

Data Science

9 – Modeling with Ensemble Learning

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Lecture 9 - topics

- Bagging
- Boosting





Ensemble Models

https://colab.research.google.com/drive/1gNx94lEp7PCV5 ZbqTijGr1I0 odmndLE





Classification and Regression Trees Disadvantages

- *Accuracy* current methods, such as support vector machines and ensemble classifiers often have 30% lower error rates than CART.
- *Instability* if we change the data a little, the tree picture can change a lot. So the interpretation is not as straightforward as it appears.

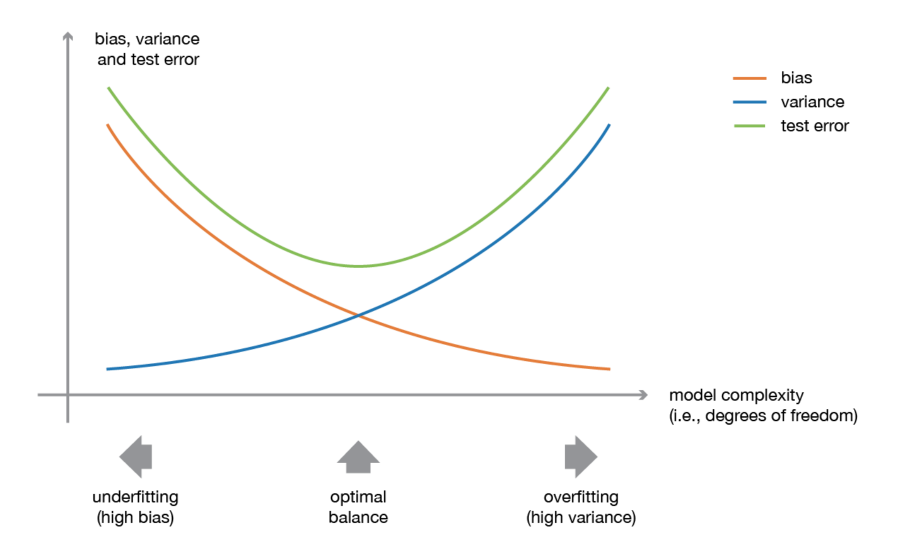
• We can do better!

Random Forests





Bias variance tradeoff







Random Forest

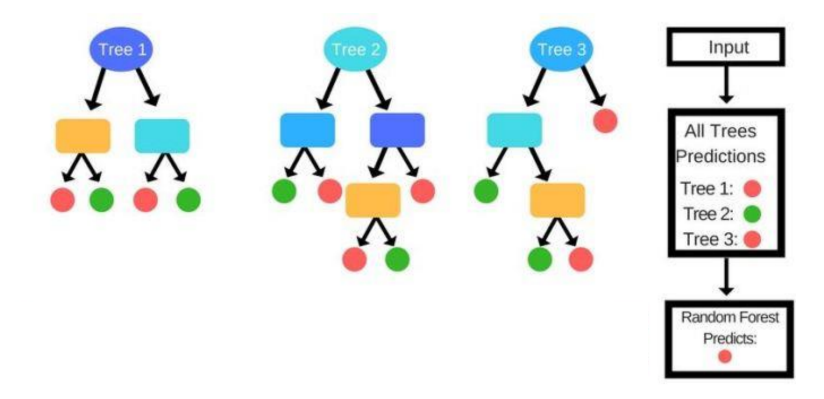
for regression and classification







Random Forest







Random forest

- An ensemble of decision tree trained by bootstrap sampling and random feature selection
- It is based on decision trees: classifiers constructed greedily using the conditional entropy
- The extension hinges on two ideas:
 - building an ensemble of trees by training on subsets of data
 - considering a reduced number of possible variables at each node





Classification and Regression Trees Pioneers

Pioneers:

