## Application of database-independent methods to assess the quality of OTU picking methods

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## Abstract

- 2 Assigning 16S rRNA gene sequences to operational taxonomic units (OTUs) allows microbial
- ecologists to overcome the inconsistencies and biases within bacterial taxonomy and provides a
- 4 strategy for clustering similar sequences that do not have representatives in a reference database. I
- 5 have applied the Matthew's correlation coefficient to assess the ability of 15 reference-independent
- and -dependent clustering algorithms to assign sequences to OTUs. This metric quantifies the
- ability of an algorithm to reflect the relationships between sequences without the use of a reference
- and can be applied to any dataset or method. The most consistently robust method was the average
- neighbor algorithm; however, for some datasets other algorithms matched its performance.

Numerous algorithms have been developed for solving the seemingly simple problem of assigning
16S rRNA gene sequences to operational taxonomic units (OTUs). These algorithms were recently
the subject of benchmarking studies performed by Westcott and myself (1, 2), He et al (3), and
Kopylova et al (4). These studies provide a thorough review of the sequencing clustering landscape,
which can be divided into three general approaches: (i) *de novo* clustering where sequences are
clustered without first mapping sequences to a reference database, (ii) closed-reference clustering
where sequences are clustered based on the references that the sequences map to, and (iii)
open reference clustering where sequences that do not map adequately to the reference are then
clustered using a *de novo* approach. Assessing the quality of the clustering assignments has been
a persistent problem in the development of clustering algorithms.

The recent analysis of Kopylova et al (4) repeated many of the benchmarking strategies employed by previous researchers. Many algorithm developers have clustered sequences from simulated 21 communities or sequencing data from synthetic communities of cultured organisms and quantify 22 how well the OTU assignments matched the organisms' taxonomy (5-16). This approach is 23 flawed because bacterial taxonomy often reflects those inconsistencies and biases within bacterial taxonomy that OTU-based methods strive to overcome. Furthermore, it is unclear how the methods 25 scale to sequences from the novel organisms we are likely to encounter in deep sequencing surveys. In a second approach, developers have compared the time and memory required to cluster sequences in a dataset (6, 13, 17, 18). These are valid parameters to assess when judging a clustering method, but indicate little regarding the quality of the OTU assignments. One could randomly assign sequences to a predetermined number of OTUs. This would be efficient. It would also poorly reflect the genetic diversity within the community. In a third approach, developers have compared the number of OTUs generated by various methods for a common dataset (4, 5). Again, 32 one could randomly cluster sequences into a target number of OTUs, but the clusterings would likely be meaningless. In a fourth approach a metric of OTU stability has been proposed as a way to assess algorithms (3). Although it is important that the methods generate reproducible OTU assignments when the initial order of the sequences is randomized, this metric ignores the possibility that the variation in assignments may be equally robust or that the assignments by a highly reproducible algorithm may be quite poor. In a final approach, some developers have

assessed the quality of clustering based on the method's ability to generate the same OTUs
generated by other methods (18, 19). Unfortunately, without the ability to ground truth any method,
such comparisons are tenuous. There is a need for an objective metric to assess the quality of
OTU assignments.

Westcott and I have proposed an unbiased and objective method for assessing the quality of OTU assignments that can be applied to any collection of sequences (1, 2). Our approach uses the observed dissimilarity between pairs of sequences and information about whether sequences were 45 clustered together to quantify how well similar sequences are clustered together and dissimilar sequences are clustered apart. To quantify the correlation between the observed and expected OTU assignments, we synthesize the relationship between OTU assignments and the distances between sequences using the Matthew's correlation coefficient (20). I have expanded our previous analysis to evaluate three hierarchical and seven greedy de novo algorithms, one open-reference clustering algorithm, and four closed-reference algorithms (Figure 1). To test these approaches 51 I applied each of them to datasets from soil (21), mouse feces (22), and two simulated datasets. 52 The simulated communities were generated by randomly selecting 10,000 16S rRNA sequences that were unique within the V4 region from the SILVA non-redundant database (4, 23). Next, 54 an even community was generated by specifying that each sequence had a frequency of 100 55 reads and a staggered community was generated by specifying that the abundance of each sequence was a randomly drawn a uniform distribution between 1 and 200. A reproducible 57 version of this manuscript and analysis has been added to the repository available at https: 58 //github.com/SchlossLab/Schloss Cluster PeerJ 2015.

