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Subsampled open-reference clustering creates consistent, comprehensive OTU definitions and scales to billions of sequences

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ABSTRACT

We present a performance-optimized algorithm, subsampled open-reference OTU picking, for assigning marker gene (e.g., 16S rRNA) sequences generated on next-generation sequencing platforms to operational taxonomic units (OTUs) for microbial community analysis. This algorithm provides benefits over de novo OTU picking (clustering can be performed largely in parallel, reducing runtime) and closed-reference OTU picking (all reads are clustered, not only those that match a reference database sequence with high similarity). Because more of our algorithm can be run in parallel relative to "classic" open-reference OTU picking, it makes open-reference OTU picking tractable on massive amplicon sequence data sets (though on smaller data sets, "classic" open-reference OTU clustering is often faster). We illustrate that here by applying it to the first 15,000 samples sequenced for the Earth Microbiome Project (1.3 billion V4 16S rRNA amplicons). To the best of our knowledge, this is the largest OTU picking run ever performed, and we estimate that our new algorithm runs in less than 1/5 the time than would be required of "classic" open reference OTU picking. We show that subsampled open-reference OTU picking yields results that

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are highly correlated with those generated by "classic" open-reference OTU picking through comparisons on three well-studied datasets. An implementation of this algorithm is provided in the popular QIIME software package, which uses uclust for read clustering. All analyses were performed using QIIME's uclust wrappers, though we provide details (aided by the open-source code in our GitHub repository) that will allow implementation of subsampled open-reference OTU picking independently of QIIME (e.g., in a compiled programming language, where runtimes should be further reduced). Our analyses should generalize to other implementations of these OTU picking algorithms. Finally, we present a comparison of parameter settings in QIIME's OTU picking workflows and make recommendations on settings for these free parameters to optimize runtime without reducing the quality of the results. These optimized parameters can vastly decrease the runtime of uclust-based OTU picking in QIIME.

Subjects Bioinformatics, Ecology, Microbiology **Keywords** OTU picking, Microbial ecology, Microbiome, Qiime, Bioinformatics

INTRODUCTION

Three high-level strategies for defining Operational Taxonomic Unit (OTU) cluster centroids have been widely applied for centroid-based greedy clustering (*Li & Godzik*, 2006; *Edgar*, 2010) of marker gene (e.g., 16S rRNA) sequences generated on next-generation sequencing platforms to facilitate microbial community analysis. These are canonically described as *de novo*, closed-reference, and open-reference OTU picking (*Navas-Molina et al.*, 2013). In each of these approaches, respectively, centroids are defined internally based only on the sequences being clustered, based only on an external, predefined database of cluster centroids, or based on a combination of the two. Each of these methods has benefits and drawbacks.

In *de novo* OTU picking, input sequences are aligned against one another, and sequences that align with greater than a user-specified percent identity are defined as belonging to the same OTU. There are many variations and free parameters in this process, such as how many alignments are performed before a sequence is assigned to an OTU or used to define a new OTU, but the common feature of these methods is that no external reference database is required. This is also the primary advantage of this method: it is not necessary to have accumulated a collection of reference sequences before working with a new marker gene. However, *de novo* OTU picking is difficult to parallelize because all processes must be able to use new OTUs that are defined by other processes. Consequently, this approach cannot scale to modern-sized data sets.

In closed-reference OTU picking, input sequences are aligned to pre-defined cluster centroids in a reference database. If the input sequence does not match any reference sequence at a user-defined percent identity threshold, that sequence is excluded. The primary advantage of closed-reference OTU picking is that it is easily parallelizable.

Because the cluster centroids are predefined, the input sequence collection can be partitioned into *n* subsets, the assignment process can be split across *n* processors, and the clustering results can be collated when all processes have completed. This dramatically reduces the "wall time" (i.e., the total time to completion as you would see it on a clock on the wall, not in terms of CPU \times hours) of this method, and makes closed-reference OTU picking a convenient strategy for extremely large datasets (e.g., as in Yatsunenko et al., 2012). Additionally, it has the convenient feature that, because OTUs are defined by a pre-existing reference, there are typically high-quality taxonomic assignments for each OTU, and a high-quality phylogenetic tree, often based on full-length sequences rather than fragments, exists and describes the relationships among those OTUs. Furthermore, because input sequences are not compared directly to one another, but rather to an external reference, the input sequences need not overlap. This is essential, for example, if performing a meta-analysis including sequences derived from different amplification products of the same marker gene, such as the V2 and V4 regions of the 16S rRNA (e.g., as in the meta-analysis performed in *Caporaso et al.*, 2010). The major drawback to closed-reference OTU picking, however, is that it cannot identify novel diversity: if a sequence has no match in the reference database, it cannot be included in the analysis, restricting analyses to already-known taxa. (Of course, the importance of this limitation decreases as the reference database increases in coverage.)

Finally, open-reference OTU picking combines the previous protocols. First, input sequences are clustered against a reference database in parallel in a closed-reference OTU picking process. However, rather than discarding sequences that fail to match the reference, these "failures" are clustered *de novo* in a serial process. Open-reference OTU picking offers benefits over both the *de novo* and closed-reference protocols. Because it includes the parallel closed-reference step, it will typically run faster than *de novo* OTU picking. And, since it includes *de novo* OTU picking of the sequences that fail to hit the reference database, all sequences are clustered, so analyses are not restricted to already-known OTUs. However, because the *de novo* clustering process is run serially, it can still be prohibitively slow for very large datasets or datasets with a substantial number of sequences that fail to hit the reference database. Because of these long runtimes, it has not yet been widely applied despite the benefits it offers.

We present a novel strategy for open-reference OTU picking that allows a larger portion of the computation to be run in parallel, which we call *subsampled open-reference OTU picking*, allowing open-reference OTU picking on very large datasets. We compare this method to "classic" open-reference OTU picking (as described in the previous paragraph) to confirm that, despite potentially slightly different OTU definitions, the summary statistics that are often used derive biological conclusions from application of these different methods to the same data set would remain the same. To achieve this, we show that alpha diversity, beta diversity, and taxonomic profiles are highly correlated between the "classic" open-reference OTU picking and subsampled open-reference OTU picking. We also compare these methods to *de novo* and closed-reference OTU picking, and explore the effect of dataset and algorithm parameters on runtime and analysis results. We note

that we specifically focus on centroid-based greedy clustering approaches in this study (e.g., as in uclust and cd-hit *Li & Godzik*, 2006; *Edgar*, 2010), not approaches that require alignment of all pairs of unique sequences (i.e., the hierarchical methods described in *Schloss & Westcott*, 2011), as the former scale better to larger data sets. However, because our full evaluation framework (metrics and data sets) and the EMP raw sequence data are all freely accessible, it is straightforward for other groups to reproduce these evaluations on alternative methods.

All analyses presented here are performed using the QIIME and pandas python packages. As far as we know, QIIME contains the only existing implementation of the sub-sampled open-reference OTU picking algorithm, but the algorithm is not QIIME-specific. Thus while our comparison is based on specific QIIME/uclust-based implementations of *de novo, closed reference, classic open reference,* and *subsampled open reference* OTU picking, our findings should be general to other implementations of these algorithms.

