are being used, say so, if not, describe your script/software/algorithm. Also please make the source code available under a code repository (Github or Bitbucket for instance).

In my view the paper is not acceptable in its current form.

Specific comments:
- At the sentence "mostly because available 16S rRNA gene reference databases were thought to provide insufficient coverage13–16." Can you please elaborate on that? What do exactly mean by that?

- "there still is no standard or consensus of best choices for variable regions."

I don't fully agree with this. Depending on your field of study, a certain consensus can usually be found. For instance, the Earth Microbiome project recommends two primer sets (V4 and the 'newer' V4-V5) - Many labs investigating soil or environmental samples in general will effectively favor these primers because they are being used by a large part of the community which readily enables inter-lab community/study comparisons.

- Concerning the OTU picking section: It is not clear how exactly you pick your OTUs. Basically, you are kind of dereplicating/clustering your raw reads data set at 100% ID and then create a one column OTU table for each sample? Please clarify.

- "Phred score, such as minimum average Phred score, maximum number of ambiguous positions, maximum bad run length, trimming and minimum read length after quality
trimming, are not utilized in NG-Tax because quality scores from the Illumina base caller have been shown to be of limited use for the identification of actual sequence errors for 16S rRNA gene amplicon studies.9,37.

Yes Q scores have their limitation, but it is unwise to not filter for reads containing Ns and reads of very poor Q scores. Some basic filtering should be implemented to at least filter for very bad data. For instance if you have a read with 10 bases with Q score lower than 10, this read should obviously be removed.

“To speed up the procedure by several orders of magnitude, 16S rRNA gene sequences from the reference database are trimmed to contain only the region amplified by the primers.”

Please specify how you generated your trimmed database of 16S rRNA genes ref. In silico PCR? A multiple alignment that was trimmed at specific coordinates?

“In the current version of NG-Tax, taxonomy is assigned to OTUs utilizing the uclust algorithm16 and the Silva_111_SSU Ref database, containing 731,863 unique full length 16S rRNA gene sequences. To ensure maximum resolution and avoid the risk of errors due to clustering-associated flaws (e.g. reference sequence error hotspots, overrepresentation of certain species and lack of robustness in cluster formation by clustering algorithms) we use the non-clustered database. To speed up the procedure by several orders of magnitude”,

Uclust is for clustering sequences/reads and not for taxonomic assignment...? Taxonomic assignment is done by other means (RDP classifier), but certainly not with uclust.

For each OTU, a taxonomic assignment is retrieved at six different identity thresholds levels (100%, 98%, 97%, 95%, 92% and 90%) and at two taxonomic levels (genus and family).

How exactly are OTUs classified? With an in-house method? The RDP classifier? Please elaborate.

Figure 1. Please add more details. Are you using open-source packages in your pipeline? If so please indicate.

Table 1: Table 1 is heavy and not really meaningful. Would fit in more appropriately in suppl. material.

Figure 3 and 4: Please find another way of displaying data of figure 3. It is simply not feasible to associate a color to a given bar graph. Maybe consider using a heatmap with hierarchical clustering or a PCA/PCoA? Typically for taxonomy stacked barplots you can't really go above 20 different colors. After that it becomes indistinguishable.

“Because the focus of NG-Tax is to retain as much biological signal as possible while minimizing the impact of any technical choice,”

But how exactly does NG-Tax retain more biological signal than other pipelines, what does that mean?
Discussion: The authors say that their pipeline outperforms Qiime, but nowhere is discussed how exactly does Qiime works. How exactly does Qiime generate OTUs, how are the reads QCed? How is the classification performed, what training sets are being used for classification? It is already known that Qiime does not perform well with default parameters (see R. Edgar’s UPARSE paper), so Qiime does not represent a gold standard, especially with default parameters.

NG-Tax pipeline availability. Please include the pipeline on a Github or bitbucket repository.

References
1. Edgar RC: UPARSE: highly accurate OTU sequences from microbial amplicon reads. *Nat Methods*. 2013; 10 (10): 996-8 PubMed Abstract | Publisher Full Text

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**Is the work clearly and accurately presented and does it cite the current literature?**
Yes

**Is the study design appropriate and is the work technically sound?**
Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**
Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**
Yes

**Are all the source data underlying the results available to ensure full reproducibility?**
Yes

**Are the conclusions drawn adequately supported by the results?**
Yes

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to state that I do not consider it to be of an acceptable scientific standard, for reasons outlined above.

Author Response 14 Nov 2018

**Javier Ramiro-Garcia**, TI Food and Nutrition (TIFN), Wageningen, The Netherlands

This paper describes a pipeline for processing 16S rRNA amplicon data. They implemented an experimental design in which they used data coming from three different HiSeq2000 runs using two variable regions (V4 and V5-V6). It is however not
clear if their data has been generated in-house or if their data was actually coming from public databases. This should be explicitly stated somewhere (unless I missed it).

To further clarify this we added the section Datasets:

Datasets:
Four synthetic communities of varying complexity were created, consisting of 16S rRNA gene amplicons of phylotypes (PTs) associated with the human GI-tract (Dataset 1). This specific setup limited the likelihood of overfitting to a particular OTU composition or distribution and allowed us to assess (1) the quantification potential, (2) noise floor and (3) the effect of richness and diversity on quality filtering parameters, thus ensuring a higher fidelity with biological samples than by using a single MC. As a reference, to assess the quality of the taxonomic classifications, full length sequences for all PTs were obtained through Sanger sequencing. Expected MCs were created by trimming the full length sequences to the sequenced region. MC1 and MC2 consisted of equimolar amounts of 17 and 55 PTs, respectively. MC3 contained 55 PTs in staggered concentrations typical for the human GI-tract, and MC4 included 50 PTs with relative abundances ranging between 0.001 and 2.49%. To account for pipetting errors, each of the four MCs was produced in triplicate. To design a pipeline that puts more focus on biology, these 12 MC templates were used to sequence the MCs with different conditions that cover most of the technical bias associated with 16S rRNA gene amplicon studies reported in literature. To this end, we:

1. Targeted either region V4 or region V5-V6,
2. Used four PCR protocols differing in the number of PCR cycles and reaction volumes.
3. PCR products were analysed in three different sequencing runs and in seven different libraries.
4. Two different library preparation protocols (with and without an additional amplification of 8 cycles) were applied (Dataset 1).

In addition the sequencing depth ranged from 2363 to 335822 reads per sample (Dataset 1). One phylotype, PT17 (*Parabacteroides*), attracted so much sequencing error in the V4 region that it was rendered undetectable although it was amplified by the primers (Supplementary Figure 1). Therefore, to test both pipelines without this sequencing anomaly, it was removed from the analysis.

