Reference-based OTU clustering for machine learning classification

Other Title Options:

A solution to the problem of inconsistent OTU clustering for machine learning classification

Optimized reference-based OTU clustering for machine learning classification

Running title: INSERT RUNNING TITLE HERE

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observation format (max 1200 words, 2 figures, 25 ref)

Abstract (250 word max)

Machine learning classification of disease based on the gut microbiome often relies on operational taxonomic units (OTUs) to quantify microbial composition. The standard methodology for quantifying taxonomic composition is clustering 16S rRNA amplicon sequences de novo into OTUs and using their abundances to train a classification model. However, OTU clusters depend on the sequences in the data set and therefore can change if new data is added. This inconsistency complicates machine learning classification because in order to use the model to classify additional samples, the OTUs have to be re-clustered to include the new sequences and the model retrained with the new OTU clusters. A new algorithm in mothur, called OptiFit, addresses this issue by clustering new sequences into the existing OTU clusters. While OptiFit is proven to produce high quality OTU clusters, it is unclear whether this method for clustering new sequence data into existing OTUs will have any impact on machine learning classification using the OTU abundances. To evaluate the potential for use in machine learning classification, we used 12 OptiFit to cluster additional data into existing OTU clusters and quantified model performance in classifying a data set containing samples from controls and people with advanced adenoma and carcinoma (SRN). We compared the model performance using OptiFit to the standard procedure of clustering all the data together. We found that OptiFit can be used to fit new sequence data to existing OTUs and perform equally in classifying SRNs. Moving forward, when OTUs are used in machine learning classification problems, 17 OptiFit can be used to avoid the need to re-cluster sequences into OTUs when classifying additional samples that were not part of the data set used to build the model.

20 Importance (150 word max)

There are instances when OTU abundance data is optimal for machine learning prediction of disease based on microbiome composition. The current methodology for OTU clustering complicates machine learning since *de novo* OTU clusters can change depending on the data in the data set. Attempting to classify additional samples that were not part of the original model build requires re-clustering OTUs and re-training the model. OptiFit provides an elegant solution to the problem of OTU inconsistency by enabling new data to be fit to the existing OTUs while maintaining machine learning classification performance. The use of OptiFit will enable a single model based on OTU abundance data to be used in classifying additional samples that were not part of the original OTU clustering.

Gut microbiome community composition is useful as a resource for machine learning prediction of diseases. including colorectal cancer (1, 2). Amplicon sequencing of the 16S rRNA gene is a reliable tool for assessing the taxonomic composition of microbial communities. Analysis of 16S rRNA amplicon sequence (16S) data generally relies on clustering of sequences based on similarity into operational taxonomic units (OTUs). The process of OTU clustering can either be reference-based or de novo. The quality of OTUs generated with reference-based clustering is dependent on the quality of the reference used, while OTUs generated with de novo clustering are optimized to the data set (3). While de novo clustering can produce high-quality OTU clusters where sequences are accurately grouped based on similarity thresholds, the resulting OTU clusters depend on the data in the data set and the addition of new data could change the overall OTU clusters. The inconsistent nature of OTU clustering complicates deployment of machine learning models since integration of additional data requires re-clustering all the data and re-training of the model. The ability to integrate new data into an existing model without re-clustering and re-training could allow for deployment of a single model that new data can be continually added to. Recently Sovacool et al introduced OptiFit: a method for fitting new data into existing OTU clusters (4). While OptiFit is proven to effectively fit new sequence data to existing OTU clusters, it is unknown if the use of OptiFit will have an impact on machine learning classification. Here we test the ability of OptiFit to cluster new sequence data into existing OTU clusters for the purpose of machine learning classification of disease based on gut microbiome composition. When using OTU abundances for machine learning classification, the current methodology is to de novo cluster all of the sequence data into OTUs with the OptiClust algorithm in mothur (5). The resulting abundance data is then split into training and testing sets, where the training set is used to tune hyperparameters and ultimately train the model. The testing set is then classified with the model and the performance of the model can be quantified (Figure 1A). However, with this methodology we would have to re-generate the OTU clusters and re-train the model if we wanted to classify additional samples. The OptiFit algorithm (4) addresses this problem by enabling new sequences to be clustered into existing OTUs. The OptiFit workflow is similar to the OptiClust workflow where the data is clustered into OTUs and used to tune hyperparameters and ultimately train the model. Then, we can use OptiFit to fit sequence data of samples not part of the original data set into the existing OTUs and use the same model to classify the samples (Figure 1B). To test how the model performance compares between these two methodologies, we used a publicly available data set of 16S sequences from stool samples of healthy subjects as well as subjects with SRN consisting of advanced adenoma and carcinoma (1). The data set was randomly split into an 80% train set and 20% test set. For the standard OptiClust workflow, the training and test sets were de novo clustered together into OTUs then the resulting abundance table was split into the training and testing set. For the OptiFit workflow, the train set was clustered *de novo* into OTUs and the remaining test
set was fit to the OTU clusters using the OptiFit algorithm. For both workflows, the abundance table of the
train set was used to tune hyperparameters and train a random forest model to classify SRN. The test set
was classified as either control or SRN using the trained models. To account for variation depending on the
split of the data, the data set was randomly split 100 times and the process repeated for each of the 100
data splits. By comparing the model performance of classifying the samples in the test data set between
the OptiFit and OptiClust algorithms, we can quantify the impact of using OptiFit on model classification
performance.