MATERIALS AND METHODS

Subsampled open-reference OTU picking algorithm

Open-reference OTU picking is preferable to the other methods presented here because it combines the advantages of closed-reference and de novo clustering. However, the de novo step of open-reference OTU picking can only be run serially, and therefore can be time-consuming for large datasets if many sequences fail to hit the reference database. To improve the runtime of open-reference OTU picking, we developed subsampled open-reference OTU picking, which incrementally increases the size of the reference database by *de novo* clustering a subset of the sequences that fail to match the reference database. The remainder of the sequences that fail to hit the reference database can then be clustered against these new cluster centroids in a parallel closed-reference OTU picking process. This allows for partial parallelization of the *de novo* clustering step and can significantly decrease runtime on large datasets, allowing open-reference OTU picking to scale to billions of input sequences (e.g., as generated in multiple Illumina HiSeq 2000 runs). It can additionally be run iteratively, so that representative sequences for the new (i.e., non-reference) OTUs can be combined with the reference database for future OTU picking runs. It is important to note that runtime is not always reduced with subsampled open-reference OTU picking. Data set and algorithm parameters have a large effect on runtime (discussed further in *Runtime differences*). This approach is similar to the Buckshot algorithm (Cutting et al., 1992; Jensen et al., 2002), initially described for semantic clustering of documents in a corpus, though we do not use the parallel hierarchical clustering approach described by Jensen et al. (2002) for initial clustering definition.

A detailed description of this workflow is illustrated in Fig. 1. It is implemented using uclust v1.2.22q (*Edgar*, 2010) for clustering in QIIME 1.6.0 (*Caporaso et al.*, 2010) and later, though any sequence clustering software that provides support for *de novo* and closed-reference clustering could be substituted for uclust in an alternate implementation. The inputs provided to this method are demultiplexed, quality-filtered sequences, and a reference sequence collection (for example, the Greengenes 13_8 97% OTU representative



sequences DeSantis et al., 2006; McDonald et al., 2012b). First, sequences are clustered in parallel using a closed-reference OTU picking workflow, where sequences are queried against the reference database at percent identity s (default 97%). If a read matches a reference sequence at greater than or equal to s% identity, it is assigned to the OTU defined by that reference sequence. These are referred to as the reference OTUs. Next, a random subsample of n% (*n* should be small, the default value in QIIME 1.8.0-dev and earlier is 0.1%) of the sequences that failed to match the reference sequence collection are clustered de novo, and the cluster centroids for all resulting OTUs are used to define a new reference sequence collection. Those OTUs are referred to as the new reference OTUs. The sequences that were not included in the random subsample that was clustered de novo then go through an additional round of parallel closed-reference OTU picking, this time where they are clustered against the new reference OTUs based on matching a sequence in the new reference sequence collection at greater than or equal to s% identity. This creation of a "new reference database" allows us to harness the parallelization of our closed-reference OTU picking pipeline, greatly decreasing the time it takes for sequences that fail to hit the initial reference database to be clustered into OTUs. In the final clustering step, sequences that fail to hit a reference sequence during this final closed-reference OTU picking step are clustered *de novo*. These are referred to as the clean-up OTUs. Finally, the reference OTUs, new reference OTUs, and clean-up OTUs are combined into a single OTU table (i.e., table of counts of OTUs on a per-sample basis, as described in *McDonald et al.* (2012a)), and this table, as well as a filtered table excluding OTUs with counts less than or equal to a user-defined threshold c, are provided to the user. By default, c = 2, so each OTU is observed at least twice (i.e., singleton OTUs are excluded). Because many more of the sequences can be clustered using closed-reference OTU picking in this workflow, it can run in far less time than classic open-reference OTU picking (see Runtime differences section below).

Evaluation of subsampled open-reference OTU picking

We validated the subsampled open-reference OTU picking workflow by comparing it to *de novo*, closed-reference, and classic (i.e., non subsampled) open-reference clustering methods on three different datasets: the Lauber "88 Soils" study (*Lauber et al., 2009*) (referred to as *88-soils* here), the Caporaso "Moving Pictures" study (*Caporaso et al., 2011*) (referred to as *moving-pictures* here), and the Costello "Whole Body" study (*Costello et al., 2009*) (referred to as *whole-body* here) using three metrics. Table 1 provides a description of the OTU picking methods being compared. First, we tested the correlation between sample alpha diversities (OTU counts, i.e., QIIME's *observed species* metric, and Phylogenetic Diversity (PD) (*Faith, 1992*)) based on subsampled open-reference OTU picking and the other OTU picking protocols. Next, we tested whether beta diversity patterns (as determined by weighted and unweighted UniFrac (*Lozupone & Knight, 2005*) distances between samples) were consistent across OTU picking protocols, based on Mantel tests (*Mantel, 1967*) with 1,000 Monte Carlo iterations. Finally, we tested whether the same taxonomic profiles were obtained on a per-sample basis using each of the OTU picking

Table 1 Meth.refer to each mpick-de-novo-absent. The exc	od definitions. Def tethod by its abbrev otus.py command. tet command/parar	initions of the OTU pickin riation for simplicity. We no ucr is applied when pick_ot neter combinations used fo	g methods ote that the cus:otu-picl or each OTU	being con both de n king_metho U picking r	npared he ovo (uc) ; od uclust. 'un are pr	rre, base and clas ref is sp ovided j	ed on the a sic openre occified in in the stud-	ubbreviatio ference OT the parame y's GitHub	ns used thi U picking ters file, ar repository	roughout the (ucr) are acc nd uc is appli (see Data Av	paper. From essed through ed when that ailability).	here, we QIIME's option is
Abbreviation	Title	Command	max_ accepts	max_ rejects	step words	word length	prefilter_ percent_id	min_ otu_size	speed_ mode	Processors	reference_ percent_id	subsample_ fraction
uc	De novo	pick_de_novo_otus.py	20	500	20	12	NA	NA	slow	1	0.97	NA
ucr	Legacy open reference	pick_de_novo_otus.py	20	500	20	12	NA	NA	slow	10	0.97	NA
ucrC	Closed reference	pick_closed_reference_otus.py	20	500	20	12	NA	NA	slow	10	0.97	NA
ucrss	Subsampled open reference	pick_open_reference_otus.py	20	500	20	12	0	1	slow	10	0.97	0.001
ucrss_wfilter	Subsampled open reference, filtered	pick_open_reference_otus.py	20	500	20	12	0.6	1	slow	10	0.97	0.001
uc_fast	De novo, fast settings	pick_de_novo_otus.py	1	8	8	8	NA	NA	fast	1	0.97	NA
ucr_fast	Legacy open reference, fast settings	pick_de_novo_otus.py	1	œ	œ	×	NA	NA	fast	10	0.97	NA
ucrC_fast	Closed reference, fast settings	pick_closed_reference_otus.py	1	8	8	8	NA	NA	fast	10	0.97	NA
ucrss_fast	Subsampled open reference, fast settings	pick_open_reference_otus.py	1	œ	∞	×	0	-	fast	10	0.97	0.001
ucrss_wfilter_fast	Subsampled open reference, filtered, fast settings	pick-open reference-otus.py	-	œ	œ	×	0.6	1	fast	10	0.97	0.001

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1 182	Title Legacy open	Command pick.de.novo.otus.py	max_ accepts 1	max_ rejects 8	step words 8	word length 8	prefilter_ percent_id 0	min_ otu_size 1	speed_ mode fast	Processors 29	reference_ percent_id 0.82	subsample_ fraction 0.001
refe fast 82% OTU 29 F	rence, settings, 5 reference Js, rocessors											
Le, ref fas 29	gacy open erence, t settings, processors	pick.de_novo_otus.py	-	œ	×	œ	0	1	fast	29	0.97	0.001
Su rei 82 0' 29 29	bsampled open ference, st settings, % reference IUs, processors	pick-open_reference-otus.py	_	ø	œ	×	0	-	fast	29	0.82	0.001
Su ref fas 29	bsampled open erence, :t settings, processors	pick.open.reference.otus.py	-	œ	œ	œ	0	1	fast	29	0.97	0.001
Su fav 29	bsampled open ference, st settings, ' processors, & subsample	pick-open_reference_otus.py	-	∞	œ	œ	0	1	fast	29	0.97	0.1

methods. It is important to note that we are not trying to assess whether one method is better than another using these metrics. Instead, we are testing whether the methods give highly correlated results.

