**Best practices for applying machine learning to bacterial 16S rRNA gene sequencing data**

Running title: Machine learning methods in microbiome studies

Begüm D. Topçuoğlu, Nicholas A. Lesniak, Jenna Wiens, Mack Ruffin, Patrick D. Schloss1

To whom correspondence should be addressed: [pschloss@umich.edu](mailto:pschloss@umich.edu)

1. Department of Microbiology and Immunology, University of Michigan, Ann Arbor, MI 48109

2. Department of Computer Science and Engineering, University or Michigan, Ann Arbor, MI 49109

3. Department of Family Medicine and Community Medicine, Penn State Hershey Medical Center, Hershey, PA

## Abstract

Machine learning (ML) modeling of the human microbiome has the potential to identify the microbial biomarkers and aid in diagnosis of many chronic diseases such as inflammatory bowel disease, diabetes and colorectal cancer. Progress has been made towards developing ML models that predict health outcomes from bacterial abundances, but rigourous ML models are scarce due to the flawed methods that call the validity of developed models into question. Furthermore, the use of black box ML models has hindered the validation of microbial biomarkers. To overcome these challenges, we benchmarked seven different ML models that use fecal 16S rRNA sequences to predict colorectal cancer (CRC) lesions (n=490 patients, 261 controls and 229 cases). To show the effect of model selection, we assessed the predictive performance, interpretability, and computational efficiency of the following models: L2-regularized logistic regression, L1 and L2-regularized support vector machines (SVM) with linear and radial basis function kernels, a decision tree, random forest, and extreme gradient boosting (XGBoost). The random forest model was best at detecting CRC lesions with an AUROC of 0.695 but it was slow to train (83.2 h) and hard to interpret. Despite its simplicity, L2-regularized logistic regression followed random forest in predictive performance with an AUROC of 0.680, and it trained much faster (12 min). In this study, we established standards for the development of modeling pipelines for microbiome-associated ML models. Additionally, we showed that ML models should be chosen based on expectations of predictive performance, interpretability and available computational resources.

## Importance (needs work)

Prediction of health outcomes using ML is rapidly being adopted by human microbiome studies. However, the developed ML models so far are overoptimistic in terms of validity and predictive performance. Without rigorous ML pipelines, we cannot trust ML models. Before we can speed up progress, we need to slow down, define and implement good ML practices.

## Background

As the number of people represented in human microbiome datasets grow, there is an increasing desire to use microbiome data to diagnose diseases. However, the structure of the human microbiome is remarkably variable between individuals to the point where it is often difficult to identify the bacterial populations that are associated with diseases using traditional statistical models. This variation is likely due to the ability of many bacterial populations to fill the same niche such that different populations cause the same disease in different individuals. Furthermore, a growing number of studies have shown that it is rare for a single bacterial species to be associated with a disease. Instead, subsets of the microbiome account for differences in health. Traditional statistical approaches do not adequately account for the variation in the human microbiome and typically consider the protective or risk effects of each bacterial population individually. Recently, machine learning models have grown in popularity among microbiome researchers because of the large amount of data that can now be generated and because the models are effective at accounting for the interpersonal microbiome variation and the ecology of the disease.

ML models are useful for understanding the variation in the structure of existing data and to apply that knowledge to make predictions about new data. Researchers have used ML models to diagnose and understand the ecological basis of diseases such as liver cirrhosis, colorectal cancer, inflammatory bowel diseases (IBD), obesity, and type 2 diabetes (1–16, 16–18). The task of diagnosing an individual with high confidence relies on a ML model that is built with rigorous methods. However, there are common methodological problems across many of these studies that need to be addressed as the field progresses. These include a lack of transparency in which methods are used and how these methods are implemented; developing and evaluating models without a separate held-out test data; large variation between the predictive performance on different folds of cross-validation; and large variation between cross-validation and testing performances. Nevertheless, the microbiome field is making progress to avoid some of these pitfalls including validating their models on independent datasets (7, 18, 19) and introducing analysis frameworks to better use ML tools (20–23). More work is needed to further improve reproducibility and minimize over-optimism for model performance.

Among microbiome researchers the lack of transparency in justifying a modelling approach has been due to an implicit assumption that more complex models are better because they are more complex. This has resulted in a trend towards using models such as random forest and neural networks (2, 11, 24–26) over simpler models such as logistic regression or other linear models (18, 22, 27). Although the more complex models may be better at incorporating non-linear relationships or yield better predictions, they are considered to be black box models because they are not inherently interpretable. These models require post hoc explanations to quantify the importance of each feature in making a prediction and they do not show the structure of how the features are used. Depending on the application of the model, researchers may choose to use different modeling approaches. For example, researchers trying to identify the populations causing a disease would likely want a more interpretable model whereas clinicians may emphasize performance. Although one may feel that they are sacrificing interpretability for performance, that tradeoff may be minimal (28, 29). Regardless, it is important for researchers to articulate why they have selected a specific modelling approach or even compare multiple approaches in the same study.