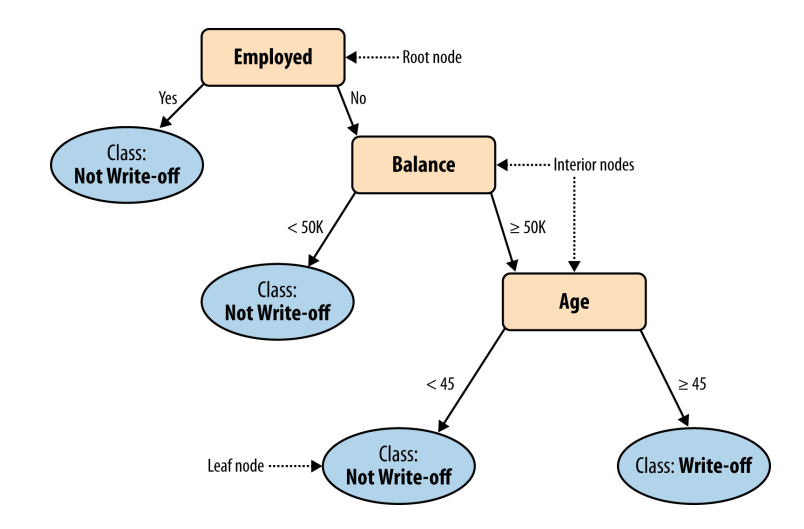
- Morgan and Sonquist (1963).
- Breiman, Friedman, Olshen, Stone (1984). CART
- Quinlan (1993). C4.5