In this section we explain how we created and sequenced the MCs. The sequencing data was generated by a sequencing company (GATC, Constance, Germany; see section Materials and Methods). The sequencing data has been submitted to the ENA repository, and we added the following sequence data availability section:

Sequence data availability:
Sequence data have been deposited in the European Nucleotide Archive[^46], accession number [ENA:PRJEB11702](http://www.ebi.ac.uk/ena/data/view/PRJEB11702) (amplicon sequencing data for all 49 samples) and [ENA:LN907729-LN907783](http://www.ebi.ac.uk/ena/data/view/PRJEB11702) (full length 16S rRNA gene sequences for all 55 Pts).

**Using this data as input, the authors developed a pipeline labeled NG-Tax, which according to them: 1) better accounts (compared to what?) for errors associated with**

[^46]: European Nucleotide Archive
a range of technical aspects of 16S rRNA amplicon sequencing and 2) improves comparability by removing technical bias and facilitating efforts towards standardization. In my view, the problem is that why their pipeline does 1) and 2) is not addressed in depth.

We agree with the reviewer that the highlighted elements were not sufficiently clear and lacked an explanation why we believe that NG-Tax performs better. Therefore we replaced this sentence with:

“This allowed for the development of NG-Tax, a pipeline that accounts for biases associated with this range of technical aspects associated with 16S rRNA gene amplicon sequencing. Therefore NG-Tax will improve comparability by removing technical bias and facilitate efforts towards standardization, by focusing on reproducibility as well as accuracy. To assess the performance regarding key output parameters such as taxonomic classification, composition and richness, and $\alpha$ and $\beta$ diversity measures, we benchmarked the results obtained with NG-Tax.”

In order to account for errors and increase comparability by removing technical bias from 16S rRNA amplicon studies, NG-Tax should fulfill the following requirements:

1. Taxonomy assignment using short reads should be comparable with the assignment using the complete 16S rRNA gene.
2. Composition profiles based on sequencing data should resemble the real composition of the biological sample.
3. $\alpha$ and $\beta$ diversity should match the expected $\alpha$ and $\beta$ diversity.
4. Results should be reproducible and therefore robust against biological variation (different sample compositions) and technical (PCR and sequencing settings) biases.

We consider that these requirements were met by NG-Tax, as supported by the following data.

Figure 2 shows the high similarity of the taxonomic classification of the V4 and V5V6 amplicon results compared to full length sequences using SILVA Incremental Aligner (SINA). The specificity and the number of hits testify to the reliability of the assignments.

Table 1 shows the low number and percentage of spurious reads.

Figure 3 shows that NG-Tax derived compositional profiles based on sequencing data accurately resemble the expected profiles.

Figure 5 quantifies the distances to the expected profiles.

Figure 6 & 7: the PCoA plots show that MCs group by type, despite all technical bias associated with 16S rRNA gene amplicons sequencing, such as PCR settings, and primer or region selection. Figure 7 shows that all within-MC pairwise comparisons and the dispersion of all pairwise comparisons are significantly smaller in NG-Tax meaning that distances within and between MC types are robust. These results could not have been achieved without a proper reduction of the aforementioned biases. This will improve comparability.
by enabling direct comparison between studies even when using slightly different approaches.

The description of the technical aspects of their pipeline in the first part of the result section only very summarily describes the general workflow of the pipeline, but nowhere do they describe how exactly OTU picking is done (see comment below). How exactly Chimera are detected? With an open-source package? In-house script? Taxonomic assignment methodology is unclear as well. The authors state that they are using uclust for taxonomic assignment, while uclust is a sequence clustering software (also see comments below).

In the revised manuscript we substantially increased the amount and detail of information on the description of the general workflow. All the critical steps, including barcode & primer filtering, OTU picking, mapping rejected reads to accepted OTUs, de novo chimera filtering, taxonomic assignment and the generation of a phylogenetic tree are now detailed in Figure 1 and explained in the user manual.

Then the authors compares their pipeline results with the ones generate by Qiime with default paramters. Qiime with its default parameters is already known to not perform optimally (See UPARSE paper, Edgar, 2013). I think that comparing with Qiime for validation is okay, but do not spend too much time dissecting the results.

We agree with the reviewer's view on the default parameters of QIIME, however, the major improvements are gained by not clustering and processing the reads per sample. Therefore, the presented results cannot be achieved with QIIME independent of the parameters we choose. Besides that, testing QIIME under different settings has been already extensively covered elsewhere and if we would change parameters, reviewers could argue that our chosen parameters are less than optimal and therefore we stayed with the default settings.

Nonetheless, as suggested by the reviewer we reproduced the QIIME analysis with a 0.1% abundance threshold, and this is now included in the supplementary data. The results using 0.1% or 0.005% are consistent and show no performance gain (“Supplementary data. QIIME beta-div results all settings”). This is also in line with the result shown by Bokulich et al 2013, supplementary material 2; pages 8, 9 and 10. This text includes a comparison of the expected composition against real sample composition using different filtering parameters. One of these parameters is OTU abundance and the plot shows that the obtained profiles do not change much using different filtering abundance thresholds.

Although we agree with the reviewer that we need not to put too much emphasis on the QIIME results, they do show the consequences when technical bias is not adequately taken care of, which makes it easier for the non-technical reader to place the results achieved by NG-Tax into context.

What the authors should focus on is, I think, on improving substantially on the technical description of their pipeline – describe each step in details. If open source packages are being used, say so, if not, describe your script/software/algorithm. Also
please make the source code available under a code repository (Github or Bitbucket for instance).

In my view the paper is not acceptable in its current form.

We thanks the reviewer for the constructive suggestions and hope that the changes introduced in the manuscript and supplementary data help to change his opinion. As suggested by the reviewer the source code is now available on Github (https://github.com/JavierRamiroGarcia/NG-Tax.git).

Specific comments:
At the sentence "mostly because available 16S rRNA gene reference databases were thought to provide insufficient coverage 13–16." Can you please elaborate on that? What do exactly mean by that?

In the past, when pyrosequencing was the standard sequencing method, open reference base OTU picking was the common strategy for two main reasons: 1) reference databases were still very small and thought to provide insufficient coverage and 2) the more computationally intense open reference methods were used because the amount of reads generated were lower than nowadays. Over time databases were more complete and the amount of data generated increased the needed computational time, so close reference OTU picking gained popularity. Currently, with new bioinformatics solutions, open reference OTU picking is gaining ground and NG-Tax is following that trend by implementing a new open reference OTU picking algorithm.