To showcase a rigorous ML pipeline and to shed light on how ML model selection can affect modeling results, we performed an empirical analysis comparing 7 modeling approaches with the same dataset and pipeline. We built three linear models with different forms of regularization: L2-regularized logistic regression and L1 and L2-regularized support vector machines (SVM) with a linear kernel. We also built four non-linear models: SVM with radial basis function kernel, a decision tree, random forest and XGBoost. We compared the predictive performance, interpretability, and computational efficiency. To demonstrate the performance of these modeling approaches and our pipeline, we used data from a previously published study that sought to classifiy individuals as having normal colons or colonic lesions based on the 16S rRNA gene sequences collected from fecal samples (3). This dataset was selected because it is a relatively large collection of individuals (N=490) connected to a clinically significant disease where there is ample evidence that the disease is driven by variation in the microbiome (1, 3, 4, 30). With this dataset we developed a framework that implementf a ML pipeline that can be used for any modeling approach, evaluates predictive performance, and demonstrates how to interpret these models. This framework can be easily applied to other host-associated and environmental microbiome datasets.

## Results

**Model selection and pipeline construction** We established a ML pipeline where we trained and validated each of the seven models using a common approach that is based on standard methods within the ML community (REFS)[Figure 1].

First, we randomly split the data into training and test sets so that the training set consisted of 80% of the full dataset while the test set was composed of the remaining 20% of the data [Figure 1]. To maintain the distribution of controls and cases that was found with the full dataset, we performed stratified splits. For example, our full dataset included 490 individuals. Of these, 261 had normal colons (53%) and 229 had a screen relevant neoplasia (SRN; 46.7%). A training set included 393 individuals, of which 209 had an SRN (53%), while the test set was composed of 97 individuals of which 52 had an SRN (54%). The training data was used to build the models and the test set was used for evaluating predictive performance.

Second, we trained seven different models using the training data. We selected models with different classification algorithms and regularization methods. Regularization is a technique that discourages overfitting by penalizing the model for learning the training data too well. For regularized logistic regression and SVM with linear kernel, we used L2 regularization to keep all potentially important features. For comparison, we also trained an L1 regularized SVM model with linear kernel. L1-regularization on microbiome data led to a sparser solution (i.e., force many coefficients to zero). To explore the potential for non-linear relationships among features to improve classification, we trained tree-based models including decision tree, random forest, and XGBoost and we trained an SVM model with non-linear kernel.

Third, fitting of these models require selecting appropriate hyperparameters. Hyperparameters are the rules that are learned from the training set in a classification algorithm. For example, in the linear models the regularization term (C) is a hyperparameter that indicates the penalty for overfitting. Similar to regularization term C, all hyperparameters are tuned to find the best model. We selected hyperparameters by performing 100 five-fold cross-validation (CV) repeats on the training set [Figure 1]. The five-fold CV was also stratified to maintain the overall case and control distribution. We chose the best hyperparameter values for each model based on its CV predictive performance using the area under the receiver operating characteristic curve (AUROC) metric [Figure S1 and S2]. The AUROC ranges from 1.0, where the model perfectly distinguishes between cases and controls, to 0.50, where the model’s predictions are no different from random chance.

Finally, we trained the full training dataset with the selected hyperparameter values and applied the model to the held-out data to evaluate the testing predictive performance of each model. The data-split, hyperparameter selection, training and testing steps were repeated 100 times to get a reliable and robust reading of model performance [Figure 1].

**Predictive performance and generalizability of the seven models.** We evaluated the predictive performance of seven models to classify individuals as having normal colons or SRNs [Figure 2]. The random forest model had significantly higher test AUROC values than the other models for detecting SRNs (Wilcoxon rank sum test, p < 0.01). The median AUROC of the random forest model was 0.695 (IQR 0.044). L2-regularized logistic regression, XGBoost, L2-regularized SVM with linear and radial basis function kernel AUROC values were not significantly different from one another and had median AUROC values of 0.68 (IQR 0.055), 0.679 (IQR 0.052), 0.678 (IQR 0.056) and 0.668 (IQR 0.056) respectively. L1-regularized SVM with linear kernel and decision tree had significantly lower AUROC values than the other ML models with median AUROC of 0.65 (IQR 0.066) and 0.601 (IQR 0.059), respectively [Figure 2]. Interestingly, these results demonstrate that the most complex model (XGBoost) did not have the best performance and that the most interpretable models (L2-regularlized logistic regression and linear SVM) performed nearly as well as random forest.

To evaluate the generalizability of each model, we compared the median cross-validation AUROC to the median testing AUROC. If the difference between the cross-validation and testing AUROCs is large, then that would indicate that the models were overfit. difference in median AUROCs was 0.021 in L1-regularized SVM with linear kernel, followed by SVM with radial basis function kernel and decision tree with a difference of 0.007 and 0.006, respectively [Figure 2]; however, these differences are relatively small and would not indicate a problem with overfitting.

To evaluate the risk for over-optimism of each model, we calculated the range for each model using 100 splits. The range among the testing AUROC values within each model varied by 0.23 on average across the seven models. If we had only done a single split, then there is a risk that we could gotten lucky or unlucky with the performance of the model. For instance, the lowest AUROC value of the random forest model was 0.59 whereas the highest was 0.81. These results showed that depending on the data-split, the testing AUROC values showed great variability [Figure 2]. Therefore, it is important to employ the hierarchical data splits that were included in our pipeline to minimize the risk of over-optimism.

**Interpretation of each ML model.** Interpretability is the degree to which humans can understand the reasons behind a model prediction (31). Because we often use ML models not just to predict a health outcome but also to learn the ecology behind a disease, model interpretation becomes crucial for microbiome studies. ML models decrease in interpretability as they increase in complexity. In this study we used two methods to interpret models with varying complexity.

