08 feature selection 2 forward

December 24, 2019

1 Kaggle Titanic survival - feature selection 2 (model forward selection)

Reducing the number of features we use can have three benefits:

- Simplifies model explanation
- Model fit may be improved by the removal of features that add no value
- Model will be faster to fit

In this notebook we will use a model-based approach whereby we incrementally add features that most increase model performance (we could use simple accuracy, but in this case we will use ROC Area Under Curve as a more thorough analysis of performance). If you are not familiar with ROC AUC, have a look at:

 $https://github.com/Michael Allen 1966/1804 _python_health care/blob/master/titanic/06_roc_sensitivity_specification and the control of the$

Two key advantages of this method are:

- It is relatively simple.
- It is tailored to the model in question.

Some key disadvantage of this method are:

- It may be slow if there are many paraemters (though the loop to select features could be limited in the number of features to select).
- The selection of features may be dependent on model meta-parameters (such as level of regularisation).
- The selection of features may not transfer between models (e.g. a model that does not allow for feature interactions may not detect features which do not add much value independently).

The machine learning model we will use to test the feature selection is as previously described:

https://github.com/MichaelAllen1966/1804_python_healthcare/blob/master/titanic/02_logistic_regression.ipython_healthcare/blo

We will also assess performance using k-fold stratification for better measurement of performance. If you are not familiar with k-fold stratification, have a look at:

https://github.com/MichaelAllen1966/1804_python_healthcare/blob/master/titanic/03_k_fold.ipynb

We will go through the following steps:

- Download and save pre-processed data
- Split data into features (X) and label (y)
- Loop through features to select the feature that most increas ROC AUC

• Plot results

https://scikit-learn.org/stable/modules/feature_selection.html#recursive-feature-elimination

```
[1]: # Hide warnings (to keep notebook tidy; do not usually do this)
import warnings
warnings.filterwarnings("ignore")
```

1.1 Load modules

A standard Anaconda install of Python (https://www.anaconda.com/distribution/) contains all the necessary modules.

```
[2]: import numpy as np
  import pandas as pd
# Import machine learning methods
  from sklearn.linear_model import LogisticRegression
  from sklearn.metrics import auc
  from sklearn.metrics import roc_curve
  from sklearn.model_selection import StratifiedKFold
  from sklearn.preprocessing import StandardScaler
```

1.2 Load data

The section below downloads pre-processed data, and saves it to a subfolder (from where this code is run). If data has already been downloaded that cell may be skipped.

Code that was used to pre-process the data ready for machine learning may be found at: https://github.com/MichaelAllen1966/1804 python healthcare/blob/master/titanic/01 preprocessing.ipynb

```
data.to_csv(data_directory + 'processed_data.csv', index=False)
```

```
[4]: data = pd.read_csv('data/processed_data.csv')
```

The first column is a passenger index number. We will remove this, as this is not part of the original Titanic passenger data.

```
[5]: # Drop Passengerid (axis=1 indicates we are removing a column rather than a row)
# We drop passenger ID as it is not original data

data.drop('PassengerId', inplace=True, axis=1)
```

1.3 Divide into X (features) and y (lables)

We will separate out our features (the data we use to make a prediction) from our label (what we are truing to predict). By convention our features are called X (usually upper case to denote multiple features), and the label (survvive or not) y.

```
[6]: X = data.drop('Survived',axis=1) # X = all 'data' except the 'survived' column y = data['Survived'] # y = 'survived' column from 'data'
```

1.4 Forward feature selection

Define data standardisation function.

```
[7]: def standardise_data(X_train, X_test):
    # Initialise a new scaling object for normalising input data
    sc = StandardScaler()

# Set up the scaler just on the training set
    sc.fit(X_train)

# Apply the scaler to the training and test sets
    train_std=sc.transform(X_train)
    test_std=sc.transform(X_test)

return train_std, test_std
```

The forward selection method:

- Keeps a list of selected features
- Keeps a list of features still available for selection
- Loops through available features:
 - Calculates added value for each feature (using stratified k-fold validation)
 - Selects feature that adds most value
 - Adds selected feature to selected features list and removes it from available features list

