# 5 Random forests and model architecture

October 6, 2021

## 1 Exercise: Random forests and model architecture

In the previous exercise, we used decision trees to predict whether a crime would be solved in San Francisco.

Recall that decision trees did a reasonable job, but they have a tendency to *overfit*, meaning that the results would degrade considerably when using the *test* set or any *unseen data*.

This time we will used *random forests* to address that overfit tendency.

We will also look at how the *model's architecture* can influence its performance.

## 1.1 Data visualization and preparation

As usual, let's take another quick look at the crime dataset, then split it into train and test sets:

```
[]: import pandas
     import numpy as np
     from sklearn.model_selection import train_test_split
     import graphing # custom graphing code. See our GitHub repo for details
     # Import the data from the .csv file
     dataset = pandas.read_csv('san_fran_crime.csv', delimiter="\t")
     # Remember to one-hot encode our crime and PdDistrict variables
     categorical features = ["Category", "PdDistrict"]
     dataset = pandas.get_dummies(dataset, columns=categorical_features,_
     →drop_first=False)
     # Split the dataset in an 90/10 train/test ratio.
     # Recall that our dataset is very large so we can afford to do this
     # with only 10% entering the test set
     train, test = train_test_split(dataset, test_size=0.1, random_state=2,__
     ⇒shuffle=True)
     # Let's have a look at the data and the relationship we are going to model
     print(dataset.head())
     print("train shape:", train.shape)
     print("test shape:", test.shape)
```

```
DayOfWeek Resolution
                                      Х
                                                  Y day_of_year
                                                                  time_in_hours \
0
                     True -122.403405
                                         37.775421
                                                               29
                                                                        11.000000
            5
                     True -122.403405
                                                               29
                                                                        11.000000
1
                                         37.775421
2
            1
                     True -122.388856
                                         37.729981
                                                              116
                                                                        14.983333
            2
                                                                5
3
                    False -122.412971
                                         37.785788
                                                                        23.833333
4
            5
                    False -122.419672
                                         37.765050
                                                                1
                                                                         0.500000
   Category_ARSON
                    Category_ASSAULT
                                        Category_BAD CHECKS
                                                               Category_BRIBERY
0
                 0
                                                            0
                 0
                                     0
                                                            0
                                                                               0
1
                 0
2
                                     0
                                                            0
                                                                               0
3
                 0
                                     0
                                                            0
                                                                               0
4
                 0
                                     0
                                                            0
                                                                               0
      PdDistrict_BAYVIEW
                            PdDistrict_CENTRAL PdDistrict_INGLESIDE
0
1
                         0
                                               0
                                                                       0
                         1
                                               0
                                                                       0
2
3
                         0
                                               0
                                                                       0
                                               0
                                                                       0
4
   PdDistrict_MISSION
                        PdDistrict_NORTHERN
                                               PdDistrict PARK
0
                     0
                                            0
                                                               0
1
2
                     0
                                            0
                                                               0
3
                     0
                                            0
                                                               0
4
                                            0
                                                               0
                      1
   PdDistrict_RICHMOND
                          PdDistrict_SOUTHERN
                                                 PdDistrict_TARAVAL
0
                                             1
                       0
                                                                   0
1
                                             1
2
                       0
                                             0
                                                                   0
3
                       0
                                             0
                                                                   0
4
                       0
                                             0
                                                                   0
   PdDistrict_TENDERLOIN
0
                         0
1
                         0
2
                         0
3
                         1
                         0
4
```

[5 rows x 54 columns] train shape: (135387, 54) test shape: (15044, 54)

Hopefully this looks familiar to you! If not, jump back and go through the previous exercise on decision trees.

#### 1.2 Model assessment code

We will use the same model assessment code as we did in the previous exercise

```
[]: from sklearn.metrics import balanced_accuracy_score
     # Make a utility method that we can re-use throughout this exercise
     # To easily fit and test out model
     features = [c for c in dataset.columns if c != "Resolution"]
     def fit_and_test_model(model):
         Trains a model and tests it against both train and test sets
         global features
         # Train the model
         model.fit(train[features], train.Resolution)
         # Assess its performance
         # -- Train
         predictions = model.predict(train[features])
         train_accuracy = balanced_accuracy_score(train.Resolution, predictions)
         # -- Test
         predictions = model.predict(test[features])
         test_accuracy = balanced_accuracy_score(test.Resolution, predictions)
         return train_accuracy, test_accuracy
     print("Ready to go!")
```

Ready to go!