We interpreted linear models (L1 and L2-regularized SVM with linear kernel and L2-regularized logistic regression) using the absolute feature weights of the trained models. We ranked the absolute values of the weights of all the OTUs for each data-split [Figure 3]. We calculated the median ranks of these features over the 100 data-splits. In the three linear models, the OTUs that had the largest median rank and drove the detection of SRNs contained sequences that were most similar to members of the Lachnospiraceae (OTU 1239, OTU 742), Roseburia (OTU00822, OTU00768), Clostridium (OTU00609), Blautia (OTU00659, OTU00012), Faecalibacterium (OTU00015), Gamella (OTU00426) and Peptostreptococcus (OTU00367) [Figure 3]. Among the five highest ranked OTUs across these three models, it was encouraging that OTU 1239 was shared among each of the models and that … were shared among both of the L2-regularized models.

We explained the feature importances in non-linear models; SVM with radial basis kernel, decision tree, random forest and XGBoost, using a method called permutation importance on the held-out test data. Permutation importance analysis is a posthoc explanation of the model where we randomly permute non-correlated features individually and groups of highly correlated features together. We then calculate how much the predictive performance of the model (i.e AUROC values) decrease when each OTU or group of OTUs is permuted randomly. We ranked the OTUs based on how much they decreased the median testing AUROC; the OTU with the largest decrease ranking highest. The top 5 OTUs with the largest negative impact on testing AUROC overlapped in tree-based models [Figure 4]. Specifically, permuting *Peptostreptococcus* (OTU00367) abundances randomly, dropped the predictive performances the most in all tree-based methods [Figure 4]. Decision tree, random forest and XGBoost models’ predictive performance dropped from 0.6 median AUROC to 0.52, from 0.69 to 0.68 and from 0.68 to 0.65, respectively [Figure 4].

To highlight the differences between the two interpretation methods, we used permutation importance to interpret linear models as well [Figure S3]. L1-regularized SVM with linear kernel picked out some of the same OTUs (OTU00822, OTU01239, OTU00609) as important in feature rankings based on weights [Figure 3] and permutation importance [Figure S3]. Similarly, L2-regularized SVM and L2-regularized logistic regression picked out some of the same OTUs in both interpretation methods, OTU00659 and OTU00012, respectively. However, for all the linear models, the rankings of these features were different due to the collinearity in microbial communities. Collinearity in a microbial dataset is when one OTU is dependant on another OTU. The feature weights of correlated OTUs are influenced by one another which makes it difficult to interpret models using feature weights. Our interpretation of L2-regularized logistic regression based on feature weight rankings, showed that an OTU that belongs to *Lachnospiraceae* (OTU00056) was not among the most important OTUs in making a prediction [Figure 3C]. However, it was picked out as important when the same model was interpreted using permutation importance [Figure S3C]. This is due to collinearity in the dataset, where *Lachnospiraceae* (OTU00056) is significantly correlated with *Feacalibacterium* (OTU00015) which has the highest ranked feature weights [Figure 3C]. We need to investigate the relationships among features to identify the true underlying factors when making a prediction.

**The computational efficiency of each ML model.**

We compared the training times of the seven ML models. As the complexity of a ML model and the number of tuned hyperparameter settings increased [Figures S1-S2], its training times increased as well [Figure 5]. Linear models trained faster than non-linear models. L1 and L2-regularized SVM with linear kernel and L2-regularized logistic regression had the shortest training times with 0.2 hours, (std ± 0.03), 0.2 hours, (std ± 0.02), and 0.2 hours, (std ± 0.02), respectively. Whereas, a decision tree, SVM with radial basis function kernel, random forest and XGBoost had training times of 4.4 hours, (std ± 0.3), 59.6 hours, (std ± 8.8), 83.2 hours, (std ± 11.3) and 155.1 hours, (std ± 1), respectively [Figure 5].

## Discussion

Microbiome studies use ML models, often with a classification task, to predict a disease but also to learn which microbes are indicators of that disease (2–11). Achieving either of these tasks have far-reaching impact on human health, however ML as a tool in microbiome studies is still at its infancy. A framework is needed to develop rigorous ML models, to identify and overcome potential pitfalls. Previous studies generated workflows to allow ML to be widely used by the microbiome researchers (20–23). This study sets-up standards for developing and evaluating rigorous ML models for microbiome data [Table 1].

We benchmarked seven ML models with different classification algorithms to show that a clearly defined ML problem that is based on the goal of the microbiome study should inform our model selection. Our results showed that if the goal of a study is to learn the ecology behind a disease and to identify microbial biomarkers, we can create ML models that are inherently interpretable and easily trained without losing predictive power. In terms of predictive performance, random forest model had the best testing AUROC values compared to the other six models. However, the second best model was L2-regularized logistic regression with a median AUROC difference of only 0.015 compared to random forest. While random forest took 83.2 hours to train, L2-regularized logistic regression trained in 12 minutes. In terms of interpretability, random forest was a more complex ML model and could only be explained using methods such as permutation importance. On the other hand, L2-regularized logistic regression was easier to interpret by ranking absolute feature weights of the trained model.

Even with interpretable models such as L2-regularized logistic regression, there are potential pitfalls when it comes to identifying biomarkers of a disease. As domain experts, we know that human-associated microbial communities have complex correlation structures that creates collinearity in the dataset. Collinearity is a severe problem and needs to be addressed to reliably interpret ML models (32). In this study we used two different methods to interpret our linear models; ranking each OTU by (1) their absolute weights in the trained models and (2) their impact on the predictive performance based on permutation importance. We observed differences in the OTU rankings between the two interpretation methods due to collinearity in the dataset. To avoid misinterpreting the models, once we identify the most important microbes, we should check for their relationships with other microbes as well. These relationships will help us generate new hypotheses about the ecology of the disease. These hypotheses needs to be tested with follow-up experiments to identify the true biomarkers of a disease.

