A Microbial-Plus Genes Random Forest Classifier for CRC Anatomical Side

August 24, 2022

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[38]: #Created by: Tyler Kolisnik
      #April 3, 2022
      # Description:
      #This code is for evaluating and finalizing a random forest model after using_
       ⇔grid search cv to find the best parameters
      #Includes scoring and validation
      # Import required packages and functions
      import sklearn
      import numpy as np
      import pandas as pd
      import rpy2.robjects as robjects
      import pickle
      import seaborn as sns
      import matplotlib.pyplot as plt
      from sklearn.ensemble import RandomForestClassifier
      from sklearn.model_selection import train_test_split
      from sklearn import metrics
      from rpy2.robjects import pandas2ri
      from sklearn.metrics import confusion_matrix, make_scorer
      from sklearn.metrics import classification_report
      from sklearn.metrics import roc_curve
      from matplotlib import pyplot
      # Increase output plot resolution
      plt.rcParams['figure.dpi'] = 300
      plt.rcParams['savefig.dpi'] = 300
      sns.set(rc={"figure.dpi":300, 'savefig.dpi':300})
      sns.set_context('notebook')
      sns.set_style("whitegrid")
      # Import the training set data (278 samples)
      pandas2ri.activate()
```

```
[2]: # Create the classifier from the best parameter set found in the grid search CV
     rforest_finalparamset = RandomForestClassifier(
         bootstrap=True,
         class_weight=None,
         criterion='gini',
         max depth=10,
         max features=0.025,
         max leaf nodes=4,
         min_impurity_decrease=0.0,
         min_samples_leaf=1,
         min_samples_split=2,
         min_weight_fraction_leaf=0.0,
         n_estimators=50,
         n_jobs=2,
         oob_score=True,
         random_state=2,
         verbose=0,
         warm_start=False
```

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[3]: # Split the dataset into training and testing

#Test Size = 0.25 instructs the model to fit on 75% of the data and test on 25%

#Stratify ensures an even distribution of samples

#Fixing random state allows for reproducibility of results

X_train, X_test, y_train, y_test = train_test_split(tpmdata2, targetdata, test_size=0.

□ 25,random_state=3,

□ stratify=targetdata,shuffle=True)

# Fit (train) the classifier to our dataset

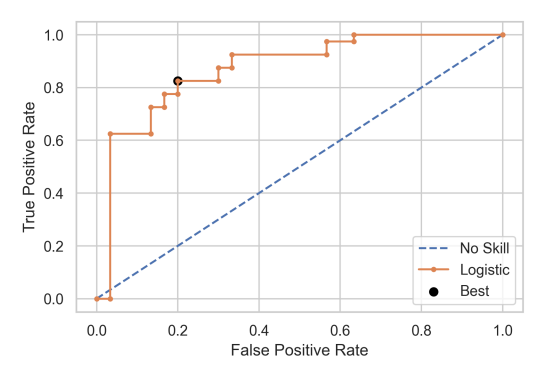
#alternatively import this from a file if you have already saved the model (. □ pkl)

rforest_finalparamset.fit(X_train,y_train)

# Load the model from disk
```

Model Accuracy: 0.8

```
[39]: # Threshold hyperparameter optimization
      #The default threshold for a binary variable is 0.5
      \#Checking the threshold allows for the optimization of the sensitivity and
       ⇔specificity
      #and a correction for imbalanced datasets
      #If a new threshold is set then you must nolonger use .predict, and must use .
       ⇔predict_proba
      # Generate prediction probabilities (necessary for AUROC/AUPRC and thresholds)
      yhat = rforest_finalparamset.predict_proba(X_test)
      # keep probabilities for the positive outcome only
      yhat = yhat[:, 1]
      # calculate roc curves
      fpr, tpr, thresholds = roc_curve(y_test, yhat)
      # calculate the q-mean for each threshold
      gmeans = np.sqrt(tpr * (1-fpr))
      # locate the index of the largest q-mean
      ix = np.argmax(gmeans)
      print('Best Threshold=%f, G-Mean=%.3f' % (thresholds[ix], gmeans[ix]))
      # plot the roc curve for the model
      pyplot.plot([0,1], [0,1], linestyle='--', label='No Skill')
      pyplot.plot(fpr, tpr, marker='.', label='Logistic')
      pyplot.scatter(fpr[ix], tpr[ix], marker='o', color='black', label='Best')
      # axis labels
      pyplot.xlabel('False Positive Rate')
      pyplot.ylabel('True Positive Rate')
      pyplot.legend()
      # show the plot
      pyplot.show()
      #With Code Adapted From: https://machinelearningmastery.com/
       → threshold-moving-for-imbalanced-classification/
```