"there still is no standard or consensus of best choices for variable regions."

I don't fully agree with this. Depending on your field of study, a certain consensus can usually be found. For instance, the Earth Microbiome project recommends two primer sets (V4 and the 'newer' V4-V5) - Many labs investigating soil or environmental samples in general will effectively favor these primers because they are being used by a large part of the community which readily enables inter-lab community/study comparisons.

We agree with the reviewer that there is a certain consensus for some projects, but still there are many publications addressing the differences in results when different primers are used. Therefore, when choosing a primer pair, whether these primers are used by the community becomes an important factor if afterwards the researcher wants to compare the results with existing studies. The idea of NG-Tax is to decrease the importance of this factor by providing comparable results across different primer sets, giving more freedom to the researcher to explore new possibilities. But as suggested by the reviewer we softened our statement and rephrased as:

“There still is no complete consensus regarding best choices for variable regions even if some initiatives like the Earth Microbiome Project are setting standards that are increasingly being adopted by the field.”
Concerning the OTU picking section: It is not clear how exactly you pick your OTUs. Basically, you are kind of dereplicating/clustering your raw reads data set at 100% ID and then create a one column OTU table for each sample? Please clarify.

We tried to make the paper as readable as possible by not adding too much technical information. Realizing that we have excessively reduced technical detail in the original manuscript, in the new version all technical details can be found in the user manual. An indication that this information can be found in the user manual is now included in the manuscript.

In NG-Tax OTUs are generated per sample using the following strategy:
For each sample reads are ranked by abundance and OTUs are added to an OTU table starting from the most abundant sequence until the read abundance is lower than a percentage defined by the user (recommended is at is 0.1%). Subsequently, the discarded reads are clustered to the OTU table allowing one mismatch.
In practical terms this is in fact guided clustering where seeds are determined by abundance. The difference with an normal clustering approach is that there is no clustering to define the seeds. This allows seeds that differ as little as one nucleotide. The clustering is applied only afterwards to compensate for potential bias due to PCR and sequencing errors. Error is sequence-specific, and hence some sequences could be affected more than others. If a species specific amplicon is more prone to PCR or sequencing errors, the relative abundance of that particular OTU will be underestimated. But after clustering, OTUs more prone to error receive a higher percentage of discarded reads than others, this differential recovery helps to reestablish the true composition that was lost due to sequence specific error rates.

“Phred score, such as minimum average Phred score, maximum number of ambiguous positions, maximum bad run length, trimming and minimum read length after quality trimming, are not utilized in NG-Tax because quality scores from the Illumina base caller have been shown to be of limited use for the identification of actual sequence errors for 16S rRNA gene amplicon studies9,37.”

Yes Q scores have their limitation, but it is unwise to not filter for reads containing Ns and reads of very poor Q scores. Some basic filtering should be implemented to at least filter for very bad data. For instance if you have a read with 10 bases with Q score lower than 10, this read should obviously be removed.

We fully agree with the reviewer. A filtering process is needed, and this is already implemented in NG-Tax. The point is that it is not based in quality score but based on abundance. Illumina have reported that 95%-97% of the reads have Q>30 (http://www.illumina.com/documents/products/technotes/technote_Q-Scores.pdf). This 3 to 5 percent of reads with lower quality will contain reads for all the different phylotypes, and within phylotypes there will be reads with errors in different positions and with different base substitutions. This decreases the probability of having exactly the same erroneous read. Therefore we expect that any specific erroneous read should be in low abundance. Subsequently, when samples are filtered by discarding low abundance sequences, those low quality reads will be removed without the need to check for quality scores.
In addition quality scores do not account for PCR errors since the base caller will give them very high scores, because according to the sequencer they are real sequences. In contrast, filtering by abundance is insensitive to the error source, and hence if the reads with PCR errors are in low abundance (especially if high fidelity taq polymerase is used), they will be also removed.

A good example of how stringent quality thresholds can bias the results can be found in Bokulich et al 2013, supplementary material 2; pages 8, 9 and 10.

“To speed up the procedure by several orders of magnitude, 16S rRNA gene sequences from the reference database are trimmed to contain only the region amplified by the primers.”

Please specify how you generated your trimmed database of 16S rRNA genes ref. In silico PCR? A multiple alignment that was trimmed at specific coordinates?

We thank the reviewer for the suggestion. This information is important and now it has been added to the user manual. NG-Tax applies an in silico PCR using the primers and a reference database given by the user. Degenerated primer positions are allowed and alternative primers with mismatches can be supplied.

"In the current version of NG-Tax, taxonomy is assigned to OTUs utilizing the uclust algorithm16 and the Silva_111_SSU Ref database, containing 731,863 unique full length 16S rRNA gene sequences. To ensure maximum resolution and avoid the risk of errors due to clustering-associated flaws (e.g. reference sequence error hotspots, overrepresentation of certain species and lack of robustness in cluster formation by clustering algorithms), we use the non-clustered database. To speed up the procedure by several orders of magnitude”,

Uclust is for clustering sequences/reads and not for taxonomic assignment...?

Taxonomic assignment is done by other means (RDP classifier), but certainly not with uclust.

For each OTU, a taxonomic assignment is retrieved at six different identity thresholds levels (100%, 98%, 97%, 95%, 92% and 90%) and at two taxonomic levels (genus and family).

How exactly are OTUs classified? With an in-house method? The RDP classifier? Please elaborate.

First we wanted to inform that uclust has been substituted by usearch in the scripts for the second version of the manuscript.

Any or at least most methods for taxonomic assignment contain two main steps. First, the read to be classified is linked to sequences in a reference database by sequence similarity, and then the taxonomic information of linked sequences, termed hits, is transferred to the sequence to be classified. Different methods can be used to perform the linking step. In our case we used usearch (previously uclust). We used dynamic thresholds to get hits at 6 different identity levels, after which the taxonomic information is transferred to the read of
unknown taxonomy by the NG-Tax classifier algorithm. Similar dynamic thresholds are used by rtax 2.

A description of how the NG-Tax classifier works:

For each OTU, usearch is used to retrieve hits for the forward and reverse reads against their respective trimmed reference database. Hits that are in common between both reads are divided in 6 identity thresholds 100, 98, 97, 95, 92, 90. A hit belongs to a certain level, for example 97, when both reads have at least a 97 percentage identity with that hit. Using the highest available identity threshold, NG-Tax assigns the consensus taxonomy to the OTU if this taxonomy is supported for at least half of the hits. Genus, Family or Order remains unassigned if the maximum identity percentage level is lower or equal to 97%, 95% and 92% respectively. The levels lower than 97% are only useful for unexplored environments; otherwise most of the OTUs are assigned at 100% identity.