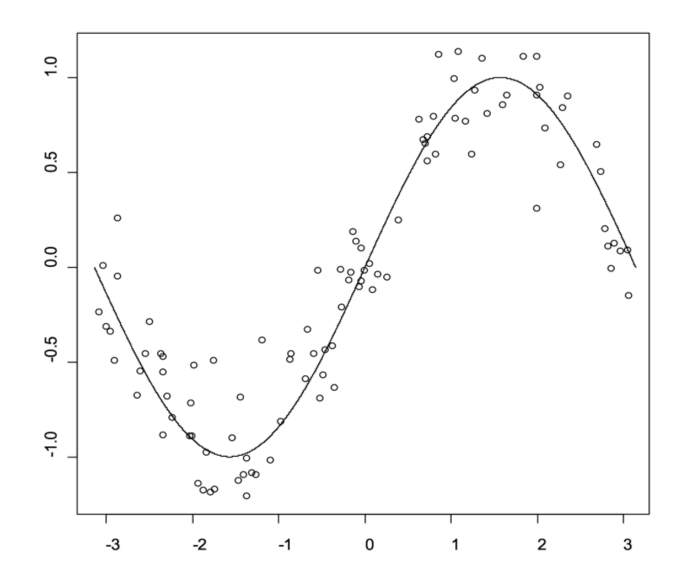
A Classification Tree







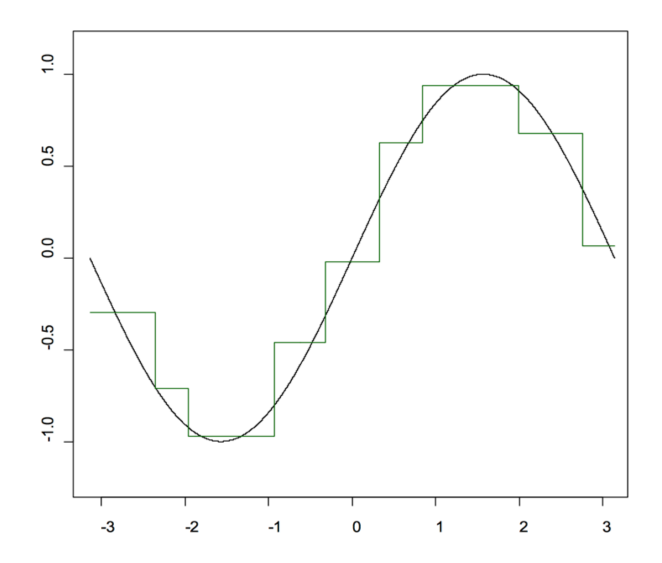
Data and Underlying Function







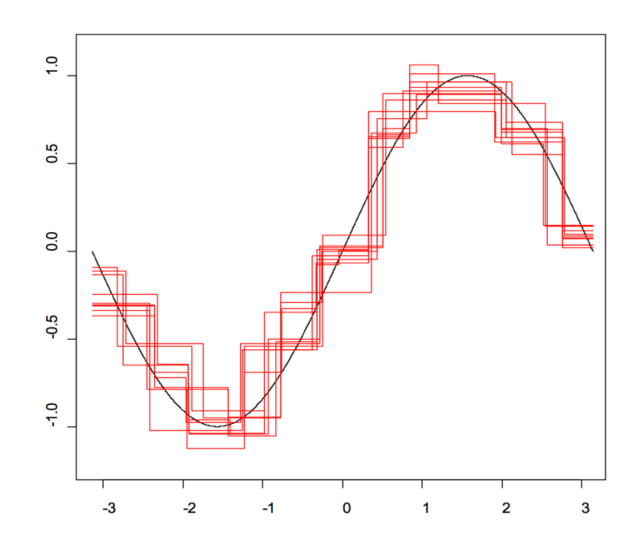
Single Regression Trees







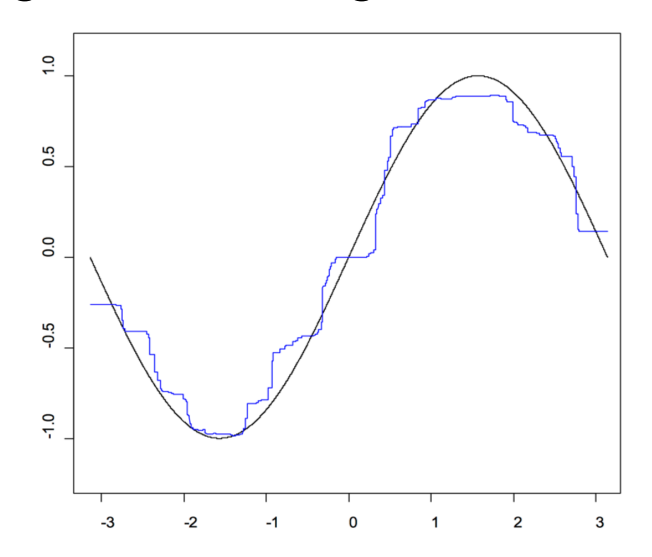
10 Regression Trees







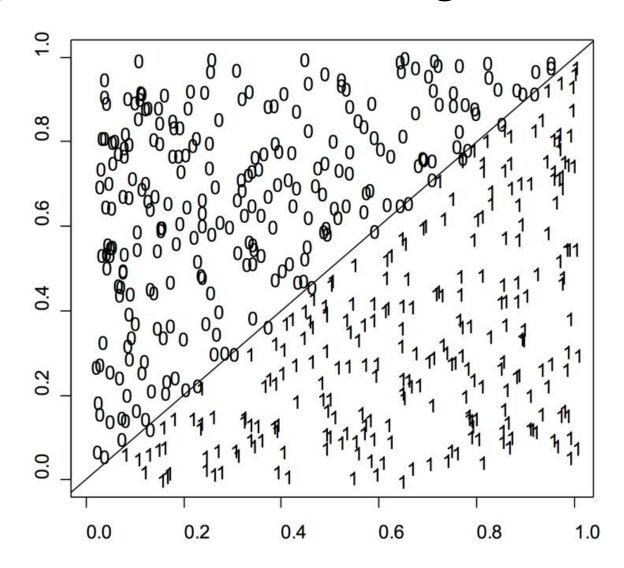
Average of 100 Regression Trees







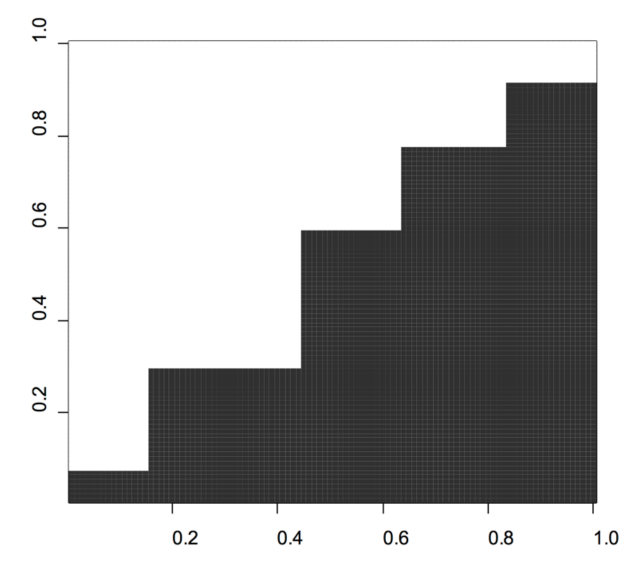
Hard problem for a single tree







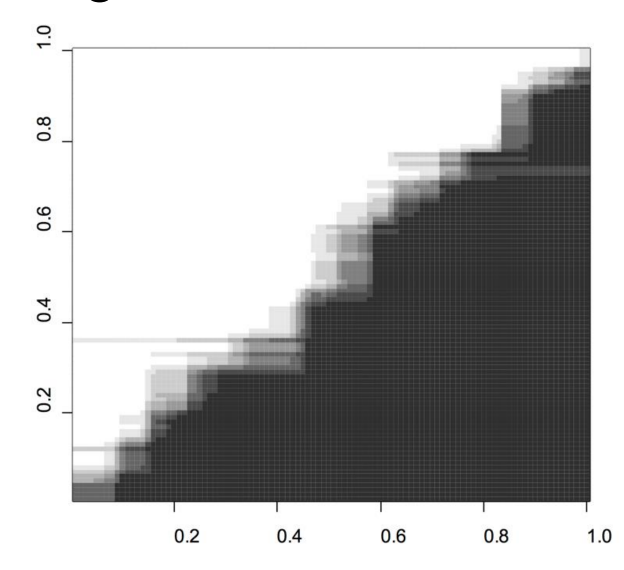
Single Tree







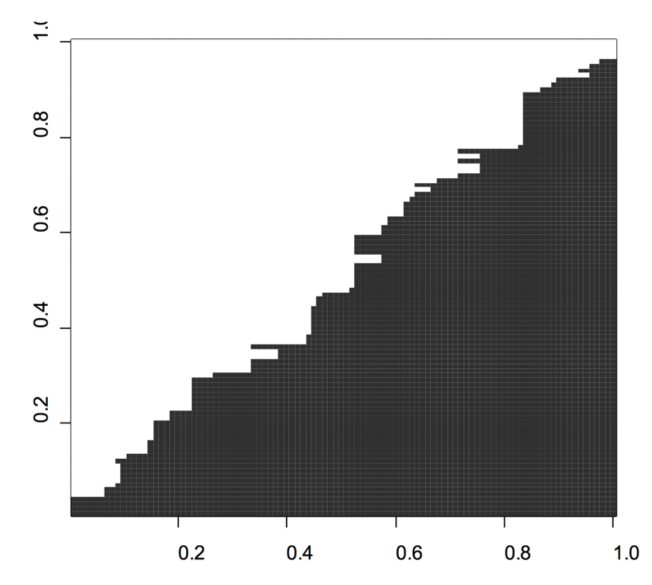
25 Averaged Trees







25 Voted Trees







Bagging (Bootstrap Aggregating)

• Fit classification or regression models to bootstrap samples from the data and combine by voting (classification) or averaging (regression)

Bootstrap sample \rightarrow $f_1(x)$

Bootstrap sample \rightarrow $f_2(x)$

Bootstrap sample \rightarrow $f_3(x)$

• • •

Bootstrap sample \rightarrow $f_M(x) \bot$

Combine $f_1(x), ..., f_M(x) \rightarrow f(x)$

 $f_i(x)$'s are "base learners"





Bagging (Bootstrap Aggregating)

- A bootstrap sample is chosen at random *with* replacement from the data. Some observations end up in the bootstrap sample more than once, while others are not included ("out of bag").
- Bagging reduces the *variance* of the base learner but has limited effect on the *bias*.
- It's most effective if we use *strong* base learners that have very little bias but high variance (unstable). E.g. trees.