0.554755 Threshold:

[40]:		precision	recall	f1-score	support
0		0.774194	0.800000	0.786885	30.000000
1		0.846154	0.825000	0.835443	40.000000
ac	ccuracv	0.814286	0.814286	0.814286	0.814286

```
0.810174   0.812500   0.811164   70.000000
      macro avg
                     0.815314   0.814286   0.814633   70.000000
      weighted avg
[41]: print("0.5 Threshold:")
      pd.DataFrame(classification report(y test, test predictions, output dict=True)).T
     0.5 Threshold:
[41]:
                    precision
                                 recall f1-score
                                                   support
                     0.863636 0.633333 0.730769
                                                      30.0
                     0.770833 0.925000 0.840909
      1
                                                      40.0
                                                       0.8
      accuracy
                     0.800000 0.800000 0.800000
      macro avg
                     0.817235 0.779167 0.785839
                                                      70.0
      weighted avg
                     0.810606 0.800000 0.793706
                                                      70.0
[42]: # Print Scoring Metrics for Training (Model Discovery) Set
      print("Accuracy:",metrics.accuracy_score(y_test,rfpredictions))
      print("Out-of-Bag Score:",rforest_finalparamset.oob_score_)
      print("F1 Score:",metrics.f1_score(y_test,rfpredictions))
      print("ROC AUC Score:",metrics.roc_auc_score(y_test,rfpredictions))
      print("Recall Score:",metrics.recall_score(y_test,rfpredictions))
      print("Precision Score:",metrics.precision_score(y_test,rfpredictions))
      confmat = confusion_matrix(y_test,rfpredictions)
      fp = confmat[0][1]
      tn = confmat[0][0]
      tp = confmat[1][1]
      fn = confmat[1][0]
      tprate=tp/(tp+fn)
      fprate=fp/(fp+tn)
      print("False Positives:",fp)
      print("False Negatives:",fn)
      print("True Positives:",tp)
      print("True Negatives:",tn)
      print("True Positive Rate:",tprate)
      print("True Negative Rate:",fprate)
      print(classification_report(y_test,rfpredictions))
     Accuracy: 0.8
     Out-of-Bag Score: 0.7355769230769231
     F1 Score: 0.840909090909091
     ROC AUC Score: 0.7791666666666667
     Recall Score: 0.925
     Precision Score: 0.77083333333333334
     False Positives: 11
     False Negatives: 3
     True Positives: 37
```

precision recall f1-score support 0 0.86 0.63 0.73 30 1 0.93 0.77 0.84 40 0.80 70 accuracy 0.79 70 macro avg 0.82 0.78 0.81 0.80 0.79 70 weighted avg [43]: # Generate AUROC/AUPRC scores and create functions for plotting curves val_auprc = sklearn.metrics. →average_precision_score(y_test,rfpredictions,pos_label=1) val_auroc = sklearn.metrics.roc_auc_score(y_test,rfpredictions) aurocscore="AUROC="+str(round(val auroc,2)) auprcscore="AUPRC="+str(round(val_auprc,2)) print(auprcscore) print(aurocscore) def plot_auroc_curve(y_test,rfpredictions,aurocscore): data_fpr, data_tpr, data_thresholds = sklearn.metrics. →roc_curve(y_test,rfpredictions) data_fpr_tpr = pd.DataFrame({'fpr':data_fpr, 'tpr':data_tpr}) p = sns.lineplot(data=data_fpr_tpr, x='fpr', y='tpr', ci=None) p.set(xlabel='False Positive Rate', ylabel='True Positive Rate') p.plot([0, 1], [0, 1], color='black', ls='--') p.text(0.2,0.5,aurocscore) def plot_auprc_curve(y_test,rfpredictions,auprcscore): data_prcsn, data_rcll, data_thrshlds = sklearn.metrics. →precision_recall_curve(y_test,rfpredictions) data_prcsn_rcll = pd.DataFrame({'prcsn':data_prcsn, 'rcll':data_rcll}) p = sns.lineplot(data=data_prcsn_rcll, x='rcll', y='prcsn',ci=None) p.set(xlabel='Recall', ylabel='Precision') p.plot([0, 1], [1, 0], color='black', ls='--') p.text(0.2,0.5,auprcscore) AUPRC=0.76 AUROC=0.78 [44]: # Plot AUROC Curve plot_auroc_curve(y_test,discpreds2,aurocscore)