As suggested by the reviewer we have included a detailed explanation of how the algorithm works in the user manual provided in the supplementary files.

Figure 1. Please add more details. Are you using open-source packages in your pipeline? If so please indicate.

We thank the reviewer for the suggestion, now we added a figure with more details. As stated in the user manual we use USEARCH and QIIME.

Table 1: Table 1 is heavy and not really meaningful. Would fit in more appropriately in suppl. material.

As suggested by the reviewer the table 1 is now supplied as supplementary material.

Figure 3 and 4: Please find another way of displaying data of Figure 3. It is simply not feasible to associate a color to a given bar graph. Maybe consider using a heatmap with hierarchical clustering or a PCA/PCoA? Typically for taxonomy stacked barplots you can’t really go above 20 different colors. After that it becomes indistinguishable.

With figure 3 and 4 we just wanted to show in one glance, how close the sample compositions resembled the expected composition and how many different taxa are found in the data. In the new version of the manuscript we have added boxplots showing distances to the expected profiles to improve interpretation. An excel file with taxonomic profiles is also added to the supplementary material for further interpretation. PCoA plots showing distances between samples and expected for both pipelines are provided in figure 6. Figure 7 shows those distances as pairwise comparisons.

"Because the focus of NG-Tax is to retain as much biological signal as possible while minimizing the impact of any technical choice,'"

But how exactly does NG-Tax retain more biological signal than other pipelines, what does that mean?
We agree with the reviewer that the sentence is confusing. Therefore it has been rephrased.

“Therefore, these indices probably provide a better estimate of the true diversity for data generated by high throughput next generation technology sequencers.

Because the aim of NG-Tax is to enhance the biological signal as much as possible by minimizing the impact of any technical choice, divergence-based α-diversity (Phylogenetic Diversity (PD) [41]) and β-diversity (Unifrac [39]) metrics were used to visualize the diversity within and between MCs (Figure 6).”

**Discussion:** The authors say that their pipeline outperforms Qiime, but nowhere is discussed how exactly does Qiime works. How exactly does Qiime generate OTUs, how are the reads QCed? How is the classification performed, what training sets are being used for classification? It is already known that Qiime does not perform well with default parameters (see R. Edgar’s UPARSE paper), so Qiime does not represent a gold standard, especially with default parameters.

In our manuscript we applied recommended settings like those described in the Bokulich paper. This paper extensively describes QIIME, the rationale behind the recommendations and the way that these choices impact the data. The scope of this manuscript was not to test QIIME under different settings. For NG-Tax analysis we also employed default and recommended settings so we thought that even if it is not optimal and has limitations, this could be a fair approach.

We also analyzed the MCs with QIIME to show that this dataset is not an exceptional case with regards to the commonly reported problems (such as many un- or poorly classified OTUs, inflated richness and diversity, taxonomic profiles that do not match the expected ones, region dependent taxonomic classification and results which are highly dependent on minor changes in the experimental procedures) are also found in this dataset. So in our mind the QIIME analysis should primarily be seen as a performance comparison. In fact we encourage researchers to use more than one method, as this will increase the amount of information they can obtain from their datasets and determine the quality of their data. This will benefit their research and by extension the whole field.

Nevertheless in an effort to increase comparability we also performed an additional analysis using QIIME with a 0.1% abundance threshold (which is conservative compared to the advised setting of 0.005%). Nevertheless this did still not reproduce the biological signal and the results obtained with 0.1% or 0.005% are consistent. These analyses have been added to the supplementary material as “Supplementary data. QIIME beta-div results all settings”.

**NG-Tax pipeline availability. Please include the pipeline on a Github or bitbucket repository.**

NG-Tax scripts were previously available as supplementary material, and as suggested by the reviewer they are now also available in Github ([https://github.com/JavierRamiroGarcia/NG-Tax.git](https://github.com/JavierRamiroGarcia/NG-Tax.git))
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**Competing Interests:** No competing interests were disclosed.

Reviewer Report 02 August 2016

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Thomas S. B. Schmidt
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In their manuscript, the authors introduce NG-Tax, an open-source software for the (meta-)analysis of 16S rRNA-based microbiome datasets. Their tool focuses on an important and so-far arguably understudied aspect of microbial ecology research: the integration of results across studies, in view of both technical and biological variation.

The approach is interesting and addresses important points. In particular, several sequencing datasets of different mock communities were generated, even using different primer sets: this is great data to benchmark on, and many (most) other papers introducing tools do not provide benchmarks on such an array of real (mock) data. In general, I feel that this is very interesting work and that NG-Tax can be a promising alternative to existing tools in the field.

However, there are several points that I feel would need to be addressed in order for the manuscript to stand tall, and for the reader to get a good understanding of how NG-Tax can be useful in practice.

**Major comments:**
- Even after reading the manuscript and online user manual repeatedly, I have to admit that
it is not completely clear to me how NG-Tax works in detail, and in which points exactly it differs from existing approaches. Based on the introduction, I gather that NG-Tax relies on closed-reference OTU picking, but this is not mentioned explicitly anywhere in the text. Also, does reference-based OTU picking in NG-Tax rely on uclust? If yes, which version and parameters were used, and how do they differ from QIIME's defaults? Also, the Background and Discussion sections do not elaborate on the various disadvantages of closed-reference approaches; most importantly, closed-ref only takes into account sequences matching the database and removes everything else. When integrating sequence data from different primer sets, this is arguably the most straightforward approach; however, the limitations should be discussed.

- I gather from the text that NG-Tax's main innovations are the use of primer-tailored reference databases and a different (more conservative) read abundance filtering scheme. It is perfectly valid to benchmark these against QIIME's default settings; however, it would be great to see how QIIME performs with similarly conservative settings, to better understand where NG-Tax's edge in performance comes from.

- Regarding taxonomy assignments, it is valid to compare NG-Tax's uclust-based approach to QIIME's uclust-based approach. However, I believe that the gold standard continues to be the RDP Classifier, and it would be interesting to see a performance comparison to this tool (on the short-read data, not only on full-length reads). Also, how does taxonomic classification by NG-Tax differ conceptually from RTAX (http://www.uio.no/english/services/it/research/hpc/abel/help/software/rtax.html)? I do believe that they are not equivalent, but the approaches appear somewhat related.