This methos uses a while lop to keep exploring features until no more are available. An alternative would be to use a for loop with a maximum number of features to select.

```
[8]: # Create list to store accuracies and chosen features
     roc_auc_by_feature_number = []
     chosen_features = []
     # Initialise chosen features list and run tracker
     available_features = list(X)
     run = 0
     number_of_features = len(list(X))
     # Loop through feature list to select next feature
     while len(available_features)> 0:
         # Track and pront progress
         print ('Feature run {} of {}'.format(run, number_of_features))
         # Convert DataFrames to NumPy arrays
         y_np = y.values
         # Reset best feature and accuracy
         best_result = 0
         best_feature = ''
         # Loop through available features
         for feature in available_features:
             # Create copy of already chosen features to avoid orginal being changed
             features to use = chosen features.copy()
             # Create a list of features from features already chosen + 1 new feature
             features_to_use.append(feature)
             # Get data for features, and convert to NumPy array
             X_np = X[features_to_use].values
             # Set up lists to hold results for each selected features
             test_auc_results = []
             # Set up k-fold training/test splits
             number_of_splits = 5
             skf = StratifiedKFold(n_splits = number_of_splits)
             skf.get_n_splits(X_np, y)
             # Loop through the k-fold splits
             for train_index, test_index in skf.split(X_np, y_np):
```

```
# Get X and Y train/test
            X_train, X_test = X_np[train_index], X_np[test_index]
            y_train, y_test = y[train_index], y[test_index]
            # Get X and Y train/test
            X_train_std, X_test_std = standardise_data(X_train, X_test)
            # Set up and fit model
            model = LogisticRegression(solver='lbfgs')
            model.fit(X_train_std,y_train)
            # Predict test set labels
            y_pred_test = model.predict(X_test_std)
            # Calculate accuracy of test sets
            accuracy_test = np.mean(y_pred_test == y_test)
            # Get ROC AUC
            probabilities = model.predict_proba(X_test_std)
            probabilities = probabilities[:, 1] # Probability of 'survived'
            fpr, tpr, thresholds = roc_curve(y_test, probabilities)
            roc_auc = auc(fpr, tpr)
            test_auc_results.append(roc_auc)
        # Get average result from all k-fold splits
        feature_auc = np.mean(test_auc_results)
        # Update chosen feature and result if this feature is a new best
        if feature_auc > best_result:
            best_result = feature_auc
            best_feature = feature
    # k-fold splits are complete
    # Add mean accuracy and AUC to record of accuracy by feature number
   roc_auc_by_feature_number.append(best_result)
    chosen_features.append(best_feature)
   available_features.remove(feature)
# Put results in DataFrame
results = pd.DataFrame()
results['feature to add'] = chosen features
results['ROC AUC'] = roc_auc_by_feature_number
```

Feature run 1 of 24 Feature run 2 of 24 Feature run 3 of 24 Feature run 4 of 24

```
Feature run 5 of 24
Feature run 6 of 24
Feature run 7 of 24
Feature run 8 of 24
Feature run 9 of 24
Feature run 10 of 24
Feature run 11 of 24
Feature run 12 of 24
Feature run 13 of 24
Feature run 14 of 24
Feature run 15 of 24
Feature run 16 of 24
Feature run 17 of 24
Feature run 18 of 24
Feature run 19 of 24
Feature run 20 of 24
Feature run 21 of 24
Feature run 22 of 24
Feature run 23 of 24
Feature run 24 of 24
```

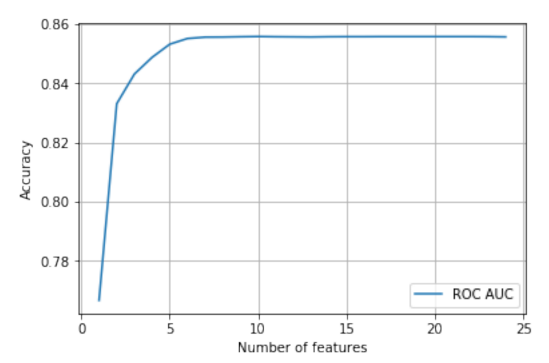
1.4.1 Show results

[9]: results

```
[9]:
            feature to add
                              ROC AUC
                      male 0.766733
     0
     1
                    Pclass 0.833036
     2
                        Age 0.843055
     3
                     SibSp 0.848686
                Embarked_S 0.853125
     4
     5
        CabinNumberImputed 0.855046
     6
             CabinLetter_C 0.855504
     7
                      SibSp
                            0.855531
     8
        CabinNumberImputed
                            0.855640
     9
                        Age
                            0.855717
     10
                 Embarked_S 0.855638
     11
                      SibSp
                            0.855598
     12
        CabinNumberImputed
                            0.855558
     13
                     Pclass
                            0.855637
     14
                      SibSp
                            0.855664
     15
                      SibSp 0.855664
                      SibSp 0.855690
     16
     17
                     SibSp 0.855690
     18
                     SibSp 0.855690
     19
                     SibSp 0.855690
```

```
20 SibSp 0.855690
21 SibSp 0.855690
22 Age 0.855663
23 Pclass 0.855584
```

1.4.2 Plot results



From the above results it looks like we could use just 5-5 features in this model. It may also be

worth examining the same method using other performance scores (such as simple accuracy, or f1) in place of ROC AUC.	