Random Forests

- Grow a **forest** of many trees. (100-500)
- Grow each tree on an independent **bootstrap sample*** from the training data.
- At each node:
 - 1. Select *m* variables at random out of all *M* possible variables (independently for each node).
 - 2. Find the best split on the selected *m* variables.
- Grow the trees to maximum depth (classification). Vote/average the trees to get predictions for new data.
- *Sample N cases at random with replacement.





Random Forests

Inherit many of the advantages of CART:

- Applicable to both regression and classification problems. Yes.
- Handle categorical predictors naturally. Yes.
- Computationally simple and quick to fit, even for large problems. Yes.
- No formal distributional assumptions (non-parametric). Yes.
- Can handle highly non-linear interactions and classification boundaries.
 Yes.
- Automatic variable selection. Yes. But need variable importance too.
- Handles missing values through surrogate variables. Using proximities.
- Very easy to interpret if the tree is small. NO!





Random Forests

Improve on CART with respect to:

- *Accuracy* Random Forests is competitive with the best known machine learning methods.
- *Instability* if we change the data a little, the individual trees may change but the forest is relatively stable because it is a combination of many trees.





Two natural questions

- Why bootstrap? (Why subsample?)
 - Bootstrapping → out-of-bag data →
 - Estimated error rate and confusion matrix
 - Variable importance
- Why trees?
 - Trees → proximities →
 - Missing value fill-in
 - Outlier detection
 - Illuminating pictures of the data (clusters, structure, outliers)





Load data

```
[1] from google.colab import drive drive.mount('/content/drive')

Mounted at /content/drive

[5] import pickle

print(y_test.shape)

(1029, 44)
(441, 44)
(1029,)
(441,)
```

pickle.load(open('/content/drive/MyDrive/GSB/hr_attrition.data','rb'))

X_train, X_test, y_train, y_test = \

[6] print(X_train.shape)

print(X_test.shape)

print(y_train.shape)





Modeling - RandomForestClassifier

```
[7] from sklearn.ensemble import RandomForestClassifier

rf = RandomForestClassifier(min_samples_leaf=20, max_depth=8, class_weight='balanced')

rf.fit(X_train,y_train)
```





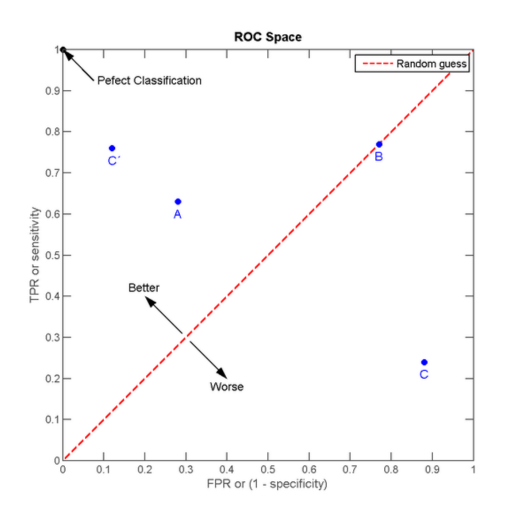
Random Forest – Feature Importance

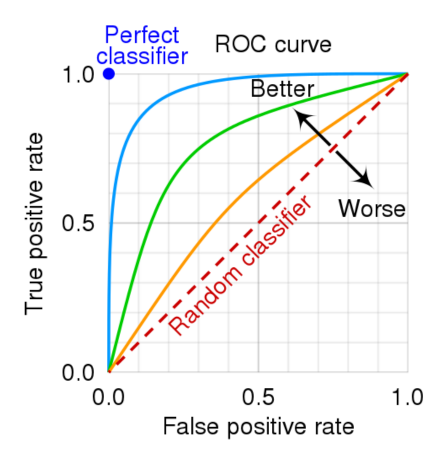
	Feature	Value
30	MonthlyIncome	0.111079
20	OverTime_Yes	0.088303
40	YearsAtCompany	0.065036
36	StockOptionLevel	0.063221
21	Age	0.057085
22	DailyRate	0.045814