True Negatives: 19

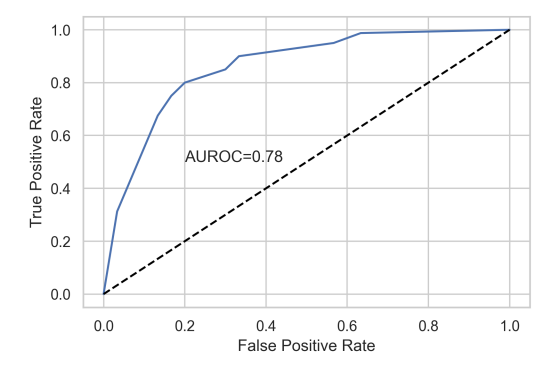
plot_auroc_curve
Save to file

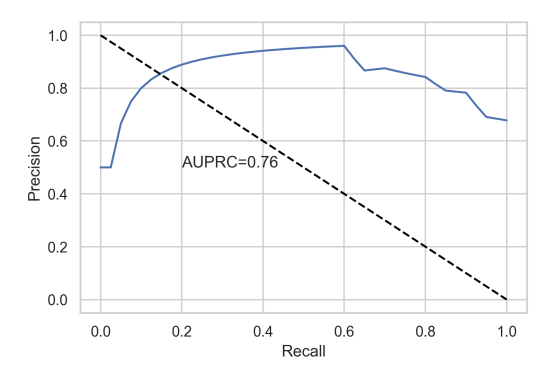
True Positive Rate: 0.925

True Negative Rate: 0.3666666666666664

#plt.savefig("/Volumes/FryShareNVME/2022-03-25-ML-Results-Side/ \Rightarrow 2022-03-27-finalized-paper-data/microbial-plusallgenes-exploration/ \Rightarrow bacteriaplusgenes-v2-AUROC-training-set.pdf")

[44]: <function __main__.plot_auroc_curve(y_test, rfpredictions, aurocscore)>





```
readRDS = robjects.r['readRDS']
      val_data=readRDS('/Volumes/FryShareNVME/2022-03-25-ML-Results-Side/
       →2022-03-27-finalized-paper-data/microbial-plusallgenes-exploration/
       $\to 2022-05-29-microbial-plus-all-genes-preprocessing workflow_SIDE/output_data/
       →2022-03-29-CPM-Microbial-PLUS-GENES-validation-set-side-30samples.rds')
      valdata=val_data[0]
      valdata2=valdata.transpose()
      valtargetdata=np.ravel(val_data[1]).astype(int)
[47]: # Run the model on the validation set
      runmodelonvaldata=rforest_finalparamset.predict(valdata2)
[48]: # Generate prediction probabilities (necessary for AUROC/AUPRC and thresholds)
      val_prob_preds = rforest_finalparamset.predict_proba(valdata2)
      # Take only the positive values
      val_preds = val_prob_preds[:, 1]
[49]: # Print Scoring Metrics for Validation Set
      print("Accuracy:",metrics.accuracy_score(runmodelonvaldata, valtargetdata))
      print("F1 Score:",metrics.f1_score(runmodelonvaldata, valtargetdata))
      print("ROC AUC Score:",metrics.roc_auc_score(runmodelonvaldata, valtargetdata))
```

[46]: # Import independent validation set (30 samples the model has never seen before)