Data availability

The raw sequence data analyzed in this study is available in the QIIME Database under study numbers 103 (88-soils), 449 (whole-body), and 550 (moving-pictures). All analyses were run with QIIME 1.8.0-dev. All commands, as well as all processed data and IPython Notebooks that illustrate how to work with that data, are available in this project's GitHub repository at https://github.com/gregcaporaso/cloaked-octo-ninja.

RESULTS AND DISCUSSION

Subsampled versus "classic" open-reference OTU picking

Alpha diversity (Table 2; whole-body PD Pearson r = 0.989; 88-soils PD Pearson r = 0.930; moving-pictures PD Pearson r = 0.996), beta diversity (Table 3; whole-body unweighted UniFrac Mantel r = 0.948; 88-soils unweighted UniFrac Mantel r = 0.939; moving-pictures unweighted UniFrac Mantel r = 0.991) and taxonomic summaries (Table 4; whole-body: r = 0.999 at phylum level, 0.999 at species level; 88-soils r = 0.999 at phylum level, r = 0.999 at species level; moving-pictures r = 0.999 at phylum level, r = 0.999 at species level) were highly correlated between classic and subsampled open-reference OTU picking. Minor differences likely arise from the non-deterministic step of rarefying all samples to even sampling depth before comparing samples. These results suggest that subsampled open-reference picking yields the same results as classic open-reference OTU picking, including identical numbers of sequences failing to hit the reference database, and therefore is a suitable replacement.

Application to the Earth Microbiome Project dataset

In order to evaluate the effectiveness of the subsampled open-reference OTU picking method on an extremely large data set, the first 15,000 samples (1.3 billion V4 16S rRNA amplicons) from the Earth Microbiome Project (EMP, Gilbert et al., 2010) were processed on the Amazon Web Services (AWS) EC2 platform. These samples were split across more than 60 studies, which were clustered iteratively. To the best of our knowledge, this is the largest OTU picking run ever completed. We created a StarCluster-based (http://star.mit. edu/cluster/) virtual cluster on AWS using between 8 and 18 M2.4xlarge spot instances (the number of instances was varied at different stages of the run). Each instance (or virtual cluster node) had 69 GB RAM and 8 cores. A total of 11,242 CPU hours were consumed to complete subsampled open-reference OTU picking (at 97% nucleotide identity), and the combined input and output files consumed 1.2 TB of disk space. (This runtime includes the pre-filtering step. The process would have completed much faster if this were disabled.) The resulting OTU table contained 5.6 million non-singleton OTUs. This is the largest number of OTUs identified, and the most comprehensive survey of microbial diversity across environment types to date, so it likely suggests the magnitude of the lower-bound on the microbial diversity of the Earth (although the accuracy is limited because some of

Table 2 Alpha diversity results. Pearson correlation coefficients (*r*) of alpha diversity for (a) 88-soils PD, (b) moving-pictures PD, (c) whole-body PD, (d) 88-soils observed species, (e) moving-pictures observed species, and (f) moving-pictures observed species.

(a)										
	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	1	0.951	0.933	0.934	0.953	0.956	0.936	0.927	0.948	0.947
ucr	0.951	1	0.902	0.931	0.93	0.946	0.94	0.903	0.952	0.944
ucrC	0.933	0.902	1	0.894	0.909	0.905	0.914	0.978	0.902	0.911
ucrss	0.934	0.931	0.894	1	0.929	0.944	0.935	0.894	0.948	0.949
ucrss_wfilter	0.953	0.93	0.909	0.929	1	0.952	0.933	0.903	0.931	0.943
uc_fast	0.956	0.946	0.905	0.944	0.952	1	0.953	0.898	0.956	0.96
ucr_fast	0.936	0.94	0.914	0.935	0.933	0.953	1	0.914	0.95	0.952
ucrC_fast	0.927	0.903	0.978	0.894	0.903	0.898	0.914	1	0.902	0.903
ucrss_fast	0.948	0.952	0.902	0.948	0.931	0.956	0.95	0.902	1	0.962
ucrss_fast_wfilter	0.947	0.944	0.911	0.949	0.943	0.96	0.952	0.903	0.962	1
(b)										

	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	1	0.996	0.993	0.996	0.996	0.995	0.996	0.992	0.996	0.996
ucr	0.996	1	0.993	0.997	0.997	0.995	0.996	0.992	0.996	0.997
ucrC	0.993	0.993	1	0.994	0.991	0.994	0.994	0.998	0.995	0.994
ucrss	0.996	0.997	0.994	1	0.996	0.996	0.997	0.994	0.997	0.997
ucrss_wfilter	0.996	0.997	0.991	0.996	1	0.994	0.995	0.991	0.996	0.996
uc_fast	0.995	0.995	0.994	0.996	0.994	1	0.997	0.994	0.997	0.996
ucr_fast	0.996	0.996	0.994	0.997	0.995	0.997	1	0.994	0.997	0.997
ucrC_fast	0.992	0.992	0.998	0.994	0.991	0.994	0.994	1	0.994	0.994
ucrss_fast	0.996	0.996	0.995	0.997	0.996	0.997	0.997	0.994	1	0.997
ucrss_fast_wfilter	0.996	0.997	0.994	0.997	0.996	0.996	0.997	0.994	0.997	1

(c)										
	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	1	0.985	0.957	0.985	0.985	0.984	0.986	0.961	0.983	0.984
ucr	0.985	1	0.956	0.99	0.989	0.988	0.987	0.96	0.987	0.986
ucrC	0.957	0.956	1	0.961	0.958	0.959	0.961	0.99	0.953	0.961
ucrss	0.985	0.99	0.961	1	0.991	0.988	0.99	0.964	0.989	0.987
ucrss_wfilter	0.985	0.989	0.958	0.991	1	0.985	0.989	0.963	0.987	0.985
uc_fast	0.984	0.988	0.959	0.988	0.985	1	0.986	0.961	0.986	0.985
ucr_fast	0.986	0.987	0.961	0.99	0.989	0.986	1	0.965	0.988	0.989
ucrC_fast	0.961	0.96	0.99	0.964	0.963	0.961	0.965	1	0.957	0.965
ucrss_fast	0.983	0.987	0.953	0.989	0.987	0.986	0.988	0.957	1	0.986
ucrss_fast_wfilter	0.984	0.986	0.961	0.987	0.985	0.985	0.989	0.965	0.986	1

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these OTUs may be artifacts of PCR or sequencing: such artifacts, e.g., chimeras, need to be identified after the OTU picking step).