- In general, the results on taxonomic classification are not discussed quantitatively. From Figures 3&4, the visual impression is that NG-Tax indeed better approximates expected taxonomic profiles than QIIME, but it is hard to quantify this from stacked bar charts. I would suggest to compute e.g. Euclidean or more sophisticated distances of classified taxonomic profiles to the expected distribution. Also, it would be interesting to see quantitative sensitivities and specificities (or F1-scores?) on the taxonomic assignments; particularly also when running on the exact same (more conservatively filtered) dataset for QIIME. Some numbers on specificity are provided in the Abstract and Conclusion sections – but I am not sure if specificity may be gained at the expense of sensitivity based on the more rigid read filtering upstream.

- As a suggestion, but certainly not as a request, I would recommend to maybe include additional, independent datasets to benchmark on. For example, Tremblay et al. (2015) have published data on mock communities sequenced with different primer sets and on different platforms. Such data could contribute to a yet more general assessment of NG-Tax performance.

Minor comments (chronologically, not in order of importance):
- Background, “The consequence of this approach is that the ‘quality’ of the clustering of the reference set propagates to reference-picked OTUs.” I believe that as such, this statement is not fully valid or supported. In fact, the negative complement is arguably true: reference-based OTU picking against a “bad” reference can never provide “good” OTUs (a garbage-in, garbage-out problem, so to say). However, even with a good reference, a bad mapping
algorithm can generate non-informative reference-based OTU sets. Schloss & Westcott have recently published a study which discusses this point, among others (Westcott & Schloss, 2015).

- **Background,** “However, in 16S rRNA gene amplicon sequencing every sequencing error could potentially lead to the false discovery of a new species.” I have two comments on this statement. First, I believe that the term “species” in this context can be misleading and I feel that the neutral term OTU or diversity unit would be more appropriate. Second, there is a large body of literature on how sequencing errors affect 16S-based diversity studies beyond the cited Bokulich et al paper (starting from Kunin et al., 2010), and it would be worth to at least mention these, although an in-depth discussion would probably lead away from this study's focus. Also, it may be worth mentioning recent algorithmic approaches to tackling this issue, such as DADA2 (Callahan et al., 2016).

- **Results & Discussion,** chimera filtering. The implemented method for chimera filtering appears a little *ad hoc* and heuristic, although the proposed approach certainly makes sense intuitively. However, given the long history of “chimera-slaying” algorithms and the quite sobering benchmark studies on them, some context would be helpful for the reader here – maybe even as a short supplement or as a reference to the user manual. For example, how is the proposed approach conceptually different from existing tools like UCHIME etc? And why was it implemented as is? What was the (empirical?) motivation to do it like this, not otherwise? Personally, I am not very convinced of the performance of chimera-filtering algorithms overall and several recent pipelines side-step the issue more or less elegantly. In the case of NG-Tax (or other reference-based OTU callers), one could even argue that if the reference database is perfectly chimera-free, a closed-reference approach would not need a chimera filtering approach at all, or only one which is based on differential mapping of a sequence to two (highly unrelated) OTUs.

- Table 1 is very large and (on the PDF) unfortunately rotated by 90 degrees. I suggest to convert it into a supplemental Excel sheet which would be more reader-friendly.

- Figure 2 has rotated horizontal axis labels, a 90deg rotated legend – maybe that's just due to formatting of the PDF. It is also difficult to read taxonomic names on the vertical axis in all-caps.

- “Consequently, these methods are more powerful than purely OTU-based methods, [...].” While I agree with this sentence to a certain extent, I believe that the statement should be supported by referring to previous work on the topic. It is not necessarily consensus that 16S “sequence often correlates with phenotypic similarity in key features”, but it is even less clear to what extent phylogenetic diversity estimators capture this signal in a useful way. Arguably, a PD-estimator of UniFrac can only be as good as their underlying tree, which in turn is based on the (representative) sequences of OTUs and thus depends on many factors in the background.

- In particular, the weighted UniFrac measure used in this study seems to be more sensitive to quite a number of factors (including sequencing errors and inflation of small clusters, not irrelevant for the points made in this study) than its unweighted sister in my personal experience, and according to a number of researchers I have talked to on this point.
However, since “personal experience” and “people I’ve talked to” are certainly not a dependable scientific source, and because performance on mock communities should not be severely impacted, I would formulate this as a suggestion and certainly not as a reviewer’s request: were the weighted UF-based results double-checked using unweighted UF and/or a non-phylogenetic method, such as Bray-Curtis?

- In the PCoA (Figure 5, A&C), it is quite hard to decide which method looks “better” purely based on visual impression, not least because the % variance explained on the axes is not equivalent. It would be good to see a more quantitative statement on which approach better recovers expected clusters from the mock communities. The most straightforward approach would be to perform MANOVA analyses, structured by the different factors to test for and then use the effect sizes to quantify the goodness of separation (or non-separation). I would suggest to run e.g. Anderson’s PERMANOVA (http://www.entsoc.org/PDF/MUVE/6_NewMethod_MANOVA1_2.pdf; implementation available through the function “adonis” in the R package vegan) or ANOSIM to this end. Alternatively, samples could be clustered based on beta div and the resulting clusterings (or dendrograms) quantitatively compared to expectations based on different factors.

- Thank you for providing Supplementary Figures 1&2; they are informative in the interpretation of the presented data.

- Similarly, thank you for providing code and data as supplements!

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Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes
If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

**Author Response 14 Nov 2018**

**Javier Ramiro-Garcia**, TI Food and Nutrition (TIFN), Wageningen, The Netherlands

In their manuscript, the authors introduce NG-Tax, an open-source software for the (meta-)analysis of 16S rRNA-based microbiome datasets. Their tool focuses on an important and so-far arguably understudied aspect of microbial ecology research: the integration of results across studies, in view of both technical and biological variation.

The approach is interesting and addresses important points. In particular, several sequencing datasets of different mock communities were generated, even using different primer sets: this is great data to benchmark on, and many (most) other papers introducing tools do not provide benchmarks on such an array of real (mock) data. In general, I feel that this is very interesting work and that NG-Tax can be a promising alternative to existing tools in the field.

We thank the reviewer for his nice comments and also his suggestions about the manuscript.

However, there are several points that I feel would need to be addressed in order for the manuscript to stand tall, and for the reader to get a good understanding of how NG-Tax can be useful in practice.