Receiver Operating Characteristics









Evaluation

AUC = 0.82

```
from sklearn.metrics import RocCurveDisplay, roc_curve, auc
y_res = rf.predict_proba(X_test)
fpr, tpr, thresholds = roc_curve(y_test, y_res[:,1])
roc_auc = auc(fpr, tpr)
display = RocCurveDisplay(fpr=fpr, tpr=tpr, roc_auc=roc_auc,
                                    estimator name='Random Forest')
display.plot()
<sklearn.metrics._plot.roc_curve.RocCurveDisplay at 0x7f2470e9aee0>
    1.0
    0.8
 True Positive Rate
    0.6
    0.4
    0.2
                                               Random Forest (AUC = 0.82)
    0.0
                      0.2
                                   0.4
          0.0
                                               0.6
                                                           0.8
                                                                        1.0
                                 False Positive Rate
```





Bagging vs Boosting

• Bagging (random forests) perform well for multi-class object detection, which tends to have a lot of statistical noise.

• Gradient Boosting performs well when you have unbalanced data such as in real time risk assessment.





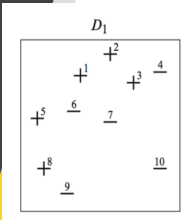
Boosting

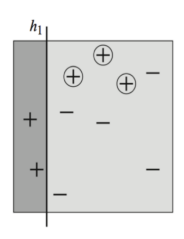


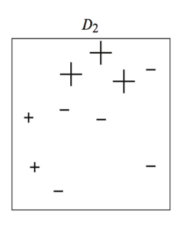


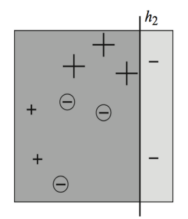
AdaBoost (Schapire and Freund, 2012)

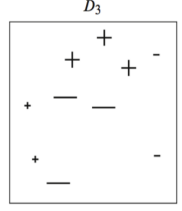
- Fit additive model $\sum_t c_t h_t(x)$
- In each stage, introduce a weak learner to compensate the shortcoming of existing weak learner

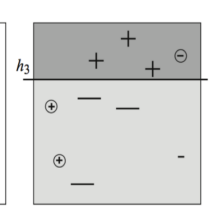










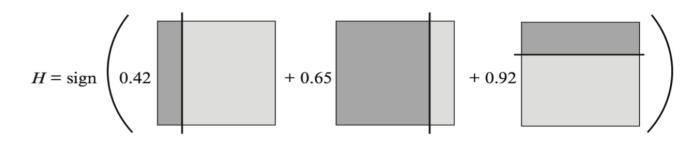






AdaBoost

$$H(x) = \sum_{t} \rho_t h_t(x)$$







Gradient Boosting

- Gradient boosting = gradient descent + boosting
- Fit additive model $\sum_t c_t h_t(x)$
- In each stage, introduce a weak learner to compensate the shortcoming of existing weak learner
- In Gradient Boosting, "shortcomings" are identified by gradients. (residual)
- Both high-weight data points and gradients tell us how to improve our model.