We first examined the quality of the resulting OTU clusters from the two algorithms using the Matthews correlation coefficient (MCC). The MCC score is quantified by examining all pairs of sequences and assessing whether they belong together in an OTU or not based on sequence similarity (5). MCC scores range between negative one and one. A score of negative one means none of the sequences in an OTU are within the similarity threshold and any sequences within the similarity threshold are not in an OTU together. An MCC score of zero essentially means the sequences are randomly clustered. An MCC score of 1 means all sequences in an OTU are within the similarity threshold and all sequence pairs within the similarity threshold are in the same OTU. To ensure that OptiFit is appropriately integrating new sequence data into the existing OTUs, we would expect that the MCC scores produced by the OptiClust and OptiFit workflows are similar. Since the data is only clustered once in the OptiClust workflow there is only one MCC score while the OptiFit workflow produces an MCC score for the OTU clusters from each data split.

Overall the MCC scores were similar between OptiClust (MCC = 0.884) and OptiFit (average MCC = 0.879) indicating that OptiFit performs as well as OptiClust when integrating new sequences into the existing OTUs.

After verifying that the quality of the OTUs was consistent between OptiClust and OptiFit, we next examined
the model performance for classifying samples in the test data set as control or SRN. To quantify model
performance we used the taxonomic abundances of the training data from the OptiClust and OptiFit
workflows to train a model to predict SRNs. Using the predicted and actual diagnosis classification,
we calculated the area under the receiver operating characteristic curve (AUROC) for each data split to
quantify model performance. During cross-validation (CV) training, the model performance was equivalent
between the two algorithms (p-value = 0.13, OptiClust mean CV AUROC = 0.694, OptiFit mean CV AUROC
= 0.697, Figure 2A). The trained model was then deployed to classify the samples of the test data as
control or SRN. The performance on the test data was equivalent between the two algorithms (p-value =
0.63, OptiClust mean test AUROC = 0.709, OptiFit mean test AUROC = 0.712, Figure 2B,C) indicating that

93 new data can be fit to existing OTU clusters without impacting model performance.