### 1.3 Decision tree

Let's quickly train a reasonably well-tuned decision tree to remind ourselves of its performance:

```
[]: import sklearn.tree
# re-fit our last decision tree to print out its performance
model = sklearn.tree.DecisionTreeClassifier(random_state=1, max_depth=10)

dt_train_accuracy, dt_test_accuracy = fit_and_test_model(model)

print("Decision Tree Performance:")
print("Train accuracy", dt_train_accuracy)
print("Test accuracy", dt_test_accuracy)
```

```
Decision Tree Performance:
Train accuracy 0.7742407145595661
Test accuracy 0.7597105242913844
```

#### 1.4 Random Forest

A random forest is a collection of decision trees that work together to calculate the label for a sample.

Trees in a random forest are trained independently, on different partitions of data, and thus develop different biases, but when combined they are less likely to overfit the data.

Let's build a very simple forest with two trees and the *default* parameters:

Random Forest Performance: Train accuracy 0.8842998107846062 Test accuracy 0.734378540999183

Our two-tree forest has done more poorly than the single tree on the test set, though has done a better job on the train set.

To some extent this should be expected. Random forests usually work with many more trees. Simply having two allowed it to overfit the training data much better than the original decision tree.

### 1.5 Altering the number of trees

Let's then build several forest models, each with a different number of trees, and see how they perform:

```
[]: import graphings
    # n_estimators states how many trees to put in the model
    # We will make one model for every entry in this list
    # and see how well each model performs
    n_estimators= [2, 5, 10, 20, 50]

# Train our models and report their performance
    train_accuracies = []
```

```
test_accuracies = []
     for n_estimator in n_estimators:
         print("Preparing a model with ", n_estimator, " trees...")
         # Prepare the model
         rf = RandomForestClassifier(n_estimators=n_estimator,
                                     random_state=2,
                                     verbose=False)
         # Train and test the result
         train_accuracy, test_accuracy = fit_and_test_model(rf)
         # Save the results
         test_accuracies.append(test_accuracy)
         train_accuracies.append(train_accuracy)
    Preparing a model with 2 trees...
    Preparing a model with 5
                               trees...
    Preparing a model with 10
                                trees...
    Preparing a model with 20
                                trees...
    Preparing a model with 50
                                trees...
[]: pandas.DataFrame(dict(num_of_Trees=n_estimators,Train=train_accuracies,_u
      →Test=test_accuracies))
[]:
       num_of_Trees
                         Train
                                    Test
     0
                   2 0.884300 0.734379
     1
                   5 0.971677 0.799896
     2
                  10 0.979708 0.800002
     3
                  20 0.992921 0.810757
                  50 0.999098 0.815017
[]: # Plot results
     graphings.line_2D(dict(Train=train_accuracies, Test=test_accuracies),
                         n_estimators,
                         label_x="Numer of trees (n_estimators)",
                         label_y="Accuracy",
                         title="Performance X number of trees", show=True)
```

The metrics look great for the *training* set, but not so much for the *test* set. More trees tended to help both but only up to a point.

We might have expected the number of trees to resolve our overfitting problem, but this was not the case! Chances are that the model is simply too complex relative to the data, allowing it to overfit the training set.

## 1.6 Altering the minimum number of samples for split parameter

Recall that decision trees have a root node, internal nodes and leaf nodes, and that the first two can be split into newer nodes with subsets of data.

If we let our model split and create too many nodes, it can become increasingly complex and start to overfit.

One way to limit that complexity is to tell the model that each node needs to have **at least** a certain number of samples, otherwise it can't split into subnodes.

In other words, we can set the model's min\_samples\_split parameter to the least number of samples required so that a node can be split.

Our default value for min\_samples\_split is only 2, so models will quickly become too complex if that parameter is left untouched.