**Major comments:**
- Even after reading the manuscript and online user manual repeatedly, I have to admit that it is not completely clear to me how NG-Tax works in detail, and in which points exactly it differs from existing approaches. Based on the introduction, I gather that NG-Tax relies on closed-reference OTU picking, but this is not mentioned explicitly anywhere in the text. Also, does reference-based OTU picking in NG-Tax rely on uclust? If yes, which version and parameters were used, and how do they differ from QIIME's defaults? Also, the Background and Discussion sections do not elaborate on the various disadvantages of closed-
reference approaches; most importantly, closed-ref only takes into account sequences matching the database and removes everything else. When integrating sequence data from different primer sets, this is arguably the most straightforward approach; however, the limitations should be discussed. Thanks for the suggestion. In the new version we included a more detailed Figure 1 including those unique aspects of NG-Tax.

We agree with the reviewer that close reference OTU picking has the disadvantage of only taking sequences into account that have a match in the database, and this is incompatible with having stable OTUs since databases change over time. For this reason, NG-Tax employs an open reference approach to remain independent of reference databases. Different clustering algorithms also lead to different OTUs, and hence no clustering process is applied and the generation of OTUs is independent for each sample. Existing open reference approaches generate OTUs for the whole study by clustering the reads from all the samples together. Then, if new samples are included to a previous study, the OTUs need to be regenerated with the reads from the previous study and new samples together, which will lead to discrepancies in the former and new composition of the samples because some of the previous OTUs may not be present in the new analysis anymore.

Instead, in NG-Tax OTUs are generated sample by sample using the following strategy: For each sample reads are ranked by read abundance and OTUs are added to an OTU table starting from the most abundant sequence until the read abundance is lower than a percentage defined by the user (recommended is at is 0.1%). Subsequently, the discarded reads are clustered to the OTU table allowing one mismatch. In practical terms it is guided clustering where seeds are determined by abundance. The differences with an normal clustering approach is that there is no clustering to define the seeds, which allows seeds that differ as little as one nucleotide. The clustering is applied only afterwards to compensate for potential bias due to PCR and sequencing errors. Error is sequence-specific, and hence some sequences could be affected more than others. If a species specific amplicon is more prone to PCR or sequencing errors, the relative abundance of that particular OTU will be underestimated. But after clustering, OTUs more prone to error receive a higher percentage of discarded reads than others, this differential recovery helps to reestablish the true composition that was lost due to sequence specific error rates.

We substituted uclust by usearch in the scripts of the new version.

I gather from the text that NG-Tax's main innovations are the use of primer-tailored reference databases and a different (more conservative) read abundance filtering scheme. It is perfectly valid to benchmark these against QIIME's default settings; however, it would be great to see how QIIME performs with similarly conservative settings, to better understand where NG-Tax's edge in performance comes from.

We think that the main innovation of NG-Tax is the way OTUs are generated. This may seem counter-intuitive because it does not follow the standard approach but it is the discerning step compared with other existing pipelines. This innovative OTU generation algorithm is the reason of the NG-Tax's edge in performance.
With QIIME those conservative thresholds cannot be used because the filtering percentage threshold is defined using the whole library and within a library there are samples that contain 20 times more reads than others. A conservative threshold like 0.1% is conservative for an average sample, not conservative for a big sample at all and extreme for small samples. Hence, OTUs present in only small samples can be discarded even if they represent 1% of that sample but less than 0.1% of the whole dataset. On the other hand, NG-Tax applies thresholds defined by sample accounting for sample heterogeneity.

In the manuscript we used the setting recommended by QIIME and described in Bokulich et al 2013\(^1\). For NG-Tax analysis we also employed recommended default settings so we thought that even if this is not optimal and has its limitations, this could be a fair approach. Nevertheless, we benchmarked with QIIME not to compare performances but rather to show that this dataset is not an exceptional case and the commonly reported problems such as many un- or poorly classified OTUs, inflated richness and diversity, taxonomic profiles that do not match the expected ones, region dependent taxonomic classification and results being highly dependent on minor changes in the experimental setup are also found in this dataset when standard approaches are used.

In Bokulich et al 2013, supplementary material 2; pages 8, 9 and 10, the authors compare expected composition against real sample composition using different parameters, one of them being OTU abundance, and the plot shows that the obtained profiles do not change much using different abundance thresholds.

Nonetheless, as suggested by the reviewer we reproduced the QIIME analysis with a 0.1% abundance threshold and this is included in the supplementary data. The results using 0.1% or 0.005% are consistent.

- Regarding taxonomy assignments, it is valid to compare NG-Tax's uclust-based approach to QIIME's uclust-based approach. However, I believe that the gold standard continues to be the RDP Classifier, and it would be interesting to see a performance comparison to this tool (on the short-read data, not only on full-length reads). Also, how does taxonomic classification by NG-Tax differ conceptually from RTAX (http://www.uio.no/english/services/it/research/hpc/abel/help/software/rtax.html)?

I do believe that they are not equivalent, but the approaches appear somewhat related.

In the manuscript we wanted to show that taxonomy assignment using short reads should be comparable with the assignment using the complete 16S rRNA gene (Figure 2). This is why we employed full length sequences. We could have included also RDP short read based taxonomy but the reads were too short for RDP, and hence genus and many times even family assignment could not be achieved with a minimum threshold value of 50%. In the supplementary data we supplied the theoretical compositions for all mock communities. The files for MC2 V4 and MC2 V5V6 contain all phylotypes and can be uploaded to the RDP classifier to verify the poor performance. In the new manuscript we substituted RDP for SILVA Incremental Aligner (SINA) to classify the full length sequences and we also updated the database in NG-Tax to SILVA 128, improving in both cases the classification.

I read the manuscript suggested by the reviewer and I can say that NG-Tax taxonomic
classification is very similar to rtax. The NG-Tax classifier works as follows:

For each OTU, usearch is used to retrieve hits for the forward and reverse reads against their respective trimmed reference database. Hits that are common between both reads are divided into 6 identity thresholds: 100, 98, 97, 95, 92, 90. A hit belongs to a certain level, for example 97, when both reads have at least a 97 percentage identity with that hit. Using the highest available identity threshold, NG-Tax assigns the consensus taxonomy to the OTU if this taxonomy is supported for at least half of the hits. Genus, Family or Order remains unassigned if the maximum identity percentage level is lower or equal to 97%, 95% and 92% respectively. These are the main differences:

rtax clusters the reference database at 99%, while NG-Tax does not.
rtax averages the percentage identity for both reads and then considers the hits that have an averaged percentage identity 0.5% lower than the maximum averaged percentage identity as valid. NG-Tax does not average the percentage identities and uses fixed values 100, 98, 97, 95, 92 and 90 as thresholds.
For the rest they are indeed very similar approaches.
Therefore we have added rtax to the references and acknowledge in the manuscript that similar dynamic identity thresholds have been already employed to assign taxonomy.
Furthermore, all the details about NG-Tax taxonomic assignment have been added to the user manual.