We tested the ability of OptiFit to integrate new data into existing OTUs for the purpose of machine learning classification of disease based on microbiome abundance. A potential problem to using OptiFit for machine learning prediction is that any sequences in the new data that do not map to the existing OTU clusters will be discarded resulting in a possible loss of information. However, we demonstrated that OptiFit can be used to fit new sequence data into existing OTU clusters and perform equally in predicting SRN compared to clustering all of the sequence data together. The ability to integrate new data into existing OTUs enables the deployment of a single machine learning model based on microbiome composition that new data can be classified with. These results are based on a single data set and disease. Further analysis is needed to determine the data size necessary to build a robust model capable of classifying diverse data sets. A robust machine learning model could be implemented as part of a non-invasive and low-cost aid in diagnosing SRN.

Materials and Methods

Data Set. Raw 16S rRNA amplicon sequence data isolated from human stool samples was downloaded from NCBI Sequence Read Archive (accession no. SRP062005) (1).; (6) This data set contains stool samples from a total of N subjects, however after preprocessing to screen for sequence quality and subsample to 10,000 reads per sample, 490 samples remained. For this analysis, samples from subjects identified in the metadata as normal, high risk normal, or adenoma were categorized as "normal" while samples from subjects identified as advanced adenoma or carcinoma were categorized as "screen relevant neoplasia" (SRN). The resulting data set consisted of 261 normal samples and 229 SRN samples.

Data Processing. The full dataset was pre-processed with mothur (v1.45) (7) using the SILVA reference database (v132) (8) to join forward and reverse reads, merge duplicate reads, align to the reference, pre-cluster, remove chimeras with UCHIME (6), assign taxonomy, and remove non-bacterial reads following the Schloss Lab MiSeq standard operating procedure described on the mothur website (https://mothur.org/wiki/miseq_sop/). 100 splits of the 490 samples were generated where 80% of the samples (392 samples) were randomly assigned to the training set and the remaining 20% (98 samples) were assigned to the test set. Using 100 splits of the data accounts for the variation that may be observed depending on the samples that are in the training or test sets. Each samples was in the training set an average of 80 times (SD=4.1) and the test set an average of 20 times (SD=4.1).

22 The data was processed through two workflows. First, the standard workflow using the OptiClust algorithm

(5). In this pathway, all of the data was clustered together with OptiClust to generate OTUs and the resulting abundance tables were split into the training and testing sets. In the second workflow, the pre-processed data was split into the training and testing sets. The training set was clustered into OTUs, then the test set was fit to the OTUs of the training set using the OptiFit algorithm (4). The OptiFit algorithm was run with method open so that any sequences that didn't map to the existing OTU clusters would form new OTUs. Any OTUs that were not in the training set were removed prior to machine learning. For both pathways, the shared files were sub-sampled to 10,000 reads per sample.

Machine Learning. Machine learning using Random Forest was conducted with the R package mikrompl (v XXXX) (9) to predict the diagnosis (SRN or normal) for the samples in the test set for each data split.

The training set was preprocessed to normalize values (scale/center), collapse correlated features, and remove features with zero-variance. The preprocessing from the training set was then applied to the test set. P values comparing model performance were calculated as previously described {}. The averaged ROC curves were plotted by taking the average and standard deviation of the sensitivity at each specificity value.

Code Availability. The analysis workflow was implemented in Snakemake (10). Scripts for analysis were written in R (11) and GNU bash (12). The software used includes mothur v1.47.0 (7), RStudio (13), the Tidyverse metapackage (14), R Markdown (15), the SRA toolkit (16), and conda (17). The complete workflow and supporting files required to reproduce this study are available at: https://github.