We were next interested in how long the de novo clustering step of classic open-reference OTU picking would take on the EMP data set, but as we'll illustrate this is an intractable problem in practice with current computer hardware. We began by applying de novo

Table 2 (continued)

(d)

	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	1	0.948	0.88	0.909	0.924	0.935	0.934	0.877	0.925	0.913
ucr	0.948	1	0.905	0.946	0.947	0.947	0.953	0.903	0.938	0.932
ucrC	0.88	0.905	1	0.926	0.888	0.882	0.908	0.973	0.91	0.896
ucrss	0.909	0.946	0.926	1	0.932	0.923	0.935	0.915	0.931	0.929
ucrss_wfilter	0.924	0.947	0.888	0.932	1	0.943	0.946	0.884	0.932	0.927
uc_fast	0.935	0.947	0.882	0.923	0.943	1	0.942	0.883	0.941	0.94
ucr_fast	0.934	0.953	0.908	0.935	0.946	0.942	1	0.908	0.943	0.932
ucrC_fast	0.877	0.903	0.973	0.915	0.884	0.883	0.908	1	0.904	0.906
ucrss_fast	0.925	0.938	0.91	0.931	0.932	0.941	0.943	0.904	1	0.953
ucrss_fast_wfilter	0.913	0.932	0.896	0.929	0.927	0.94	0.932	0.906	0.953	1
(e)										
	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	1	0.992	0.984	0.992	0.992	0.989	0.99	0.978	0.989	0.99
ucr	0.992	1	0.994	0.998	0.998	0.992	0.997	0.991	0.997	0.997
ucrC	0.984	0.994	1	0.995	0.995	0.984	0.993	0.997	0.994	0.994
ucrss	0.992	0.998	0.995	1	0.998	0.992	0.997	0.991	0.997	0.997
ucrss_wfilter	0.992	0.998	0.995	0.998	1	0.992	0.997	0.991	0.997	0.997
uc₋fast	0.989	0.992	0.984	0.992	0.992	1	0.993	0.981	0.992	0.992
ucr_fast	0.99	0.997	0.993	0.997	0.997	0.993	1	0.992	0.998	0.998
ucrC_fast	0.978	0.991	0.997	0.991	0.991	0.981	0.992	1	0.993	0.992
ucrss_fast	0.989	0.997	0.994	0.997	0.997	0.992	0.998	0.993	1	0.998
ucrss_fast_wfilter	0.99	0.997	0.994	0.997	0.997	0.992	0.998	0.992	0.998	1
(f)										
	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	1	0.986	0.971	0.986	0.986	0.993	0.988	0.972	0.988	0.987
ucr	0.986	1	0.984	0.995	0.995	0.987	0.993	0.98	0.993	0.993
ucrC	0.971	0.984	1	0.985	0.984	0.97	0.981	0.992	0.98	0.979
ucrss	0.986	0.995	0.985	1	0.995	0.987	0.993	0.981	0.993	0.992
ucrss_wfilter	0.986	0.995	0.984	0.995	1	0.986	0.993	0.979	0.992	0.992
uc_fast	0.993	0.987	0.97	0.987	0.986	1	0.989	0.972	0.99	0.988
ucr_fast	0.988	0.993	0.981	0.993	0.993	0.989	1	0.981	0.994	0.994
ucrC_fast	0.972	0.98	0.992	0.981	0.979	0.972	0.981	1	0.982	0.979
ucrss_fast	0.988	0.993	0.98	0.993	0.992	0.99	0.994	0.982	1	0.995
ucrss fast wfilter	0.987	0.993	0.979	0.992	0.992	0.988	0 994	0.979	0.995	1

clustering using the "fast" uclust parameter settings to the representative sequences from the 5.6 million non-singleton OTUs from the run described above. These representative sequences represent the full alpha diversity of the EMP data set (a property known to be important to runtime of de novo and open reference OTU clustering) but the data set contains only 5.6 m sequences, so is feasible to cluster de novo. We then subsampled this to contain between 10% and 80% of those sequences, in steps of 10% with 10 iterations at each step, and compiled the runtime for each clustering run. Figure 2

Table 3 Beta diversity results. Mantel correlation coefficients (*r*) of beta diversity for (a) 88-soils unweighted UniFrac, (b) moving-pictures unweighted UniFrac, (c) whole-body unweighted UniFrac, (d) 88-soils weighted UniFrac, (e) moving-pictures weighted UniFrac, and (f) moving-pictures weighted UniFrac.

1	``
(a)
	•••

(b)

	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	0.935	0.908	0.944	0.942	0.939	0.945	0.909	0.943	0.941
ucr	NA	NA	0.915	0.94	0.945	0.934	0.942	0.918	0.944	0.949
ucrC	NA	NA	NA	0.917	0.91	0.926	0.913	0.95	0.917	0.92
ucrss	NA	NA	NA	NA	0.94	0.938	0.945	0.914	0.938	0.942
ucrss_wfilter	NA	NA	NA	NA	NA	0.934	0.943	0.907	0.942	0.941
uc_fast	NA	NA	NA	NA	NA	NA	0.938	0.92	0.939	0.941
ucr_fast	NA	NA	NA	NA	NA	NA	NA	0.909	0.946	0.947
ucrC_fast	NA	NA	NA	NA	NA	NA	NA	NA	0.917	0.924
ucrss_fast	NA	NA	NA	NA	NA	NA	NA	NA	NA	0.945
ucrss_fast_wfilter	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA

	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	0.992	0.974	0.988	0.988	0.992	0.991	0.977	0.991	0.992
ucr	NA	NA	0.982	0.992	0.991	0.991	0.992	0.984	0.993	0.993
ucrC	NA	NA	NA	0.986	0.985	0.973	0.982	0.994	0.981	0.981
ucrss	NA	NA	NA	NA	0.99	0.988	0.992	0.987	0.992	0.991
ucrss_wfilter	NA	NA	NA	NA	NA	0.986	0.99	0.986	0.99	0.991
uc_fast	NA	NA	NA	NA	NA	NA	0.991	0.976	0.992	0.991
ucr_fast	NA	NA	NA	NA	NA	NA	NA	0.983	0.993	0.992
ucrC_fast	NA	NA	NA	NA	NA	NA	NA	NA	0.982	0.983
ucrss_fast	NA	NA	NA	NA	NA	NA	NA	NA	NA	0.993
ucrss_fast_wfilter	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA

(c)

. ,										
	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	0.935	0.891	0.938	0.936	0.93	0.926	0.889	0.933	0.925
ucr	NA	NA	0.899	0.948	0.95	0.934	0.931	0.895	0.941	0.927
ucrC	NA	NA	NA	0.908	0.899	0.878	0.885	0.952	0.897	0.878
ucrss	NA	NA	NA	NA	0.953	0.938	0.936	0.905	0.945	0.928
ucrss_wfilter	NA	NA	NA	NA	NA	0.937	0.94	0.894	0.941	0.932
uc_fast	NA	NA	NA	NA	NA	NA	0.942	0.872	0.939	0.938
ucr_fast	NA	NA	NA	NA	NA	NA	NA	0.888	0.939	0.948
ucrC_fast	NA	NA	NA	NA	NA	NA	NA	NA	0.891	0.879
ucrss_fast	NA	NA	NA	NA	NA	NA	NA	NA	NA	0.933
ucrss_fast_wfilter	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA

(continued on next page)

illustrates the relationship between runtime and input sequence count, along with the results of a regression analysis presenting median runtime as a function of sequence count ($r^2 = 0.98, p = 8e-6$).