I replicated the benchmarking approach that I have used previously to assess the ability of an algorithm to correctly group sequences that are similar to each other and split sequences that are dissimilar to each other using the MCC (1, 2). When I compared the MCC values calculated using the ten *de novo* algorithms with the four datasets, the average neighbor algorithm reliably performed as well or better than the other methods (Figure 1). For the murine dataset, the MCC values for the VSEARCH (AGC: 0.76 and DGC: 0.78) and USEARCH-based (AGC: 0.76 and DGC: 0.77) algorithms, Sumaclust (0.76), and average neighbor (0.76) were similarly high. For each of the other datasets, the MCC value for the average neighbor algorithm was at least 5%

higher than the next best method. Swarm does not use a traditional distance-based criteria to cluster sequences into OTUs and instead looks for natural subnetworks in the data. When I used the distance threshold that gave the best MCC value for the Swarm data, the MCC values were generally not as high as they were using the average neighbor algorithm. The one exception was for the soil dataset. Among the reference-based methods, all of the MCC values suffer because when sequences that are at least 97% similar to a reference are pooled, the sequences within an OTU could be as much as 6% different from each other. The effect of this is observed in the MCC values that were calculated for the OTUs assigned by these methods generally being lower than those observed using the *de novo* approaches (Figure 1). It is also important to note that the MCC values for the closed-reference OTUs are inflated because sequences were removed from the analysis if there was not a reference sequence that was more than 97% similar to the sequence. Given the consistent quality of the clusterings formed by the average neighbor algorithm, these results confirm the conclusion from the previous analysis that researchers should use the average 80 neighbor algorithm or calculate MCC values for several methods and use the clustering that gives 81 the best MCC value (2).

Next, I investigated the ability of the reference-based methods to properly assign sequences 83 to OTUs. The full-length 16S rRNA gene sequences in the default reference taxonomy that accompanies QIIME are less than 97% similar to each other. Within the V4 region, however, many of the sequences were more similar to each other and even identical to each other. As a result, we previously found that there was a dependence between the ordering of sequences in the reference 87 database and the OTU assignments with USEARCH and VSEARCH (2). To explore this further, we analyzed the 32,106 unique sequences from the murine dataset with randomized databases. VSEARCH always found matches for 27,737 murine sequences, the reference matched to those 90 sequences differed between randomizations. For USEARCH there were between 28,007 and 91 28,111 matches depending on the order of the reference. In the updated analysis we found that SortMeRNA resulted in between 23,912 and 28,464 matches. Using NINJA-OPS with different 93 orderings of the reference sequences generated the same 28,499 matches. These results point 94 to an additional problem with closed-reference clustering, which is the inability for the method to assign sequences to OTUs when a similar reference sequence does not exist in the database. For

the well-characterized murine microbiota, NINJA-OPS did the best by finding relatives for 88.8% of the unique murine sequences. As indicated by the variation in the number of sequences that matched a reference sequence, these methods varied in their sensitivity and specificity to find the best reference sequence. Of the closed-reference methods, NINJA-OPS had the best sensitivity (99.7%) and specificity (79.7%) while SortMeRNA had the worst sensitivity (95.7%) and VSEARCH had the worst specificity (60.3%). Reference-based clustering algorithms are much faster than *de novo* approaches, but do not generate OTUs that are as robust.

Although the goal of Kopylova et al (4) was to compare various clustering algorithms, they also 104 studied these algorithms in the broader context of raw sequence processing, screening for chimeras, 105 and removal of singletons. Each of these are critical decisions in a comprehensive pipeline. By 106 including these steps, they confounded their analysis of how best to cluster sequences into OTUs. 107 The effect of differences in MCC values on one's ability to draw inferences is unclear and admittedly may be relatively minor for some datasets. Because of this uncertainty, researchers should use 109 the most reliable methods available in case the differences in clustering do effect the conclusions 110 that can be drawn from a particular dataset. Through the use of objective criteria that measure the quality of the clusterings, independent of taxonomy or database, researchers will be able to 112 evaluate which clustering algorithm is the best fit for their data. 113

Figure 1. Comparison of OTU quality generated by multiple algorithms applied to four 114 datasets. The nearest, average, and furthest neighbor clustering algorithms were used as implemented in mothur (v.1.37)(24). Abundance (AGC) and Distance-based greedy clustering 116 (DGC) were implemented using USEARCH (v.6.1) and VSEARCH (v.1.5.0)(3, 5, 25). Other de 117 novo clustering algorithms included Swarm (v.2.1.1)(6, 7), OTUCLUST (v.0.1)(26), and Sumaclust 118 (v.1.0.20). The MCC values for Swarm were determined by selecting the distance threshold that 119 generated the maximum MCC value for each dataset. The USEARCH and SortMeRNA (v.2.0) 120 closed-reference clusterings were performed using QIIME (v.1.9.1) (27, 28). Closed-reference 121 clustering was also performed using VSEARCH (v.1.5.0) and NINJA-OPS (v.1.3.2) (16). The order 122 of the sequences in each dataset was randomized thirty times and the intra-method range in MCC 123 values was smaller than the plotting symbol. MCC values were calculated using mothur.