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179 Figures

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H = hyperparameter settings

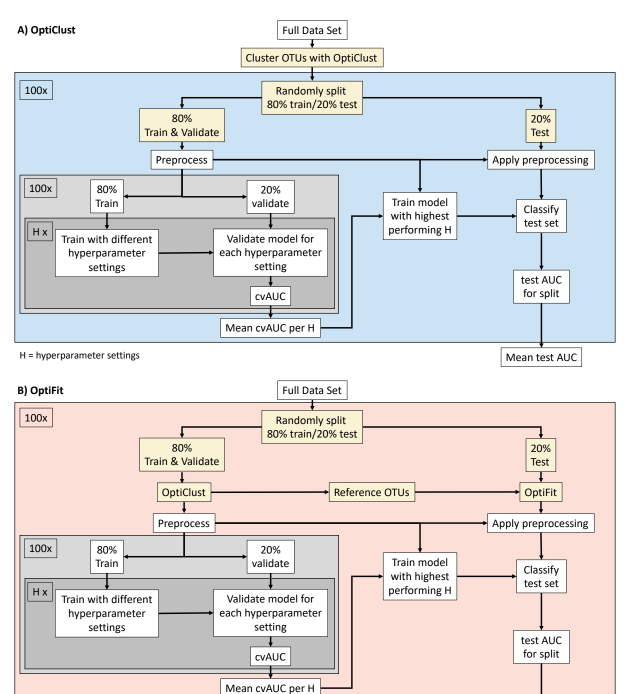


Figure 1: Workflows. A) OptiClust workflow: The full data set was clustered into OTUs using the OptiClust algorithm in mothur. The data was then split into two sets where 80% of the samples were assigned to the training set and 20% to the testing set. The training set was preprocessed with mikropml to normalize values (scale/center), collapse correlated features, and remove features with zero-variance. Using mikropml, the

Mean test AUC

training set was split into train and validate sets to compare results using different hyperparameter settings. The highest performing hyperparameter setting was then used to train the model with the full training set. The preprocessing scale from the training set was applied to the test data set, then the trained model was 188 used to classify the samples in the test set. Based on the actual classification and predicted classification, the area under the receiver operating characteristic curve (AUROC) was calculated to summarize model 190 performance. The entire process was repeated 100 times to account for variability depending on the split of the data resulting in a total of 100 AUROC values summarizing the performance of the standard OptiClust 192 workflow. B) OptiFit workflow: The data set was first split into two sets where 80% of the samples were 193 assigned to the training set and 20% to the testing set. The training set was then clustered into OTUs 194 using the OptiClust algorithm in mothur. The resulting abundance data was preprocessed with mikropml 195 to normalize values (scale/center), collapse correlated features, and remove features with zero-variance. Using mikropml, the training set was split into train and validate sets to compare results using different 197 hyperparameter settings. The highest performing hyperparameter setting was then used to train the model with the full training set. The OptiFit algorithm in mothur was used to cluster the left out testing data set 199 using the OTUs of the training set as a reference. The preprocessing scale from the training set was 200 applied to the test data set, then the trained model was used to classify the samples in the test set. Based 201 on the actual classification and predicted classification, the area under the receiver operating characteristic 202 curve (AUROC) was calculated to summarize model performance. The entire process was repeated 100 203 times to account for variability depending on the split of the data resulting in a total of 100 AUROC values summarizing the performance of the new OptiFit workflow.

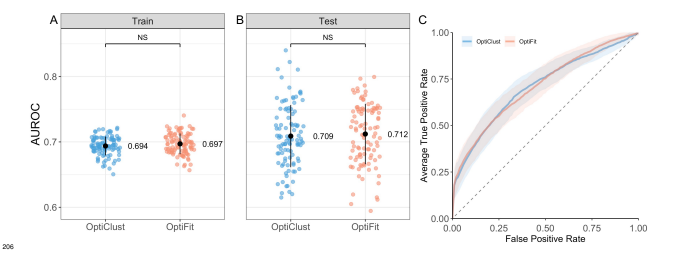


Figure 2: Model Performance. A) Area under the receiver operating characteristic (AUROC) curve during cross-validation for the OptiClust and OptiFit workflows. Mean and standard deviation of the AUROC is represented by the black dot and whiskers. Mean AUROC is printed to the right of the points. **B)** AUROC on the test data for the OptiClust and OptiFit workflows. Mean and standard deviation of the AUROC is represented by the black dot and whiskers. Mean AUROC is printed to the right of the points. **C)** Receiver operating characteristic (ROC)Averaged ROC curves