Gradient

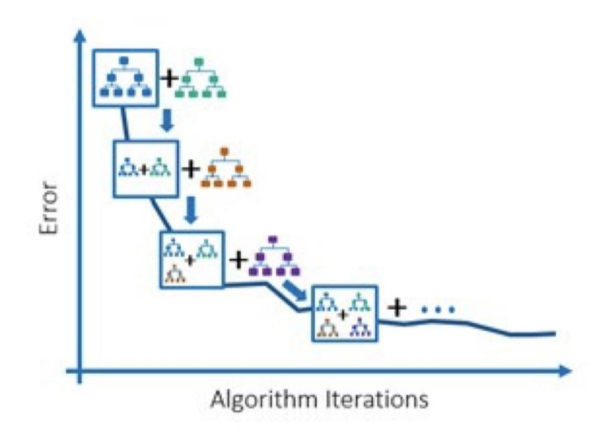
• Gradient is calculated from the derivative of the loss function.

$$Loss(y_i, \hat{y}) = (y_i - \hat{y})^2 \qquad -\nabla_{Loss}(\hat{y}) = 2 \cdot (y_i - \hat{y})$$





Gradient boosting







Gradient boosting algorithm

Algorithm 2 Gradient boosting.

```
let F_0 be a "dummy" constant model for m = 1, ..., M for each pair (x_i, y_i) in the training set compute the pseudo-residual R(y_i, F_{m-1}(x_i)) = negative gradient of the loss train a regression sub-model h_m on the pseudo-residuals add h_m to the ensemble: F_m(x) = F_{m-1}(x) + \eta \cdot h_m(x) return the ensemble F_M
```





Modeling – GradientBoostingClassifier

```
[13] from sklearn.ensemble import GradientBoostingClassifier

gbc = GradientBoostingClassifier(min_samples_leaf=20, n_estimators=200)

gbc.fit(X_train, y_train)
```

```
▼ GradientBoostingClassifier

GradientBoostingClassifier(min_samples_leaf=20, n_estimators=200)
```





Feature importance

	Feature	Value
30	MonthlyIncome	0.128732
20	OverTime_Yes	0.103441
22	DailyRate	0.058398
21	Age	0.055890
25	EnvironmentSatisfaction	0.046479
40	YearsAtCompany	0.045058
36	StockOptionLevel	0.043650
23	DistanceFromHome	0.041710
32	NumCompaniesWorked	0.039302
31	MonthlyRate	0.038362

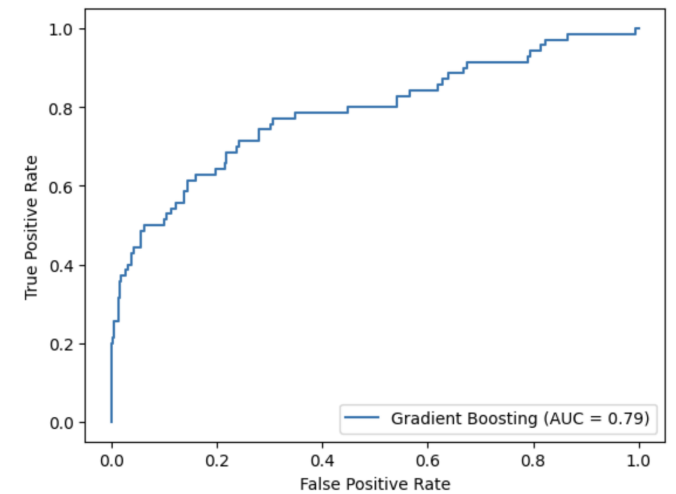






Evaluation

<sklearn.metrics._plot.roc_curve.RocCurveDisplay at 0x7f246e09a130>







eXtreme Gradient Boosting (xgboost)

xgboost is an optimized version of gradient boosting with these added features

- Regularization xgboost provides both L1 and L2 regularization in model training
- Sparsity awareness Incorporate missing data as criteria in feature selection
- Parallelization allow parallelism of algorithm in either parallel or distributed environment
- Cache aware access the learning algorithm efficiently utilizes the cache





Prepare parameters

•

```
[63] import xgboost as xgb
     param = {
         'eta': 0.3,
         'max_depth': 6,
         'objective': 'binary:logistic'
     steps = 100
```





XGBoost Parameters

XGBoost Parameters

Before running XGBoost, we must set three types of parameters: general parameters, booster parameters and task parameters.