## 25 References

- 1. **Schloss PD**, **Westcott SL**. 2011. Assessing and improving methods used in operational taxonomic unit-based approaches for 16S rRNA gene sequence analysis. Applied and Environmental Microbiology **77**:3219–3226. doi:http://doi.org/10.1128/aem.02810-10.
- 2. Westcott SL, Schloss PD. 2015. De novo clustering methods outperform reference-based
   methods for assigning 16S rRNA gene sequences to operational taxonomic units. PeerJ 3:e1487.
   doi:http://doi.org/10.7717/peerj.1487.
- 3. He Y, Caporaso JG, Jiang X-T, Sheng H-F, Huse SM, Rideout JR, Edgar RC, Kopylova E,
  Walters WA, Knight R, Zhou H-W. 2015. Stability of operational taxonomic units: An important
  but neglected property for analyzing microbial diversity. Microbiome 3. doi:http://doi.org/10.1186/
  s40168-015-0081-x.
- 4. Kopylova E, Navas-Molina JA, Mercier C, Xu ZZ, Mahé F, He Y, Zhou H-W, Rognes T,
  Caporaso JG, Knight R. 2016. Open-source sequence clustering methods improve the state of
  the art. mSystems 1:e00003–15. doi:http://doi.org/10.1128/msystems.00003-15.
- 5. Edgar RC. 2013. UPARSE: Highly accurate OTU sequences from microbial amplicon reads.
   Nature Methods 10:996–998. doi:http://doi.org/10.1038/nmeth.2604.
- 6. Mahé F, Rognes T, Quince C, Vargas C de, Dunthorn M. 2014. Swarm: Robust and fast clustering method for amplicon-based studies. PeerJ 2:e593. doi:http://doi.org/10.7717/peerj.593.
- 7. **Mahé F**, **Rognes T**, **Quince C**, **Vargas C de**, **Dunthorn M**. 2015. Swarm v2: Highly-scalable and high-resolution amplicon clustering. PeerJ **3**:e1420. doi:http://doi.org/10.7717/peerj.1420.
- 8. **Barriuso J**, **Valverde JR**, **Mellado RP**. 2011. Estimation of bacterial diversity using next generation sequencing of 16S rDNA: A comparison of different workflows. BMC Bioinformatics **12**:473. doi:http://doi.org/10.1186/1471-2105-12-473.
- 9. **Bonder MJ**, **Abeln S**, **Zaura E**, **Brandt BW**. 2012. Comparing clustering and pre-processing in taxonomy analysis. Bioinformatics **28**:2891–2897. doi:http://doi.org/10.1093/bioinformatics/bts552.

- 150 10. **Chen W**, **Zhang CK**, **Cheng Y**, **Zhang S**, **Zhao H**. 2013. A comparison of methods for clustering 16S rRNA sequences into OTUs. PLoS ONE **8**:e70837. doi:http://doi.org/10.1371/journal. pone.0070837.
- 11. **Huse SM**, **Welch DM**, **Morrison HG**, **Sogin ML**. 2010. Ironing out the wrinkles in the rare biosphere through improved OTU clustering. Environmental Microbiology **12**:1889–1898. doi:http: //doi.org/10.1111/j.1462-2920.2010.02193.x.
- 12. **May A**, **Abeln S**, **Crielaard W**, **Heringa J**, **Brandt BW**. 2014. Unraveling the outcome of 16S rDNA-based taxonomy analysis through mock data and simulations. Bioinformatics **30**:1530–1538. doi:http://doi.org/10.1093/bioinformatics/btu085.
- 13. **Cai Y**, **Sun Y**. 2011. ESPRIT-tree: Hierarchical clustering analysis of millions of 16S rRNA pyrosequences in quasilinear computational time. Nucleic Acids Research **39**:e95–e95. doi:http://doi.org/10.1093/nar/gkr349.
- 14. Sun Y, Cai Y, Huse SM, Knight R, Farmerie WG, Wang X, Mai V. 2011. A large-scale benchmark study of existing algorithms for taxonomy-independent microbial community analysis.