In this study, we also established a rigorous ML pipeline to use 16S rRNA sequence counts to predict a binary health outcome. First of all, we used a held-out test set to illustrate the difference between cross-validation and testing AUROC values. When the difference between cross-validation and test performance is low, this suggest the models are not overfit and that they will perform similar with similar data. In all the models, the difference between median cross-validation and testing AUROC values did not exceed 0.021 which suggests that these models are generalizable and can be used to test similar new data. Furthermore, we performed the initial 80%-20% random datasplit 100 times in our ML pipeline. Depending on how the data is split, there is the chance of being overoptimistic about the predictive performance of a model. We showed that there was variability in AUROC values between different random data-splits in each of the models we tested. Our results showed that the testing AUROC values varied 0.23 on average between different data-splits. The randomization and resampling of the initial data-split to create a held-out test set is a crucial step in the ML pipeline to develop robust ML models and to report reliable performance metrics. Additionally, we performed a full grid search for hyperparameter settings when training our ML models. Default hyperparameter settings in previously developed ML packages in R, Python, and Matlab programming languages are inadequate for effective application of classification algorithms and need to be optimized for each new dataset used to generate a model. In the example of L1-regularized SVM with linear kernel [Figure S1], the model showed large variability between different regularization coefficients (C) and was susceptible to performing poorly if the wrong regularization coefficient was assigned to the model by default. And finally, we used the AUROC metric in our study to evaluate the predictive performance of the ML models. AUROC is always random at the value 0.5 and is a robust metric when a dataset is imbalanced.

We used a balanced CRC dataset to develop ML models with a binary classification task. We did not evaluate multicategory classification methods or regression analyses to predict non-binary outcomes. However, the principles highlighted throughout this study [Table 1] apply to all ML modeling tasks with microbiome data. The models we built were generalizable despite the high number of features microbiome datasets usually have. The generalization performance of ML models depends on sample size. The more complex the model, the more data it will need. Our dataset had 490 samples, however microbiome studies that have smaller sample sizes would benefit from using less complex models. Our analysis was limited to shallow learning methods and did not explore deep learning methods such as neural networks. Microbiome datasets often suffer from having high dimensionality but low sample sizes which makes deep learning models prone to overfitting. There are studies that address overcoming these challenges in biomedical datasets (11, 33, 34), however sudies that estalish frameworks with microbiome data are lacking. This would be an interesting direction for future work in microbiome studies.

Paragraph about deploying a model.

This study highlighted the need to make educated choices at every step of developing a ML model with microbiome data. Model selection should be done with a solid understanding of model complexity and interpretability, rigorous ML pipelines should be built with cross-validation for hyperparameter tuning and with a held-out test set for evaluating predictive performance and models should be interpreted while considering collinearity in datasets. The right methods will help us achieve the level of validity and accountability we want from models built for patient health.

## Materials and Methods

**Data collection and study population.** The data used for this analysis are stool bacterial abundances and clinical information of the patients recruited by Great Lakes-New England Early Detection Research Network study. These data were obtained from Sze et al (35). The stool samples were provided by recruited adult participants who were undergoing scheduled screening or surveillance colonoscopy. Colonoscopies were performed and fecal samples were collected from participants in four locations: Toronto (ON, Canada), Boston (MA, USA), Houston (TX, USA), and Ann Arbor (MI, USA). Patients’ colonic health was labeled by colonoscopy with adequate preparation and tissue histopathology of all resected lesions. Patients with an adenoma greater than 1 cm, more than three adenomas of any size, or an adenoma with villous histology were classified as advanced adenoma. Study had 172 patients with normal colonoscopies, 198 with adenomas and 120 with carcinomas. Of the 198 adenomas, 109 were identified as advanced adenomas. Stool provided by the patients was used for 16S rRNA gene sequencing to measure bacterial population abundances. The bacterial abundance data was generated by Sze et al, by processing 16S rRNA sequences in Mothur (v1.39.3) using the default quality filtering methods, identifying and removing chimeric sequences using VSEARCH and assigning to OTUs at 97% similarity using the OptiClust algorithm (36–38).

**Data definitions and pre-processing.**

The colorectal health of the patient was defined as two encompassing classes; Normal or Screen Relevant Neoplasias (SRNs). Normal class includes patients with non-advanced adenomas or normal colons whereas SRN class includes patients with advanced adenomas or carcinomas. The study had 261 normal and 229 SRN samples. The bacterial abundances are the features used to predict colorectal health of the patients. For each patient, we had 6920 features (fecal bacterial abundances) and a two-class label that defines their colorectal health (normal or SRN colorectal lesions as defined by colonoscopies). We established modeling pipelines for a binary prediction task Bacterial abundances are discrete data in the form of Operational Taxonomic Unit (OTU) counts. OTU counts were set to the size of our smallest sample and were subsampled at the same distances. They were then transformed by scaling to a [0-1] range.

**Model training and evaluation.**

Models were trained using the machine learning wrapper caret package (v.6.0.81) in R (v.3.5.0). Within the caret package, we have made modifications to L2-regularized SVM with linear kernel function **svmLinear3** and developed a L1-regularized SVM with linear kernel function **svmLinear4** to calculate decision values instead of predicted probabilities. These changes are available at <https://github.com/SchlossLab/Topcuoglu_ML_XXXX_2019/>.