pandas2ri.activate()

```
print("Recall Score:",metrics.recall score(runmodelonvaldata, valtargetdata))
print("Precision Score:", metrics.precision_score(runmodelonvaldata, ___
  ⇔valtargetdata))
confmat = confusion_matrix(runmodelonvaldata,valtargetdata)
fp = confmat[0][1]
tn = confmat[0][0]
tp = confmat[1][1]
fn = confmat[1][0]
tprate=tp/(tp+fn)
fprate=fp/(fp+tn)
print("False Positives:",fp)
print("False Negatives:",fn)
print("True Positives:",tp)
print("True Negatives:",tn)
print("True Positive Rate:",tprate)
print("True Negative Rate:",fprate)
print(classification_report(runmodelonvaldata, valtargetdata))
Accuracy: 0.86666666666667
F1 Score: 0.8823529411764706
ROC AUC Score: 0.8947368421052632
Recall Score: 0.7894736842105263
Precision Score: 1.0
False Positives: 0
False Negatives: 4
True Positives: 15
True Negatives: 11
True Positive Rate: 0.7894736842105263
True Negative Rate: 0.0
              precision
                         recall f1-score
                                              support
           0
                   0.73
                             1.00
                                       0.85
                                                    11
           1
                   1.00
                             0.79
                                       0.88
                                                    19
                                       0.87
                                                    30
    accuracy
  macro avg
                   0.87
                             0.89
                                       0.86
                                                    30
                             0.87
                                       0.87
weighted avg
                   0.90
                                                    30
```

```
[50]: # Generate validation set AUROC/AUPRC scores

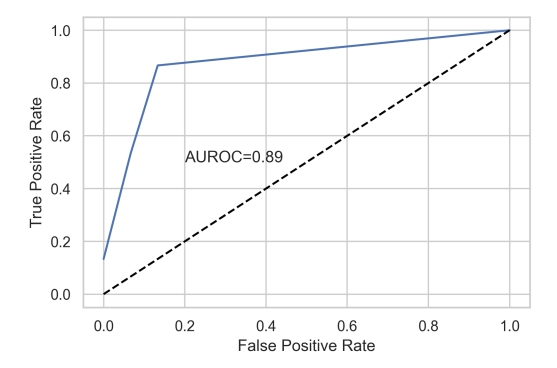
val_auprc = sklearn.metrics.average_precision_score(runmodelonvaldata,_u

valtargetdata,pos_label=1)

val_auroc = sklearn.metrics.roc_auc_score(runmodelonvaldata, valtargetdata)
aurocscore="AUROC="+str(round(val_auroc,2))
```

```
auprcscore="AUPRC="+str(round(val_auprc,2))
print(auprcscore)
print(aurocscore)
```

AUPRC=0.92 AUROC=0.89

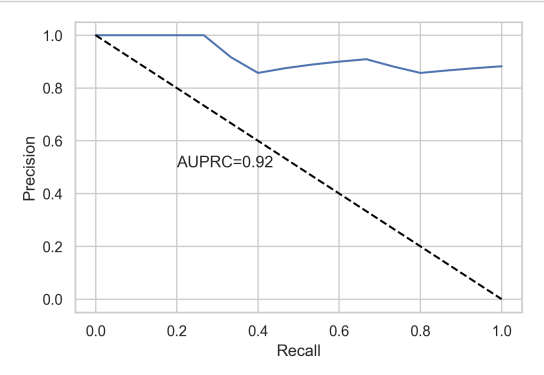


```
[52]: # Plot AUPRC Curve
plot_auprc_curve(valtargetdata,val_preds,auprcscore)
# Save to file
```

plt.savefig("/Volumes/FryShareNVME/2022-03-25-ML-Results-Side/

\$\times 2022-03-27-finalized-paper-data/microbial-plusallgenes-exploration/

\$\times bacteriaplusgenes-v2-AUPRC-validation-set.pdf")\$



0.570738 Threshold:

```
[53]:
                  precision
                               recall f1-score
                                                  support
     0
                   0.866667   0.866667   0.866667   15.000000
     1
                    0.866667 0.866667 0.866667
                                                15.000000
     accuracy
                   0.866667 0.866667 0.866667
                                                0.866667
     macro avg
                   0.866667
                             0.866667
                                      0.866667
                                                30.000000
     weighted avg
                   0.866667   0.866667   0.866667   30.000000
[54]: # Print Classification Report for 0.5 Threshold (Default)
     print("0.5 Threshold:")
     pd.
       →DataFrame(classification_report(valtargetdata, valpredictions, output_dict=True)).
     0.5 Threshold:
[54]:
                   precision
                               recall f1-score
                                                  support
                   1.000000 0.733333 0.846154 15.000000
     0
     1
                    0.789474 1.000000 0.882353 15.000000
     accuracy
                   0.866667 0.866667
                                      0.866667
                                                0.866667
     macro avg
                    weighted avg
                   [27]: # Save the feature importance scores of the random forest model mapped to the
      ⇔features (genes)
     ftnames=rforest_finalparamset.feature_names_in_
     ftimportances=rforest finalparamset.feature importances
     df=pd.DataFrame({ 'feature_names':ftnames,'feature_importances':ftimportances,})
     #df.shape
     # Save to file
     df.to csv("/Volumes/FryShareNVME/2022-03-25-ML-Results-Side/
      -2022-03-27-finalized-paper-data/microbial-plusallgenes-exploration/
       →2022-04-05-bacteriaplusgenes-v2-crc-rf-final-feature-importance-scores.csv")
[29]: # Save the model to disk
     filename = '/Volumes/FryShareNVME/2022-03-25-ML-Results-Side/
       -2022-03-27-finalized-paper-data/microbial-plusallgenes-exploration/
       -2022-04-03-microbialplusallgenes-v2-crc-side-random-forest-model.pkl
     pickle.dump(rforest_finalparamset, open(filename, 'wb'))
```