In the subsampled open-reference OTU picking run on the EMP dataset, 660 million sequences failed to hit the reference database, and therefore need to be clustered de novo

Table 3 (continued)

(d)

	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	0.896	0.936	0.951	0.901	0.925	0.937	0.924	0.956	0.902
ucr	NA	NA	0.896	0.889	0.966	0.891	0.939	0.895	0.901	0.947
ucrC	NA	NA	NA	0.919	0.914	0.906	0.928	0.984	0.931	0.896
ucrss	NA	NA	NA	NA	0.9	0.917	0.947	0.903	0.949	0.899
ucrss_wfilter	NA	NA	NA	NA	NA	0.885	0.938	0.911	0.899	0.94
uc_fast	NA	NA	NA	NA	NA	NA	0.909	0.898	0.919	0.874
ucr_fast	NA	NA	NA	NA	NA	NA	NA	0.92	0.952	0.96
ucrC_fast	NA	NA	NA	NA	NA	NA	NA	NA	0.918	0.89
ucrss_fast	NA	NA	NA	NA	NA	NA	NA	NA	NA	0.918
ucrss_fast_wfilter	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
(e)										
	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc₋fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	0.971	0.949	0.97	0.973	0.972	0.977	0.949	0.974	0.966
ucr	NA	NA	0.928	0.952	0.952	0.957	0.958	0.928	0.96	0.954
ucrC	NA	NA	NA	0.96	0.94	0.948	0.934	0.999	0.965	0.932
ucrss	NA	NA	NA	NA	0.938	0.965	0.955	0.96	0.98	0.932
ucrss_wfilter	NA	NA	NA	NA	NA	0.946	0.966	0.941	0.951	0.967
uc_fast	NA	NA	NA	NA	NA	NA	0.97	0.948	0.971	0.949
ucr_fast	NA	NA	NA	NA	NA	NA	NA	0.934	0.967	0.967
ucrC_fast	NA	NA	NA	NA	NA	NA	NA	NA	0.965	0.932
ucrss_fast	NA	NA	NA	NA	NA	NA	NA	NA	NA	0.951
ucrss_fast_wfilter	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
(f)										
	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	0.947	0.896	0.934	0.943	0.96	0.939	0.898	0.904	0.936
ucr	NA	NA	0.9	0.924	0.95	0.951	0.92	0.904	0.871	0.944
ucrC	NA	NA	NA	0.886	0.924	0.907	0.911	0.994	0.831	0.939
ucrss	NA	NA	NA	NA	0.944	0.92	0.917	0.882	0.918	0.911
ucrss_wfilter	NA	NA	NA	NA	NA	0.933	0.918	0.926	0.897	0.932
uc_fast	NA	NA	NA	NA	NA	NA	0.955	0.909	0.889	0.966
ucr_fast	NA	NA	NA	NA	NA	NA	NA	0.91	0.936	0.951
ucrC_fast	NA	NA	NA	NA	NA	NA	NA	NA	0.83	0.94
ucrss_fast	NA	NA	NA	NA	NA	NA	NA	NA	NA	0.866

clustering in open-reference OTU picking. While it is obviously problematic to use a regression model trained on 5.6 million sequences to extrapolate the runtime on 660 million sequences, we feel that this can give us an idea of the magnitude of the runtime for the serial de novo clustering of the full dataset. Our regression model projects that the serial de novo clustering of sequences that fail to hit the reference data set would require approximately 150 days to run (in wall time). In contrast, the subsampled open-reference OTU picking run presented here (which included the pre-filtering step) ran in just under

NA NA NA

NA NA

NA

NA

NA

NA

ucrss_fast_wfilter NA

 Table 4 Taxonomic profile results. Pearson correlation coefficients (r) of taxonomic summaries for (a) 88-soils at phylum level, (b) 88-soils at genus level, (c) moving-pictures at phylum level, (d) movingpictures at genus level, (e) whole-body at phylum level, and (f) whole-body at genus level.

(a)										
	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	1	0.983	1	1	1	1	0.981	1	1
ucr	NA	NA	0.983	1	1	1	1	0.981	1	1
ucrC	NA	NA	NA	0.983	0.983	0.983	0.983	0.999	0.983	0.983
ucrss	NA	NA	NA	NA	1	1	1	0.981	1	1
ucrss_wfilter	NA	NA	NA	NA	NA	1	1	0.981	1	1
uc_fast	NA	NA	NA	NA	NA	NA	1	0.981	1	1
ucr_fast	NA	NA	NA	NA	NA	NA	NA	0.981	1	1
ucrC_fast	NA	NA	NA	NA	NA	NA	NA	NA	0.981	0.981
ucrss_fast	NA	NA	NA	NA	NA	NA	NA	NA	NA	1
ucrss_fast_wfilter	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA

(b)

	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	0.939	0.85	0.939	0.939	1	0.94	0.84	0.94	0.94
ucr	NA	NA	0.821	1	1	0.94	0.998	0.923	0.998	0.998
ucrC	NA	NA	NA	0.821	0.821	0.85	0.82	0.818	0.82	0.82
ucrss	NA	NA	NA	NA	1	0.94	0.998	0.923	0.998	0.998
ucrss_wfilter	NA	NA	NA	NA	NA	0.94	0.998	0.923	0.998	0.998
uc_fast	NA	NA	NA	NA	NA	NA	0.94	0.84	0.94	0.94
ucr_fast	NA	NA	NA	NA	NA	NA	NA	0.921	1	1
ucrC_fast	NA	NA	NA	NA	NA	NA	NA	NA	0.921	0.921
ucrss_fast	NA	NA	NA	NA	NA	NA	NA	NA	NA	1
ucrss_fast_wfilter	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA

(c)

	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc₋fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	1	0.997	1	1	1	1	0.997	1	0.998
ucr	NA	NA	0.997	1	1	1	1	0.997	1	0.998
ucrC	NA	NA	NA	0.997	0.997	0.997	0.997	1	0.997	0.998
ucrss	NA	NA	NA	NA	1	1	1	0.997	1	0.998
ucrss_wfilter	NA	NA	NA	NA	NA	1	1	0.997	1	0.999
uc_fast	NA	NA	NA	NA	NA	NA	1	0.997	1	0.998
ucr_fast	NA	NA	NA	NA	NA	NA	NA	0.997	1	0.998
ucrC_fast	NA	NA	NA	NA	NA	NA	NA	NA	0.997	0.997
ucrss_fast	NA	NA	NA	NA	NA	NA	NA	NA	NA	0.998
ucrss_fast_wfilter	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA

(continued on next page)

30 days of wall time. This illustrates that while on relatively small data sets the performance enhancement of subsampled relative to classic open-reference OTU picking is either non-existence or modest (discussed in *Run-time differences*), on datasets at the current upper limit of size, the increased parallelizability of subsampled open-reference OTU picking makes open-reference OTU picking far more tractable.