For L2-regularized logistic regression, L1 and L2 support vector machines (SVM) with linear and radial basis function kernels we tuned the **cost** hyperparameter which determines the regularization strength where smaller values specify stronger regularization. For SVM with radial basis function kernel we also tuned **sigma** hyperparameter which determines the reach of a single training instance where for a high value of sigma, the SVM decision boundary will be dependent on the points that are closest to the decision boundary. For the decision tree model, we tuned the **depth of the tree** where deeper the tree, the more splits it has. For random forest, we tuned the **number of features** to consider when looking for the best tree split. For XGBoost, we tuned for **learning rate** and the **fraction of samples** to be used for fitting the individual base learners.For hyperparameter selection, we started with a granular grid search. Then we narrowed and fine-tuned the range of each hyperparameter. The range of the grid depends on the ML task and ML model. A full grid search needs to be performed to avoid variability in testing performance. We can use hyper-band to help us with our hyperparameter selection (39).

The computational burden during model training due to model complexity was reduced by parallelizing segments of the ML pipeline. In this study we have parallelized each data-split which allowed 100 data-splits to be processed through the ML pipeline at the same time for each model. We can further parallelize the cross-validation step for each hyperparameter setting.

**Permutation importance workflow.** We created a Spearman’s rank-order correlation matrix, corrected for multiple pairwise comparisons. We then defined correlated OTUs as having perfect correlation (correlation coef=1 and p<0.01). Non-correlated OTUs were permuted individually whereas correlated ones were grouped together and permuted at the same time.

**Statistical analysis workflow.** Data summaries, statistical analysis, and data visualizations were performed using R (v.3.5.0) with the tidyverse package (v.1.2.1). We compared the AUROC values of the seven ML models by Wilcoxon rank sum tests to determine the best predictive performance.

**Code availability.** The code for all sequence curation and analysis steps including an Rmarkdown version of this manuscript is available at <https://github.com/SchlossLab/Topcuoglu_ML_XXXX_2019/>.

**Figure 1. Machine learning pipeline showing predictive model training and evaluation flowchart.**  We split the data 80%/20% stratified to maintain the overall label distribution, performed five-fold cross-validation on the training data to select the best hyperparameter setting and then using these hyperparameters to train all of the training data. The model was evaluated on a held-out set of data (not used in selecting the model). Abbreviations: cvAUROC, cross-validation area under the receiver operating characteristic curve

**Figure 2. Generalization and classification performance of ML models using AUROC values of all cross validation and testing performances.**  The median AUROC for diagnosing individuals with SRN using bacterial abundances was higher than chance (depicted by horizontal line at 0.50) for all the ML models. Predictive performance of random forest model was higher than other ML models. The boxplot shows quartiles at the box ends and the statistical median as the horizontal line in the box. The whiskers show the farthest points that are not outliers. Outliers are data points that are not within 3/2 times the interquartile ranges. Abbreviations: SRN, screen-relevant neoplasias; AUROC, area under the receiver operating characteristic curve; SVM, support vector machine; XGBoost, extreme gradient boosting

**Figure 3. Interpretation of the linear ML models.** The absolute feature weights of (A) L2 logistic regression coefficients (B) L1 SVM with linear kernel (C) L2 SVM with linear kernel were ranked from highest rank 1 to 100 for each data-split. The feature ranks of the highest ranked five OTUs based on their median ranks are shown here. Similar OTUs had the largest impact on the predictive performance of L2 logistic regression and L2 SVM with linear kernel. Abbreviations: SVM, support vector machine; OTU, Operational Taxonomic Unit.

**Figure 4. Interpretation of the non-linear ML models.** (A) SVM with radial basis kernel (B) decision tree (C) random forest (D) XGBoost feature importances were explained using permutation importance using held-out test set. The gray rectangle and the dashed line show the IQR range and median of the base testing AUROC without any permutation performed. The colors of the box plots stand for the unique OTUs that are shared among the different models; pink for OTU0008, salmon for OTU0050, yellow for OTU00367, blue for OTU00110, green for OTU00361 and red for OTU00882. For all the tree-based models, a *Peptostreptococcus* species (OTU00367) had the largest impact on predictive performance of the model. Abbreviations: SVM, support vector machine; OTU, Operational Taxonomic Unit; RBF, radial basis kernel; OTU, Operational Taxonomic Unit.

**Figure 5. Computational efficiency of seven ML models.** The training times for of each data-split showed the differences in computational efficiency of the seven models. The median training time in hours was the highest for XGBoost and shortest for L1-regularized SVM with linear kernel. The boxplot shows quartiles at the box ends and the statistical median as the horizontal line in the box. The whiskers show the farthest points that are not outliers. Outliers are data points that are not within 3/2 times the interquartile ranges. Abbreviations: AUROC, area under the receiver operating characteristic curve; SVM, support vector machine; XGBoost, extreme gradient boosting.

**Figure S1. Hyperparameter setting performances for linear models.** (A) L2 logistic regression (B) L1 SVM with linear kernel (C) L2 SVM with linear kernel mean cross-validation AUROC values when different hyperparameters are used in training the model. The differences in AUROC values when hyperparameters change show that hyperparameter tuning is a crucial step in building a ML model.

**Figure S2. Hyperparameter setting performances for non-linear models.** (A) Decision tree (B) Random forest (C) SVM with radial basis kernel (D) XGBoost mean cross-validation AUROC values when different hyperparameters are used in training the model. The differences in AUROC values when hyperparameters change show that hyperparameter tuning is a crucial step in building a ML model.