- **General parameters** relate to which booster we are using to do boosting, commonly tree or linear model
- **Booster parameters** depend on which booster you have chosen
- **Learning task parameters** decide on the learning scenario. For example, regression tasks may use different parameters with ranking tasks.
- **Command line parameters** relate to behavior of CLI version of XGBoost.

https://xgboost.readthedocs.io/en/latest/parameter.html





Prepare data

DMatrix is a internal data structure that used by XGBoost which is optimized for both memory efficiency and training speed.

Parameters

- **data** (os.PathLike/string/numpy.array/scipy.sparse/pd.DataFrame/) dt.Frame/cudf.DataFrame Data source of DMatrix. When data is string or os.PathLike type, it represents the path libsvm format txt file, csv file (by specifying uri parameter 'path_to_csv?format=csv'), or binary file that xgboost can read from.
- label (list, numpy 1-D array or cudf.DataFrame, optional) Label of the training data.
- **missing** (*float*, *optional*) Value in the input data which needs to be present as a missing value. If None, defaults to np.nan.
- **weight** (*list, numpy 1-D array or cudf.DataFrame , optional*) Weight for each instance.

Note

For ranking task, weights are per-group.

In ranking task, one weight is assigned to each group (not each data point). This is because we only care about the relative ordering of data points within each group, so it doesn't make sense to assign weights to individual data points.

- **silent** (boolean, optional) Whether print messages during construction
- **feature_names** (*list*, *optional*) Set names for features.
- **feature_types** (*list*, *optional*) Set types for features.
- **nthread** (*integer, optional*) Number of threads to use for loading data from numpy array. If -1, uses maximum threads available on the system.





Modeling

```
[63] import xgboost as xgb
     param = {
         'eta': 0.3,
         'max_depth': 6,
         'objective': 'binary:logistic'
     steps = 100
```





Evaluation

```
fpr, tpr, thresholds = roc_curve(y_test, p)
roc_auc = auc(fpr, tpr)
display = RocCurveDisplay(fpr=fpr, tpr=tpr, roc_auc=roc_auc,
                                     estimator_name='xgboost')
display.plot()
<sklearn.metrics._plot.roc_curve.RocCurveDisplay at 0x7f245a17c250>
    1.0
    0.8
 True Positive Rate
    0.6
    0.4
    0.2
                                                       xgboost (AUC = 0.79)
    0.0 -
          0.0
                       0.2
                                   0.4
                                                0.6
                                                             0.8
                                                                          1.0
                                  False Positive Rate
```





Feature importance

```
sorted(model.get score(importance type='total gain').items(),
        key = lambda x:x[1], reverse = True)
[('MonthlyIncome', 167.65269470214844),
 ('OverTime_Yes', 119.00450897216797),
 ('DailyRate', 108.45510864257812),
 ('MonthlyRate', 103.09760284423828),
 ('Age', 94.56500244140625),
 ('DistanceFromHome', 87.48432159423828),
 ('HourlyRate', 86.21327209472656),
 ('EnvironmentSatisfaction', 65.32698059082031),
 ('NumCompaniesWorked', 58.19672775268555),
 ('StockOptionLevel', 56.460105895996094),
 ('YearsAtCompany', 49.44914245605469),
 ('YearsSinceLastPromotion', 49.413124084472656),
 ('TotalWorkingYears', 44.08100509643555),
 ('PercentSalaryHike', 43.2405891418457),
 ('RelationshipSatisfaction', 43.197994232177734),
```





Summary

- Ensemble learning allows us to generalize the machine learning model to further improve the model performance
- Bagging is robust to noise
- Boosting allows us to squeeze the performance by modeling the residuals





Lab

- Select your own classification problem data from Kaggle.com website
- Prepare the data for modeling (either classification or regression)
- Build the model for the problem
- Evaluate the modeling result





End of Lecture 5