Table 4 (continued)

(d)

	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	0.964	0.929	0.964	0.963	0.999	0.923	0.882	0.923	0.92
ucr	NA	NA	0.963	1	0.999	0.967	0.954	0.923	0.954	0.951
ucrC	NA	NA	NA	0.963	0.963	0.934	0.925	0.917	0.925	0.925
ucrss	NA	NA	NA	NA	0.999	0.967	0.954	0.923	0.954	0.951
ucrss_wfilter	NA	NA	NA	NA	NA	0.966	0.953	0.923	0.953	0.952
uc_fast	NA	NA	NA	NA	NA	NA	0.927	0.887	0.927	0.924
ucr_fast	NA	NA	NA	NA	NA	NA	NA	0.885	1	0.997
ucrC_fast	NA	NA	NA	NA	NA	NA	NA	NA	0.885	0.884
ucrss_fast	NA	NA	NA	NA	NA	NA	NA	NA	NA	0.997
ucrss_fast_wfilter	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
(e)										
	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	1	0.999	1	1	1	1	0.998	1	1
ucr	NA	NA	0.999	1	1	1	1	0.998	1	1
ucrC	NA	NA	NA	0.999	0.999	0.999	0.999	0.999	0.999	0.999
ucrss	NA	NA	NA	NA	1	1	1	0.998	1	1
ucrss_wfilter	NA	NA	NA	NA	NA	1	1	0.998	1	1
uc_fast	NA	NA	NA	NA	NA	NA	1	0.998	1	1

(**f**)

ucr_fast

ucrC_fast

ucrss_fast

ucrss_fast_wfilter

NA

	uc	ucr	ucrC	ucrss	ucrss_wfilter	uc_fast	ucr_fast	ucrC_fast	ucrss_fast	ucrss_fast_wfilter
uc	NA	0.959	0.9	0.959	0.959	1	0.913	0.879	0.913	0.913
ucr	NA	NA	0.918	1	1	0.957	0.967	0.871	0.967	0.967
ucrC	NA	NA	NA	0.918	0.918	0.896	0.893	0.935	0.892	0.893
ucrss	NA	NA	NA	NA	1	0.957	0.967	0.871	0.967	0.967
ucrss_wfilter	NA	NA	NA	NA	NA	0.957	0.967	0.871	0.967	0.967
uc_fast	NA	NA	NA	NA	NA	NA	0.912	0.876	0.912	0.912
ucr_fast	NA	NA	NA	NA	NA	NA	NA	0.855	1	1
ucrC_fast	NA	NA	NA	NA	NA	NA	NA	NA	0.854	0.855
ucrss_fast	NA	NA	NA	NA	NA	NA	NA	NA	NA	1
ucrss_fast_wfilter	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA

NA

NA

NA

NA

NA

NA

NA

NA

0.998

NA

NA

NA

1

0.998

NA

NA

Run-time differences

The speed improvements of subsampled open-reference OTU picking arise from the fact that a larger portion of the clustering process can be parallelized. When not run in parallel, or run in parallel over only a few (e.g., 3) CPUs, classic open-reference OTU picking is likely to be faster. Similarly, for smaller data sets (e.g., less than a few million sequences), especially if most sequences have a match in the reference database (e.g., with human gut microbiome data), classic open-reference OTU picking will achieve similar runtimes to

1

1

NA

0.998



Table 5 Runtime comparison. Comparison of runtimes (as seconds of wall time) for each method on each data set.

	88-soil	Moving-picture	Whole-body
uc	1220	27748	1095
ucr	1358	46576	1082
ucrC	226	28572	388
ucrss	1493	47207	1212
ucrss_wfilter	1885	76061	2088
uc_fast	914	23510	489
ucr_fast	1052	19371	621
ucrC_fast	44	2428	68
ucrss_fast	1021	23710	707
ucrss_fast_wfilter	1525	52811	1661

subsampled open-reference clustering (Table 5). However, in these cases, the results are still highly correlated, so if in doubt of which method will be faster, subsampled open-reference OTU picking is a reasonable choice as the summary statistics of interest (often alpha diversity, beta diversity and taxonomic profiles) are very unlikely to be different between the two methods.

When more sequences fail to hit the reference database, subsampled open-reference OTU picking becomes faster than classic open-reference OTU picking (Table 6). To illustrate this, we clustered the moving-pictures sequences against the 82% and 97% Greengenes reference OTUs at 97% identity using subsampled and classic open-reference OTU picking on 29 processors. When clustering against the 82% OTUs, 52.1 million failed to hit the reference, while when clustering against the 97% OTUs 3.4 million sequences failed to hit the reference. Subsampled open-reference OTU picking ran in 4000 s less wall

Table 6 Runtime comparisons (subsampled open-reference OTU picking variants). Comparison of runtimes (as seconds of wall time) for subsampled and "classic" open-reference OTU picking methods with variations on the default parameters.

Abbreviation	Moving-picture
ucr_fast_O29_r82	21737
ucr_fast_O29_r97	16241
ucrss_fast_O29_r82	17812
ucrss_fast_O29_r97	16169
ucrss_fast_O29_s1	14911

time than classic open-reference clustering (in a single run of each on a system dedicated for this run time comparison) against the 82% OTUs, and in 72 s less time against the 97% OTUs, illustrating that as more sequences fail to hit the reference, subsampled open-reference OTU picking offers more of an advantage. This runtime difference would be even larger if the job were split over more processors.

Another parameter that can affect runtime of subsampled open-reference OTU picking is the size of the random subsample that is selected. The optimal setting for this parameter is affected by the size of the dataset being clustered and the diversity of the sequences that fail to match the reference database. On small datasets, or datasets with a lot of novel diversity, a large fraction (e.g., 1%) is better than a small fraction (e.g., 0.001%), but as the data set increases in size a large fraction can result in far more time spent performing de novo clustering of the sequences that initially fail to hit the reference database. We recommend using the default (0.1% in QIIME 1.8.0-dev and earlier), which was chosen to reduce runtime on larger datasets where optimized runtime is more important. As this parameter setting approaches zero, subsampled open-reference OTU picking becomes more like classic open-reference OTU picking, in that more of the reads that fail to hit the reference database are clustered de novo serially, and at the limit of 0% of sequences subsampled, the subsampled open reference OTU picking becomes classic open-reference OTU picking. The summary statistics investigated here are highly correlated between classic and subsampled open-reference OTU picking, suggesting that this parameter setting will not affect those statistics, but can affect runtime.

Pre-filtering

QIIME's open-reference OTU picking workflow optionally includes a pre-filtering step, where sequences are searched against the reference database with low percent identity (the default in QIIME 1.8.0 and earlier is 60%), and sequences that fail to match are discarded from the analysis. The goal of this process is to discard sequences that are likely not representatives of the marker gene, such as host genomic sequences or products of non-specific amplification. This process is functionally similar to closed-reference OTU picking (sequence reads are searched against a pre-defined reference database), and therefore is easily run in parallel.

We show that alpha diversity (Table 2; whole-body PD Pearson r = 0.991; 88-soils PD Pearson r = 0.930; moving-pictures PD Pearson r = 0.996), beta diversity (Table 3; whole-body unweighted UniFrac Mantel r = 0.953; 88-soils unweighted UniFrac Mantel r = 0.940; moving-pictures unweighted UniFrac Mantel r = 0.990) and taxonomic summaries (Table 4; whole-body: r = 1.000 at phylum level, r = 1.000 at species level; 88-soils r = 1.000 at phylum level, r = 1.000 at species level; 0.999 at species level) are highly correlated between the pre-filtered and non-pre-filtered results, when pre-filtering is performed at percent identity of 60%. Despite nearly identical results, the pre-filtering process results in vastly increased runtimes. Consequently, we no longer recommend pre-filtering of sequences prior to open-reference OTU picking. Rather, contaminant sequences should be discarded after OTU picking. This feature is now disabled by default starting with QIIME 1.8.0-dev.