**Figure S3. Interpretation of the linear ML models with permutation importance.** (A) L1-regularized SVM with linear kernel (B) L2-regularized SVM with linear kernel and (C) L2-regularized logistic regression were interpreted using permutation importance using held-out test set. The gray rectangle and the dashed line show the IQR range and median of the base testing AUROC without any permutation performed. Abbreviations: SVM, support vector machine; OTU, Operational Taxonomic Unit; RBF, radial basis kernel; OTU, Operational Taxonomic Unit.

## References

1. **Zeller G**, **Tap J**, **Voigt AY**, **Sunagawa S**, **Kultima JR**, **Costea PI**, **Amiot A**, **Böhm J**, **Brunetti F**, **Habermann N**, **Hercog R**, **Koch M**, **Luciani A**, **Mende DR**, **Schneider MA**, **Schrotz-King P**, **Tournigand C**, **Tran Van Nhieu J**, **Yamada T**, **Zimmermann J**, **Benes V**, **Kloor M**, **Ulrich CM**, **Knebel Doeberitz M von**, **Sobhani I**, **Bork P**. 2014. Potential of fecal microbiota for early-stage detection of colorectal cancer. Mol Syst Biol **10**. doi:[10.15252/msb.20145645](https://doi.org/10.15252/msb.20145645).

2. **Zackular JP**, **Rogers MAM**, **Ruffin MT**, **Schloss PD**. 2014. The human gut microbiome as a screening tool for colorectal cancer. Cancer Prev Res **7**:1112–1121. doi:[10.1158/1940-6207.CAPR-14-0129](https://doi.org/10.1158/1940-6207.CAPR-14-0129).

3. **Baxter NT**, **Koumpouras CC**, **Rogers MAM**, **Ruffin MT**, **Schloss PD**. 2016. DNA from fecal immunochemical test can replace stool for detection of colonic lesions using a microbiota-based model. Microbiome **4**. doi:[10.1186/s40168-016-0205-y](https://doi.org/10.1186/s40168-016-0205-y).

4. **Baxter NT**, **Ruffin MT**, **Rogers MAM**, **Schloss PD**. 2016. Microbiota-based model improves the sensitivity of fecal immunochemical test for detecting colonic lesions. Genome Medicine **8**:37. doi:[10.1186/s13073-016-0290-3](https://doi.org/10.1186/s13073-016-0290-3).

5. **Hale VL**, **Chen J**, **Johnson S**, **Harrington SC**, **Yab TC**, **Smyrk TC**, **Nelson H**, **Boardman LA**, **Druliner BR**, **Levin TR**, **Rex DK**, **Ahnen DJ**, **Lance P**, **Ahlquist DA**, **Chia N**. 2017. Shifts in the fecal microbiota associated with adenomatous polyps. Cancer Epidemiol Biomarkers Prev **26**:85–94. doi:[10.1158/1055-9965.EPI-16-0337](https://doi.org/10.1158/1055-9965.EPI-16-0337).

6. **Pasolli E**, **Truong DT**, **Malik F**, **Waldron L**, **Segata N**. 2016. Machine learning meta-analysis of large metagenomic datasets: Tools and biological insights. PLoS Comput Biol **12**. doi:[10.1371/journal.pcbi.1004977](https://doi.org/10.1371/journal.pcbi.1004977).

7. **Sze MA**, **Schloss PD**. 2016. Looking for a signal in the noise: Revisiting obesity and the microbiome. mBio **7**. doi:[10.1128/mBio.01018-16](https://doi.org/10.1128/mBio.01018-16).

8. **Walters WA**, **Xu Z**, **Knight R**. 2014. Meta-analyses of human gut microbes associated with obesity and IBD. FEBS Lett **588**:4223–4233. doi:[10.1016/j.febslet.2014.09.039](https://doi.org/10.1016/j.febslet.2014.09.039).

9. **Vázquez-Baeza Y**, **Gonzalez A**, **Xu ZZ**, **Washburne A**, **Herfarth HH**, **Sartor RB**, **Knight R**. 2018. Guiding longitudinal sampling in IBD cohorts. Gut **67**:1743–1745. doi:[10.1136/gutjnl-2017-315352](https://doi.org/10.1136/gutjnl-2017-315352).

10. **Qin N**, **Yang F**, **Li A**, **Prifti E**, **Chen Y**, **Shao L**, **Guo J**, **Le Chatelier E**, **Yao J**, **Wu L**, **Zhou J**, **Ni S**, **Liu L**, **Pons N**, **Batto JM**, **Kennedy SP**, **Leonard P**, **Yuan C**, **Ding W**, **Chen Y**, **Hu X**, **Zheng B**, **Qian G**, **Xu W**, **Ehrlich SD**, **Zheng S**, **Li L**. 2014. Alterations of the human gut microbiome in liver cirrhosis. Nature **513**:59–64. doi:[10.1038/nature13568](https://doi.org/10.1038/nature13568).

11. **Geman O**, **Chiuchisan I**, **Covasa M**, **Doloc C**, **Milici M-R**, **Milici L-D**. 2018. Deep learning tools for human microbiome big data, pp. 265–275. *In* Balas, VE, Jain, LC, Balas, MM (eds.), Soft computing applications. Springer International Publishing.