We will now use the best performing model above, then try it with different min\_samples\_split values and compare the results:

We will use 20 as best performing model

```
[]: # Shrink the training set temporarily to explore this
     # setting with a more normal sample size
     full trainset = train
     train = full_trainset[:1000] # limit to 1000 samples
     min_samples_split = [2, 10, 20, 50, 100, 500]
     # Train our models and report their performance
     train_accuracies = []
     test_accuracies = []
     for min_samples in min_samples_split:
         print("Preparing a model with min_samples_split = ", min_samples)
         # Prepare the model
         rf = RandomForestClassifier(n estimators=20,
                                     min_samples_split=min_samples,
                                     random_state=2,
                                     verbose=False)
         # Train and test the result
         train_accuracy, test_accuracy = fit_and_test_model(rf)
         # Save the results
         test_accuracies.append(test_accuracy)
         train_accuracies.append(train_accuracy)
     # Plot results
```

```
Preparing a model with min_samples_split = 2
Preparing a model with min_samples_split = 10
Preparing a model with min_samples_split = 20
Preparing a model with min_samples_split = 50
Preparing a model with min_samples_split = 100
Preparing a model with min_samples_split = 500
```

As you can see above, small restrictions on the model's complexity - by limiting its ability to split nodes - reduce the gap between training and test performance. If this is subtle, it does so without damaging test performance at all.

By limiting the model complexity we address overfitting, improving its ability to generalize and make accurate predictions on *unseen* data.

Notice that using min\_samples\_split=20 gave us the best result for the *test* set, and that higher values worsened outcomes.

## 1.7 Altering the model depth

A related method to limit the trees is restricting max\_depth. This is equivalent to max\_depth we used for our decision tree, earlier. Its default value is None, which means nodes can be expanded until all leaves are *pure* (all samples in it have the same label) or have less samples than the value set for min\_samples\_split.

Whether max\_depth, or min\_samples\_split is more appropriate depends on the nature of your dataset, including its size. Usually we need to experiment to find the best settings. Let's investigate max\_depth as though we only had 500 crime samples available for our training set.

```
[]: # Shrink the training set temporarily to explore this
    # setting with a more normal sample size
    full_trainset = train
    train = full_trainset[:500] # limit to 500 samples

max_depths = [2, 4, 6, 8, 10, 15, 20, 50, 100]

# Train our models and report their performance
    train_accuracies = []
    test_accuracies = []

for max_depth in max_depths:
        print("Preparing a model with max_depth = ", max_depth)
```

```
# Prepare the model
    rf = RandomForestClassifier(n_estimators=20,
                                 max_depth=max_depth,
                                 random_state=2,
                                 verbose=False)
    # Train and test the result
    train_accuracy, test_accuracy = fit_and_test_model(rf)
    # Save the results
    test_accuracies.append(test_accuracy)
    train accuracies.append(train accuracy)
# Plot results
graphings.line_2D(dict(Train=train_accuracies, Test=test_accuracies),
                     max_depths,
                     label_x="Maximum depth (max_depths)",
                     label_y="Accuracy",
                     title="Performance", show=True)
# Rol back the trainset to the full set
train = full_trainset
Preparing a model with max_depth =
```

```
Preparing a model with max_depth = 2
Preparing a model with max_depth = 4
Preparing a model with max_depth = 6
Preparing a model with max_depth = 8
Preparing a model with max_depth = 10
Preparing a model with max_depth = 15
Preparing a model with max_depth = 20
Preparing a model with max_depth = 50
Preparing a model with max_depth = 100
```

The plot above tells us that our model actually **benefits** from a higher value for max\_depth, up to the limit of 15.

Increasing depth beyond this point begins to harm test performence, as it constrains the model too much for it to generalize.

As usual, it is important to evaluate different values when setting model parameters and defining its architecture.

## 1.8 An optimised model

Properly optimizing a model on a dataset this large can take many hours - more than you need to commit to this exercise just to learn. If you would like to run a model that has already been optimized for the full dataset, you can run the code below, and compare its performance to everything we have seen so far.

This is optional - just note that the model may take 1 - 2 minutes to train due to its size and the sheer volume of data

Training model. This may take 1 - 2 minutes

```
[]: pandas.DataFrame(data, columns = ["Model", "Train sensitivity", "Test⊔ ⇒sensitivity"])
```

```
[]: Model Train sensitivity Test sensitivity
0 Decision tree 0.774241 0.759711
1 Final random forest 1.000000 0.719738
```

As you can see, fine tuning the model's parameters resulted in a significant improvement in the *test* set results.

## 1.9 Summary

In this exercise we covered the following topics:

- Random forest models and how they differ from decision trees
- How we can change a model's architecture by setting different parameters and changing their values
- The importance of trying several combinations of parameters and evaluate these changes to improve performance

In the future you will see that different models have architectures where you can fine tune the parameters. Experimentation is needed to achieve the best possible results.