One case where pre-filtering may prove useful is in the preparation of sequence data where there is a large amount of contamination of non-marker-gene sequence, for example host genomic contamination. In this case, pre-filtering can be useful to remove those sequences prior to clustering. Note that if you suspect that your sample may contain human genomic contaminant sequences, it is important to filter them out before analysis or data deposition due to Institutional Review Board or other ethical concerns related to release of human DNA sequences.

Clustering parameters

We also investigated the effect of clustering parameters on the same summary statistics, as these can have a considerable effect on runtime. We compared uclust's default settings (referred to in QIIME as "fast mode") with the default settings in QIIME 1.8.0 and earlier ("slow mode"). We again compared the methods based on the degree to which they resulted in correlated alpha diversity (Table 2), beta diversity (Table 3), and taxonomic results (Table 4), and found that all results were highly correlated between fast and slow modes. This suggests that while fast mode will occasionally make suboptimal OTU assignments, the effects are subtle enough to be unnoticeable in downstream ecological analyses. We therefore recommend using the "fast" settings for decreased runtime, and these are now the default in QIIME 1.8.0-dev.

We do recommend using the "slow" settings if clustering sequences to build reference OTUs (for example, as is performed when building the Greengenes reference OTU collection *McDonald et al.*, 2012b) because suboptimal OTU assignments can have further reaching consequences. For example, "splitting" an OTU (i.e., defining two sequences that are within *s*% identity of each other as the centroids of two different *s*% OTUs), which is always a possibility in greedy clustering algorithms, is more common with the "fast" settings than with the "slow" settings. If this occurs in a single study, the downstream effects are limited to that study and are likely only to be problematic if the split OTU is of key significance to the system being investigated. However, a split OTU when defining reference OTUs is more problematic, because those definitions will be used in many studies, increasing the chance that the split OTU will be problematic for someone. For this application, the processing step is typically only run once per database release (which is relatively infrequent). Therefore, the longer runtime is preferable to less accurate OTU definitions in this particular application. If splitting and lumping of OTUs is of concern on your dataset, you may want to experiment with the "slow" parameter settings, which are still accessible in QIIME and we also recommend exploring the use of Oligotyping (*Eren et al., 2013*).

Consistent OTU definitions across runs: iterative open-reference OTU picking

Subsampled open-reference clustering, as implemented in QIIME, provides new identifiers for sequences that fail to match the reference database, allowing OTUs to be directly compared across clustering runs (although sequences clustered against this expanded reference sequence collection do need to be from the same gene fragment as the sequences used to expand the reference sequence collection). These OTUs can also be used in iterative OTU picking, which is useful in studies where sequence data is continuously accumulating, for example in routine monitoring of microbial communities in human subjects (e.g., patients monitored over time), the built-environment, or during environmental clean-up.

CONCLUSIONS

Taken together, the reduced runtime of subsampled open-reference OTU picking relative to classic open-reference OTU picking on large datasets, and the benefits that open-reference OTU picking offers over full *de novo* OTU picking (vastly decreased runtime) and closed-reference OTU picking (all sequences are clustered, not only those that match the reference collection), we recommend subsampled open-reference OTU picking when a reference collection is available.

Because the metrics provided here show that the same summary statistics are derived from the four OTU picking protocols, an interesting question is whether de novo or open-reference OTU picking offers any benefit over closed-reference OTU picking. The primary motivation for using methods that incorporate previously unknown OTUs (i.e., those that are not represented in the reference database) such as *de novo* and open-reference OTU picking is that OTUs not represented in the reference database might best illustrate a biological pattern of interest. For example, in the 88-soils data analyzed here, 1 of the top 10 OTUs identified as significantly different across sample pH is an OTU that is not represented in the reference database (Table 8) (this OTU was classified as in the Actinomycetales order by QIIME's uclust-based taxonomy classifier). Similarly, for the whole-body data set, 2 of the top 10 OTUs identified as significantly different across body sites were not represented in the reference database (these were classified as Prevotella melaninogenica and Veillonella parvula by QIIME's uclust-based taxonomy classifier). On the other hand, in the moving-pictures data analyzed here, all of the top 10 OTUs identified as significantly different across body site were OTUs represented in the reference database. Table 7 illustrates the fraction of OTUs not represented in the reference database by environment based on the Earth Microbiome Project dataset. We expect that using OTU picking methods that incorporate new OTUs is more important in samples where this fraction is higher.

 Table 7 OTU counts by environment. Comparison of OTUs with closed-reference and open-reference OTU picking by biome in the Earth Microbiome Project dataset.

	Average de novo OTUs (10K sequences per sample)	SD de novo OTUs (10K sequences per sample)	Average Reference OTUs (10k sequences per sample)	SD Reference OTUs (10k sequences per sample)	% novel diversity (10k seqs per sample)	% error novel diversity (10K seqs per sample)	Number of samples
Environmental Biome							
Mangrove biome	2,169	1,159	354	73	0.86	0.46	7
Tropical humid forests	2,398	260	397	35	0.858	0.094	26
Tundra biome	1,771	403	312	117	0.85	0.201	110
Deserts and xeric shrubland biome	3,917	127	707	15	0.847	0.028	7
Taiga	2,598	102	505	35	0.837	0.035	4
Marine biome	2,040	1,048	484	410	0.808	0.446	890
Aquatic biome	714	299	177	199	0.801	0.403	762
Freshwater biome	768	541	194	120	0.798	0.576	375
Warm deserts and semideserts	2,386	473	607	147	0.797	0.166	97
Tropical and subtropical moist broadleaf forest biome	3,072	125	846	18	0.784	0.032	2
Temperate needle-leaf forests or woodlands	2,836	159	785	132	0.783	0.057	21
Polar biome	1,721	886	483	218	0.781	0.414	277
Tropical and subtropical coniferous forest biome	1,993	256	579	94	0.775	0.106	3
Mixed island systems	1,552	618	511	203	0.752	0.315	124
Marginal sea	1,795	325	611	225	0.746	0.164	7
Temperate coniferous forest biome	2,504	1,206	885	201	0.739	0.361	19
Mediterranean forests, woodlands, and shrub biome	695	361	275	195	0.717	0.424	371
Large river biome	1,844	629	743	369	0.713	0.282	5
Terrestrial biome	2,714	222	1,138	163	0.705	0.072	627
Nest of bird	821	276	355	138	0.698	0.262	313
Temperate broadleaf and mixed forest biome	1,910	491	879	235	0.685	0.195	14
Temperate grasslands	2,745	290	1,315	164	0.676	0.082	696
Animal-associated habitat	758	329	376	240	0.668	0.359	1036
Mammalia-associated habitat	973	357	583	222	0.625	0.27	1918
Cold-winter (continental) deserts and semideserts	847	210	551	215	0.606	0.215	102
Temperate grasslands, savannas, and shrubland biome	1,688	272	1,497	275	0.53	0.121	85
Human-associated habitat	292	242	590	366	0.331	0.498	1597

Table 8 Significantly different OTUs by environmental metadata. Top 10 OTUs identified as significantly different across (a) binned pH in 88-soils,(b) body site in moving-pictures, and (c) body site in whole-body.