12. **Thaiss CA**, **Itav S**, **Rothschild D**, **Meijer MT**, **Levy M**, **Moresi C**, **Dohnalová L**, **Braverman S**, **Rozin S**, **Malitsky S**, **Dori-Bachash M**, **Kuperman Y**, **Biton I**, **Gertler A**, **Harmelin A**, **Shapiro H**, **Halpern Z**, **Aharoni A**, **Segal E**, **Elinav E**. 2016. Persistent microbiome alterations modulate the rate of post-dieting weight regain. Nature **540**:544–551. doi:[10.1038/nature20796](https://doi.org/10.1038/nature20796).

13. **Dadkhah E**, **Sikaroodi M**, **Korman L**, **Hardi R**, **Baybick J**, **Hanzel D**, **Kuehn G**, **Kuehn T**, **Gillevet PM**. 2019. Gut microbiome identifies risk for colorectal polyps. BMJ Open Gastroenterology **6**:e000297. doi:[10.1136/bmjgast-2019-000297](https://doi.org/10.1136/bmjgast-2019-000297).

14. **Flemer B**, **Warren RD**, **Barrett MP**, **Cisek K**, **Das A**, **Jeffery IB**, **Hurley E**, **O‘Riordain M**, **Shanahan F**, **O‘Toole PW**. 2018. The oral microbiota in colorectal cancer is distinctive and predictive. Gut **67**:1454–1463. doi:[10.1136/gutjnl-2017-314814](https://doi.org/10.1136/gutjnl-2017-314814).

15. **Montassier E**, **Al-Ghalith GA**, **Ward T**, **Corvec S**, **Gastinne T**, **Potel G**, **Moreau P**, **Cochetiere MF de la**, **Batard E**, **Knights D**. 2016. Pretreatment gut microbiome predicts chemotherapy-related bloodstream infection. Genome Medicine **8**:49. doi:[10.1186/s13073-016-0301-4](https://doi.org/10.1186/s13073-016-0301-4).

16. **Ai L**, **Tian H**, **Chen Z**, **Chen H**, **Xu J**, **Fang J-Y**. 2017. Systematic evaluation of supervised classifiers for fecal microbiota-based prediction of colorectal cancer. Oncotarget **8**:9546–9556. doi:[10.18632/oncotarget.14488](https://doi.org/10.18632/oncotarget.14488).

17. **Dai Z**, **Coker OO**, **Nakatsu G**, **Wu WKK**, **Zhao L**, **Chen Z**, **Chan FKL**, **Kristiansen K**, **Sung JJY**, **Wong SH**, **Yu J**. 2018. Multi-cohort analysis of colorectal cancer metagenome identified altered bacteria across populations and universal bacterial markers. Microbiome **6**:70. doi:[10.1186/s40168-018-0451-2](https://doi.org/10.1186/s40168-018-0451-2).

18. **Mossotto E**, **Ashton JJ**, **Coelho T**, **Beattie RM**, **MacArthur BD**, **Ennis S**. 2017. Classification of paediatric inflammatory bowel disease using machine learning. Scientific Reports **7**. doi:[10.1038/s41598-017-02606-2](https://doi.org/10.1038/s41598-017-02606-2).

19. **Wong SH**, **Kwong TNY**, **Chow T-C**, **Luk AKC**, **Dai RZW**, **Nakatsu G**, **Lam TYT**, **Zhang L**, **Wu JCY**, **Chan FKL**, **Ng SSM**, **Wong MCS**, **Ng SC**, **Wu WKK**, **Yu J**, **Sung JJY**. 2017. Quantitation of faecal fusobacterium improves faecal immunochemical test in detecting advanced colorectal neoplasia. Gut **66**:1441–1448. doi:[10.1136/gutjnl-2016-312766](https://doi.org/10.1136/gutjnl-2016-312766).

20. **Statnikov A**, **Henaff M**, **Narendra V**, **Konganti K**, **Li Z**, **Yang L**, **Pei Z**, **Blaser MJ**, **Aliferis CF**, **Alekseyenko AV**. 2013. A comprehensive evaluation of multicategory classification methods for microbiomic data. Microbiome **1**:11. doi:[10.1186/2049-2618-1-11](https://doi.org/10.1186/2049-2618-1-11).

21. **Knights D**, **Costello EK**, **Knight R**. 2011. Supervised classification of human microbiota. FEMS Microbiology Reviews **35**:343–359. doi:[10.1111/j.1574-6976.2010.00251.x](https://doi.org/10.1111/j.1574-6976.2010.00251.x).

22. **Wirbel J**, **Pyl PT**, **Kartal E**, **Zych K**, **Kashani A**, **Milanese A**, **Fleck JS**, **Voigt AY**, **Palleja A**, **Ponnudurai R**, **Sunagawa S**, **Coelho LP**, **Schrotz-King P**, **Vogtmann E**, **Habermann N**, **Niméus E**, **Thomas AM**, **Manghi P**, **Gandini S**, **Serrano D**, **Mizutani S**, **Shiroma H**, **Shiba S**, **Shibata T**, **Yachida S**, **Yamada T**, **Waldron L**, **Naccarati A**, **Segata N**, **Sinha R**, **Ulrich CM**, **Brenner H**, **Arumugam M**, **Bork P**, **Zeller G**. 2019. Meta-analysis of fecal metagenomes reveals global microbial signatures that are specific for colorectal cancer. Nature Medicine **25**:679. doi:[10.1038/s41591-019-0406-6](https://doi.org/10.1038/s41591-019-0406-6).

23. **Vangay P**, **Hillmann BM**, **Knights D**. 2019. Microbiome learning repo (ML repo): A public repository of microbiome regression and classification tasks. Gigascience **8**. doi:[10.1093/gigascience/giz042](https://doi.org/10.1093/gigascience/giz042).