- In general, the results on taxonomic classification are not discussed quantitatively. From Figures 3&4, the visual impression is that NG-Tax indeed better approximates expected taxonomic profiles than QIIME, but it is hard to quantify this from stacked bar charts. I would suggest to compute e.g. Euclidean or more sophisticated distances of classified taxonomic profiles to the expected distribution. Also, it would be interesting to see quantitative sensitivities and specificities (or F1-scores?) on the taxonomic assignments; particularly also when running on the exact same (more conservatively filtered) dataset for QIIME. Some numbers on specificity are provided in the Abstract and Conclusion sections – but I am not sure if specificity may be gained at the expense of sensitivity based on the more rigid read filtering upstream.

As suggested by the reviewer, distances between compositional profiles and expected profiles are now shown in Figure 5. Distances between taxonomic profiles were calculated as the sum of the weighted differences. Given two taxonomical profiles x and y, for each taxa i, we defined the difference in abundance as \( \text{difi}(x,y) = (x_i - y_i) \) and a weighing factor \( w_i(x,y) = \frac{(x_i - y_i)}{\text{avg}(x_i + y_i)} \). Weighted difference was the result of multiplying the difference in abundance by its weighting factor. This weighting factor is useful to take into account the relative change and not only the absolute change, because a 1% absolute change becomes a 200% or 20% relative change depending on whether the expected abundance is 0.5% or 5% respectively. We performed t tests to compare the performance of NG-Tax versus QIIME from a quantitative point of view.
We have also included an Excel spreadsheet with compositional profiles in the supplementary data.
Figure 2 shows specificity of the taxonomical assignments and has been modified to improve readability.
The QIIME analysis at 0.1% abundance threshold can be found in the supplementary
As a suggestion, but certainly not as a request, I would recommend to maybe include additional, independent datasets to benchmark on. For example, Tremblay et al. (2015) have published data on mock communities sequenced with different primer sets and on different platforms. Such data could contribute to a yet more general assessment of NG-Tax performance.

We thank the reviewer for the suggestion, but including new datasets will imply a rewrite of a big part of the manuscript. We consider that 49 samples can give an idea of NG-Tax performance. Additionally, we would like to mention that NG-Tax has been the reference method for 16S rRNA gene amplicon analysis in our lab for more than two years and has been used in more than 30 manuscripts that have been submitted or are in preparation. One of these manuscripts was published before this manuscript. Since then another fifteen studies using NG-Tax have been published. These studies contain biological samples that belong to very different and specific environments and were sequenced both on MiSeq and HiSeq instruments. These will contribute to the assessment of NG-Tax performance, however these were not included in the current manuscript since they are accessible on the aforementioned publications.

Minor comments (chronologically, not in order of importance):

- Background, “The consequence of this approach is that the ‘quality’ of the clustering of the reference set propagates to reference-picked OTUs.” I believe that as such, this statement is not fully valid or supported. In fact, the negative complement is arguably true: reference-based OTU picking against a “bad” reference can never provide “good” OTUs (a garbage-in, garbage-out problem, so to say). However, even with a good reference, a bad mapping algorithm can generate non-informative reference-based OTU sets. Schloss & Westcott have recently published a study which discusses this point, among others (Westcott & Schloss, 2015).

With this sentence we did not imply that only ‘good quality’ is transferred from the clustered databases to the OTUs, we meant both, pros and cons are transferred. In fact, we agree that references have their limitations and clustered databases also contain bias due to clustering. For this reason NG-Tax employs a de novo OTU picking with no references or clustering involved.

- Background, “However, in 16S rRNA gene amplicon sequencing every sequencing error could potentially lead to the false discovery of a new species.” I have two comments on this statement. First, I believe that the term “species” in this context can be misleading and I feel that the neutral term OTU or diversity unit would be more appropriate. Second, there is a large body of literature on how sequencing errors affect 16S-based diversity studies beyond the cited Bokulich et al paper (starting from Kunin et al., 2010), and it would be worth to at least mention these, although an in-depth discussion would probably lead away from this study’s focus. Also, it may be worth mentioning recent algorithmic approaches to tackling this issue, such as DADA2 (Callahan et al., 2016).

As suggested by the reviewer we rephrased the sequence to avoid the use of “species”. Now we stated: “However, in 16S rRNA gene amplicon sequencing every sequencing error could...
potentially lead to an incorrect OTU classification which may ultimately lead to the false discovery of a new phylotype.”

We added Kunin et al 2010 and Callahan et al. 2016 to the references but we just wanted to point out that sequencing error is an important factor in 16S analysis rather than make an in-depth discussion about it.

- **Results & Discussion, chimera filtering.** The implemented method for chimera filtering appears a little *ad hoc* and heuristic, although the proposed approach certainly makes sense intuitively. However, given the long history of “chimera-slaying” algorithms and the quite sobering benchmark studies on them, some context would be helpful for the reader here – maybe even as a short supplement or as a reference to the user manual. For example, how is the proposed approach conceptually different from existing tools like UCHIME etc? And why was it implemented as is? What was the (empirical?) motivation to do it like this, not otherwise? Personally, I am not very convinced of the performance of chimera-filtering algorithms overall and several recent pipelines side-step the issue more or less elegantly. In the case of NG-Tax (or other reference-based OTU callers), one could even argue that if the reference database is perfectly chimera-free, a closed-reference approach would not need a chimera filtering approach at all, or only one which is based on differential mapping of a sequence to two (highly unrelated) OTUs.

First, we would like to recall that NG-Tax is not reference-based. We fully agree with the reviewer opinion about ‘chimera-slaying’ algorithms. Chimera detectors are often validated using in-silico datasets generated by determining an initial set of valid sequences and a chimera formation pattern. This pattern or “rule” for chimera formation is commonly defined by considering that any two sequences in the initial dataset are equally probable to lead to a chimera and any nucleotide is equally probable to be the point in which these two sequences merge to form the chimera. It is conceivable that maybe the initial set is not representative of the sequences present in a specific real biological sample, not every pair of sequences has the same probability to form chimeras and not all the nucleotides may have the same odd to be the merging point of two sequences. Many different sequence sets can be selected as initial valid sequences and also many different chimera formation patterns can be chosen, but it is very difficult to really determine whether our choices mimic the way in which chimeras are formed in real sequencing data and therefore it is hard to verify if those in-silico created chimeras represent the chimeras that can be found in real sequencing samples. We consider that the proper validation should be using the real sequencing samples. If the chimera detection algorithm works, we would expect a very small number of non-assigned reads (since most chimeras should be aberrant). In case we have positive controls like MC, sequencing profiles and diversity should resemble the expected ones, and this is exactly what we observe with the results of NG-Tax.