(a)		
	Taxonomy	Test-statistic
OTU		
113212	k_Bacteria;p_Acidobacteria;c_DA052;o_Ellin6513;f_;g_;s_	55.859
1123837	kBacteria;pActinobacteria;cRubrobacteria;oRubrobacterales;fRubrobacteraceae; gRubrobacter;s	50.433
New.Reference OTU22	kBacteria;pActinobacteria;cActinobacteria;oActinomycetales;f;g;s	49.172
252012	k_Bacteria;p_Proteobacteria;c_Gammaproteobacteria;o_Xanthomonadales;f_Sinobacteraceae;g_;s_	48.65
843189	kBacteria;pAcidobacteria;cSolibacteres;oSolibacterales;fSolibacteraceae; gCandidatus Solibacter;s	47.006
1127423	k_Bacteria;p_Acidobacteria;c_Acidobacteriia;o_Acidobacteriales;f_Koribacteraceae;g_;s_	43.87
1129210	k_Bacteria;p_Acidobacteria;c_Acidobacteriia;o_Acidobacteriales;f_Koribacteraceae;g_;s_	43.804
831520	kBacteria;pActinobacteria;cRubrobacteria;oRubrobacterales; fRubrobacteraceae;gRubrobacter;s	43.625
1139779	k_Bacteria;p_Proteobacteria;c_Alphaproteobacteria	41.863
804187	kBacteria;pAcidobacteria;c[Chloracidobacteria];oRB41;f;g;s	41.151

(b)

	Taxonomy	Test-statistic
OTU		
368134	k_Bacteria;p_Firmicutes;c_Bacilli;o_Bacillales;f_Staphylococcaceae;g_Staphylococcus;s_epidermidis	1599.696
3154070	k_Bacteria;p_Bacteroidetes;c_Bacteroidia;o_Bacteroidales;f_Bacteroidaceae;g_Bacteroides;s_uniformis	1625.703
1000986	k_Bacteria;p_Actinobacteria;c_Actinobacteria;o_Actinomycetales;f_Corynebacteriaceae;g_Corynebacterium;s_	1630.009
1992	k_Bacteria;p_Bacteroidetes;c_Bacteroidia;o_Bacteroidales;f_Bacteroidaceae;g_Bacteroides;s_	1728.164
4304475	k_Bacteria;p_Bacteroidetes;c_Bacteroidia;o_Bacteroidales;f_Bacteroidaceae;g_Bacteroides;s_	1545.445
191238	k_Bacteria;p_Firmicutes;c_Clostridia;o_Clostridiales;f_Lachnospiraceae;g_Coprococcus;s_	1546.436
187665	k_Bacteria;p_Firmicutes;c_Clostridia;o_Clostridiales;f_Lachnospiraceae;g_;s_	1474.529
4396297	k_Bacteria;p_Firmicutes;c_Clostridia;o_Clostridiales;f_Lachnospiraceae;g_;s_	1585.015
3903651	k_Bacteria;p_Firmicutes;c_Clostridia;o_Clostridiales;f_Ruminococcaceae;g_Oscillospira;s_	1670.188
3472078	k_Bacteria;p_Bacteroidetes;c_Bacteroidia;o_Bacteroidales;f_Bacteroidaceae;g_Bacteroides;s_	1783.488

(continued on next page)

In conclusion, this paper presents the performance-optimized subsampled openreference OTU picking algorithm, now available in QIIME. This method can be applied iteratively to define stable OTUs across sequencing runs, and achieves nearly identical results to "classic" open-reference OTU picking (i.e., not including the subsampling step). It enables massive sequencing projects such as the Earth Microbiome Project to use open-reference OTU picking in far less time than is possible with classic open-reference OTU picking, which will facilitate our exploration of microbial diversity. Further, the iterative nature of the process (which is also possible with classic open-reference OTU picking) enables progressively expanding datasets, as might be generated in clinical laboratories as microbiome-based medical treatment becomes a reality, to cluster OTUs using OTU definitions from previous clustering runs as reference sequences. This (c)

Table 8 (continued)

(0)	Taxonomy	Test-Statistic
OTU		
4326219	k_Bacteria;p_Proteobacteria;c_Epsilonproteobacteria;o_Campylobacterales; f_Campylobacteraceae; g_Campylobacter;s_	363.881
New.CleanUp. Reference OTU222	kBacteria;pBacteroidetes;cBacteroidia;oBacteroidales;fPrevotellaceae;gPrevotella; smelaninogenica	358.02
4325533	k_Bacteria;p_Bacteroidetes;c_Bacteroidia;o_Bacteroidales;f_Rikenellaceae;g_;s_	349.852
New.CleanUp. Reference OTU17550	kBacteria;pFirmicutes;cClostridia;oClostridiales;fVeillonellaceae;gVeillonella;sparvula	337.656
316732	k_Bacteria;p_Firmicutes;c_Clostridia;o_Clostridiales;f_Lachnospiraceae;g_Lachnospira;s_	337.309
4346374	k_Bacteria;p_Bacteroidetes;c_Bacteroidia;o_Bacteroidales;f_Bacteroidaceae;g_Bacteroides;s_uniformis	331.433
4458959	k_Bacteria;p_Firmicutes;c_Clostridia;o_Clostridiales;f_Veillonellaceae;g_Veillonella	329.772
3866487	k_Bacteria;p_Firmicutes;c_Clostridia;o_Clostridiales;f_Lachnospiraceae;g_Oribacterium;s_	323.488
4391641	kBacteria;pProteobacteria;cGammaproteobacteria;oPasteurellales; fPasteurellaceae;gHaemophilus; sparainfluenzae	312
175751	k_Bacteria;p_Firmicutes;c_Clostridia;o_Clostridiales;f_Lachnospiraceae;g_;s_	305.531

avoids re-clustering all sequences every time new sequences are generated, thereby vastly decreasing computational costs.

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ADDITIONAL INFORMATION AND DECLARATIONS

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Competing Interests

The authors declare there are no competing interests.

Author Contributions

- Jai Ram Rideout performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, reviewed drafts of the paper.
- Yan He performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, wrote the paper, prepared figures and/or tables, reviewed drafts of the paper.
- Jose A. Navas-Molina and Luke K. Ursell performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, wrote the paper, reviewed drafts of the paper.
- William A. Walters, Sean M. Gibbons, John Chase, Daniel McDonald, Antonio Gonzalez and Adam Robbins-Pianka performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, reviewed drafts of the paper.
- Jose C. Clemente conceived and designed the experiments, contributed reagents/materials/analysis tools, prepared figures and/or tables, reviewed drafts of the paper.
- Jack A. Gilbert, Susan M. Huse and Hong-Wei Zhou conceived and designed the experiments, contributed reagents/materials/analysis tools, reviewed drafts of the paper.
- Rob Knight conceived and designed the experiments, contributed reagents/materials/analysis tools, wrote the paper, reviewed drafts of the paper.
- J. Gregory Caporaso conceived and designed the experiments, performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, wrote the paper, prepared figures and/or tables, reviewed drafts of the paper.

Data Deposition

The following information was supplied regarding the deposition of related data: https://github.com/gregcaporaso/cloaked-octo-ninja.

QIIME Database:

whole-body: ftp://thebeast.colorado.edu/pub/QIIME_DB_Public_Studies/study_449_split_library_seqs_and_mapping.zip

moving-pictures: ftp://thebeast.colorado.edu/pub/QIIME_DB_Public_Studies/study_ 550_split_library_seqs_and_mapping.zip

88-soils: ftp://thebeast.colorado.edu/pub/QIIME_DB_Public_Studies/study_103_split_library_seqs_and_mapping.zip.

Supplemental Information

Supplemental information for this article can be found online at http://dx.doi.org/ 10.7717/peerj.545#supplemental-information.

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