  Briefings in Bioinformatics 13:107–121. doi:http://doi.org/10.1093/bib/bbr009.
- 15. White JR, Navlakha S, Nagarajan N, Ghodsi M-R, Kingsford C, Pop M. 2010. Alignment and
   clustering of phylogenetic markers implications for microbial diversity studies. BMC Bioinformatics
   11:152. doi:http://doi.org/10.1186/1471-2105-11-152.
- 16. **Al-Ghalith GA**, **Montassier E**, **Ward HN**, **Knights D**. 2016. NINJA-OPS: Fast accurate marker gene alignment using concatenated ribosomes. PLOS Computational Biology **12**:e1004658. doi:http://doi.org/10.1371/journal.pcbi.1004658.
- 17. Sun Y, Cai Y, Liu L, Yu F, Farrell ML, McKendree W, Farmerie W. 2009. ESPRIT: Estimating species richness using large collections of 16S rRNA pyrosequences. Nucleic Acids Research 37:e76–e76. doi:http://doi.org/10.1093/nar/gkp285.
- 18. Rideout JR, He Y, Navas-Molina JA, Walters WA, Ursell LK, Gibbons SM, Chase J,
  McDonald D, Gonzalez A, Robbins-Pianka A, Clemente JC, Gilbert JA, Huse SM, Zhou

- H-W, Knight R, Caporaso JG. 2014. Subsampled open-reference clustering creates consistent, comprehensive OTU definitions and scales to billions of sequences. PeerJ 2:e545. doi:http://doi.org/10.7717/peerj.545.
- 19. **Schmidt TSB**, **Rodrigues JFM**, **Mering C von**. 2014. Limits to robustness and reproducibility in the demarcation of operational taxonomic units. Environ Microbiol **17**:1689–1706. doi:http://doi.org/10.1111/1462-2920.12610.
- 20. **Matthews B**. 1975. Comparison of the predicted and observed secondary structure of t4 phage lysozyme. Biochimica et Biophysica Acta (BBA) Protein Structure **405**:442–451. doi:http://doi.org/10.1016/0005-2795(75)90109-9.
- 21. Roesch LFW, Fulthorpe RR, Riva A, Casella G, Hadwin AKM, Kent AD, Daroub SH,
  Camargo FAO, Farmerie WG, Triplett EW. 2007. Pyrosequencing enumerates and contrasts soil
  microbial diversity. The ISME Journal. doi:http://doi.org/10.1038/ismej.2007.53.
- Schloss PD, Schubert AM, Zackular JP, Iverson KD, Young VB, Petrosino JF. 2012.
   Stabilization of the murine gut microbiome following weaning. Gut Microbes 3:383–393. doi:http://doi.org/10.4161/gmic.21008.
- Pruesse E, Quast C, Knittel K, Fuchs BM, Ludwig W, Peplies J, Glockner FO. 2007.
   SILVA: A comprehensive online resource for quality checked and aligned ribosomal RNA sequence
   data compatible with ARB. Nucleic Acids Research 35:7188–7196. doi:http://doi.org/10.1093/nar/
   gkm864.
- 24. Schloss PD, Westcott SL, Ryabin T, Hall JR, Hartmann M, Hollister EB, Lesniewski RA,
  Oakley BB, Parks DH, Robinson CJ, Sahl JW, Stres B, Thallinger GG, Horn DJV, Weber CF.
  2009. Introducing mothur: Open-source, platform-independent, community-supported software
  for describing and comparing microbial communities. Applied and Environmental Microbiology
  75:7537–7541. doi:http://doi.org/10.1128/aem.01541-09.
- 25. Rognes T, Mahé F, Flouri T, McDonald; D. 2015. Vsearch: VSEARCH 1.4.0. Zenodo.
- 201 26. Albanese D, Fontana P, Filippo CD, Cavalieri D, Donati C. 2015. MICCA: A complete

- and accurate software for taxonomic profiling of metagenomic data. Sci Rep **5**:9743. doi:http: //doi.org/10.1038/srep09743.
- 27. **Kopylova E**, **Noe L**, **Touzet H**. 2012. SortMeRNA: Fast and accurate filtering of ribosomal RNAs in metatranscriptomic data. Bioinformatics **28**:3211–3217. doi:http://doi.org/10.1093/
- 28. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, Fierer N, Peña AG, Goodrich JK, Gordon JI, Huttley GA, Kelley ST, Knights D, Koenig JE, Ley RE, Lozupone CA, McDonald D, Muegge BD, Pirrung M, Reeder J, Sevinsky JR, Turnbaugh PJ, Walters WA, Widmann J, Yatsunenko T, Zaneveld J, Knight R. 2010. QIIME allows analysis of high-throughput community sequencing data. Nature Methods 7:335–336. doi:http://doi.org/10. 1038/nmeth.f.303.