We think that there are no perfect chimera-free databases, and a valid OTU can be found in the reference database and at the same time be a perfect combination of 2 other OTUs, especially for regions with lower variability (V4). If all those 3 OTUs are present in the same sample, how can we know whether it is a chimera or real?

In our opinion chimera detection is the weakest step in 16S pipelines because there is no
satisfactory solution to the problem mentioned above. So the prevention against chimeras should come from the experimental design by reducing the PCR cycles and selecting regions of high variability. Chimera removal has many limitations and human supervision is recommended. For this reason, we decided to simplify the chimera detection as much as possible so the researcher can quickly identify why an OTU has been discarded. And also apply stringent parameters (100% identity) to avoid false positives. False negatives should be easier to detect afterwards since most of the chimeras should be aberrant.

In the manuscript of UCHIME they stated that “UCHIME searches for a chimeric alignment between a query sequence (Q) and two candidate parents (A and B)” and “candidate parents are required to have abundance at least $\lambda$ times that of the query sequence, on the assumption that a chimera has undergone fewer rounds of amplification and will therefore be less abundant than its parents. The parameter $\lambda$ is called the abundance skew, and by default $\lambda=2$”, so NG-Tax approach is very similar to de novo UCHIME approach, but NG-Tax treats forward and reverse reads separately.

Table 1 is very large and (on the PDF) unfortunately rotated by 90 degrees. I suggest to convert it into a supplemental Excel sheet which would be more reader-friendly.

As suggested by the reviewer Table 1 is now supplied as an excel spreadsheet in the supplementary material.

Figure 2 has rotated horizontal axis labels, a 90deg rotated legend – maybe that's just due to formatting of the PDF. It is also difficult to read taxonomic names on the vertical axis in all-caps.

As suggested by the reviewer we modified Figure 2 to increase readability.

“Consequently, these methods are more powerful than purely OTU-based methods, [...].” While I agree with this sentence to a certain extent, I believe that the statement should be supported by referring to previous work on the topic. It is not necessarily consensus that 16S “sequence often correlates with phenotypic similarity in key features”, but it is even less clear to what extent phylogenetic diversity estimators capture this signal in a useful way. Arguably, a PD-estimator of UniFrac can only be as good as their underlying tree, which in turn is based on the (representative) sequences of OTUs and thus depends on many factors in the background.

Taxonomic assignment of the OTUs suffers from the same problems raised by the reviewer. Not always does 16S rRNA gene sequence similarity correlate with phenotypic similarity and the taxonomical assignment is as good as the reference database and the classifier employed. But having a composition barplot with OTUs named by number rather than by taxonomy would mean that all the information provided by the nucleotide sequence is discarded. This information may not be perfect but we cannot neglect that this information transformed into taxonomical assignment is useful at least to some extent.

The same criterion was applied to evaluate diversity. We used phylogenetic methods, which retain the information of the nucleotide sequence. We acknowledge the limitations but we argue that a sample containing 5 OTUs with a 99% pairwise sequence identity should not be given the same (potential) diversity that a sample containing 5 OTUs with less than 85% pairwise sequence identity. We consider that phylogenetic methods are more powerful because they use all information available, however, we should not over extrapolate the results. 16S rRNA gene amplicon sequencing should be taken as exploratory approach, whereas metagenomic and metatranscriptomic sequencing provides a more suitable and
precise approach if we really want to focus on microbial functionality.

- In particular, the weighted UniFrac measure used in this study seems to be more sensitive to quite a number of factors (including sequencing errors and inflation of small clusters, not irrelevant for the points made in this study) than its unweighted sister in my personal experience, and according to a number of researchers I have talked to on this point. However, since “personal experience” and “people I’ve talked to” are certainly not a dependable scientific source, and because performance on mock communities should not be severely impacted, I would formulate this as a suggestion and certainly not as a reviewer’s request: were the weighted UF-based results double-checked using unweighted UF and/or a non-phylogenetic method, such as Bray-Curtis?

As suggested by the reviewer we have included the unweighted UniFrac and Bray-Curtis analysis in the supplementary material. The results obtained by all three methods are in concordance.

- In the PCoA (Figure 5, A&C), it is quite hard to decide which method looks “better” purely based on visual impression, not least because the % variance explained on the axes is not equivalent. It would be good to see a more quantitative statement on which approach better recovers expected clusters from the mock communities. The most straightforward approach would be to perform MANOVA analyses, structured by the different factors to test for and then use the effect sizes to quantify the goodness of separation (or non-separation). I would suggest to run e.g. Anderson’s PERMANOVA (implementation available through the function “adonis” in the R package vegan) or ANOSIM to this end. Alternatively, samples could be clustered based on beta div and the resulting clusterings (or dendrograms) quantitatively compared to expectations based on different factors.

Thanks for the suggestion. In the new version of the manuscript we performed a more quantitative analysis of the sequencing data and the expected MC. We performed a permanova analysis under MC type factor and it was significant for both pipelines meaning that some of the variance is explained by the Mock type. But to really evaluate accuracy and reproducibility and compare pipelines performances we used pairwise distances and t tests (Figure 7 and Dataset 1).

- Thank you for providing Supplementary Figures 1&2; they are informative in the interpretation of the presented data.

Thank you.

- Similarly, thank you for providing code and data as supplements!

Thank you.

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Competing Interests: No competing interests were disclosed.

Comments on this article

Version 1

Author Response 07 Feb 2018

Javier Ramiro-Garcia, TI Food and Nutrition (TIFN), Wageningen, The Netherlands

A new version of NG-Tax can be downloaded from https://github.com/JavierRamiroGarcia/NG-Tax

Competing Interests: No competing interests were disclosed.

Author Response 25 Jul 2016

Javier Ramiro-Garcia, Wageningen University and Research Centre, The Netherlands

The proper link to download the pipeline is:

http://www.systemsbiology.nl/NG-Tax/

We will correct the link in version 2 of the paper.
Sorry for the inconveniences.

Javier Ramiro-Garcia
**Competing Interests:** No competing interests were disclosed.

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