24. **Galkin F**, **Aliper A**, **Putin E**, **Kuznetsov I**, **Gladyshev VN**, **Zhavoronkov A**. 2018. Human microbiome aging clocks based on deep learning and tandem of permutation feature importance and accumulated local effects. bioRxiv. doi:[10.1101/507780](https://doi.org/10.1101/507780).

25. **Reiman D**, **Metwally A**, **Dai Y**. 2017. Using convolutional neural networks to explore the microbiome, pp. 4269–4272. *In* 2017 39th annual international conference of the IEEE engineering in medicine and biology society (EMBC).

26. **Fioravanti D**, **Giarratano Y**, **Maggio V**, **Agostinelli C**, **Chierici M**, **Jurman G**, **Furlanello C**. 2017. Phylogenetic convolutional neural networks in metagenomics. arXiv:170902268 [cs, q-bio].

27. **Thomas AM**, **Manghi P**, **Asnicar F**, **Pasolli E**, **Armanini F**, **Zolfo M**, **Beghini F**, **Manara S**, **Karcher N**, **Pozzi C**, **Gandini S**, **Serrano D**, **Tarallo S**, **Francavilla A**, **Gallo G**, **Trompetto M**, **Ferrero G**, **Mizutani S**, **Shiroma H**, **Shiba S**, **Shibata T**, **Yachida S**, **Yamada T**, **Wirbel J**, **Schrotz-King P**, **Ulrich CM**, **Brenner H**, **Arumugam M**, **Bork P**, **Zeller G**, **Cordero F**, **Dias-Neto E**, **Setubal JC**, **Tett A**, **Pardini B**, **Rescigno M**, **Waldron L**, **Naccarati A**, **Segata N**. 2019. Metagenomic analysis of colorectal cancer datasets identifies cross-cohort microbial diagnostic signatures and a link with choline degradation. Nature Medicine **25**:667. doi:[10.1038/s41591-019-0405-7](https://doi.org/10.1038/s41591-019-0405-7).

28. **Rudin C**. 2018. Please stop explaining black box models for high stakes decisions. arXiv:181110154 [cs, stat].

29. **Rudin C**, **Ustun B**. 2018. Optimized scoring systems: Toward trust in machine learning for healthcare and criminal justice. Interfaces **48**:449–466. doi:[10.1287/inte.2018.0957](https://doi.org/10.1287/inte.2018.0957).

30. **Knights D**, **Parfrey LW**, **Zaneveld J**, **Lozupone C**, **Knight R**. 2011. Human-associated microbial signatures: Examining their predictive value. Cell Host Microbe **10**:292–296. doi:[10.1016/j.chom.2011.09.003](https://doi.org/10.1016/j.chom.2011.09.003).

31. **Miller T**. 2017. Explanation in artificial intelligence: Insights from the social sciences. arXiv:170607269 [cs].

32. **Dormann CF**, **Elith J**, **Bacher S**, **Buchmann C**, **Carl G**, **Carré G**, **Marquéz JRG**, **Gruber B**, **Lafourcade B**, **Leitão PJ**, **Münkemüller T**, **McClean C**, **Osborne PE**, **Reineking B**, **Schröder B**, **Skidmore AK**, **Zurell D**, **Lautenbach S**. 2013. Collinearity: A review of methods to deal with it and a simulation study evaluating their performance. Ecography **36**:27–46. doi:[10.1111/j.1600-0587.2012.07348.x](https://doi.org/10.1111/j.1600-0587.2012.07348.x).

33. **Kocheturov A**, **Pardalos PM**, **Karakitsiou A**. 2019. Massive datasets and machine learning for computational biomedicine: Trends and challenges. Ann Oper Res **276**:5–34. doi:[10.1007/s10479-018-2891-2](https://doi.org/10.1007/s10479-018-2891-2).

34. **Kim M**, **Oh I**, **Ahn J**. 2018. An improved method for prediction of cancer prognosis by network learning. Genes **9**:478. doi:[10.3390/genes9100478](https://doi.org/10.3390/genes9100478).

35. **Sze MA**, **Schloss PD**. 2018. Leveraging existing 16S rRNA gene surveys to identify reproducible biomarkers in individuals with colorectal tumors. mBio **9**:e00630–18. doi:[10.1128/mBio.00630-18](https://doi.org/10.1128/mBio.00630-18).

36. **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**, **Van Horn DJ**, **Weber CF**. 2009. Introducing mothur: Open-Source, Platform-Independent, Community-Supported Software for Describing and Comparing Microbial Communities. ApplEnvironMicrobiol **75**:7537–7541.

37. **Westcott SL**, **Schloss PD**. 2017. OptiClust, an Improved Method for Assigning Amplicon-Based Sequence Data to Operational Taxonomic Units. mSphere **2**. doi:[10.1128/mSphereDirect.00073-17](https://doi.org/10.1128/mSphereDirect.00073-17).

38. **Rognes T**, **Flouri T**, **Nichols B**, **Quince C**, **Mahé F**. 2016. VSEARCH: A versatile open source tool for metagenomics. PeerJ **4**:e2584. doi:[10.7717/peerj.2584](https://doi.org/10.7717/peerj.2584).

39. **Li L**, **Jamieson K**, **DeSalvo G**, **Rostamizadeh A**, **Talwalkar A**. 2016. Hyperband: A novel bandit-based approach to hyperparameter optimization. arXiv:160306560 [cs, stat].