Homework 2

Due: Wed Nov. 6 @ 11:59pm

In this homework we will be performing model evaluation, model selection and feature selection in both a regression and classification setting.

The data we will be looking at are a small set of home sales data from as we might see on a real-estate website.

Instructions

Follow the comments below and fill in the blanks () to complete.

Please 'Restart and Run All' prior to submission.

Out of 65 points total.

Part 0: Environment Setup

```
In [1]: # 1. (2pts) Set up our environment with comman libraries and plotting.

# Note: generally we would do all of our imports here but some imports

# have been left till later where they are used.

# Import numpy, pandas and matplotlib. pylab

import numpy as np

import pandas as pd

import matplotlib. pylab as plt

# Execute the matplotlib magic function to display plots inline

matplotlib inline

# Setting a seed for numpy random

np. random. seed (123)
```

Part 1: Regression

In Part 1 we will try to predict a real value home sale price using several models.

```
[2]: # 2. (3pts) Load and prepare our data.
      # Read in the csv file house sales subset.csv
      df = pd. read csv('../data/house sales subset.csv')
      df. head()
      # Extract the dataframe X which should contains the first 5 columns:
            'SqFtTotLiving x1000', 'SqFtLot x1000', 'SqFtDriveway x1000', 'Bathrooms', 'Bedroo
      X = df. drop(['AdjSalePrice x100000'], axis = 1)
      #also
      #df. loc[:, 'SqFtTotLiving x1000': 'Bedrooms']. head()
      # Extract the series y r which should contain only the last column AdjSalePrice x100000
           Note: the 'r' here is denote the different target for regression vs. classificati
      y_r = df[['AdjSalePrice x100000']]
[3]: | # 3. (4pts) Create a held-aside set.
      # Import train test split from sklearn.model selection
      from sklearn. model selection import train test split
      # Split into 80% train and 20% test using train test split
      # Use random_state=42 for grading consistency
      X_train_r, X_test_r, y_train_r, y_test_r = train_test_split(X,
                                                                  test size = .2,
                                                                  random state=42)
      # Print out the length of y_test_r divided by the length y_r to confirm our test se
           Only show 2 significant digits (eg: 0.11).
      print(f' \{len(y test r)/len(y r): 0.2f\}')
```

0.20

Part 1.1 Baseline Regressor

```
[4]: | # 4. (4pt) Create a Dummy Regressior for baseline comparison
          # Import the DummyRegressor model from sklearn
         from sklearn.dummy import DummyRegressor
          # Instantiate a dummy model using strategy="median"
         dummy r = DummyRegressor(strategy="median")
          # Train the dummy model on the training set created above
         dummy_r.fit(X_train_r, y_train_r)
          # Calculate and print the training set R2 score of the trained model.
         dummy r training r2 = dummy r.score(X train r, y train r)
         print('dummy training set R2: {:.2f}'.format(dummy r training r2))
         dummy training set R2: -0.06
In [5]: | # 5. (4pts) Use 5-fold Cross Validation to get a set of negative-MSE scores
         # Import cross val score from sklearn.
         from sklearn.model selection import cross val score
          # Generate 5-fold cross valication neg mean squared error scores
              for the Dummy model on the training set.
         dummy r negmse cvscores = cross val score(dummy r, X train r, y train r,
                                                    cv = 5, scoring='neg mean squared error')
In [6]:
         # 6. (4pts) Since we'll need to convert from negative-MSE to RMSE several times
                write a function that takes in a list of negative-MSE scores
               and returns positive mean RMSE and 2 times the standard deviation
         import math
         def negmse to rmse (negmse cvscores):
              # Flip the cv scores from negative to positive
             mse cvscores = abs(negmse cvscores)
              # Transform the cv scores from MSE to RMSE
             rmse cvscores = np. sqrt (mse cvscores)
              # Calculate the mean RMSE over rmse cyscores
             rmse mean = np.mean(rmse cvscores)
              # Calculate 2 standard deviations of rmse cyscores
             rmse 2std = 2*np. std (rmse cyscores)
             return (rmse mean, rmse 2std)
```

```
In [7]: # 7. (2pts) Use our negmse_to_rmse function to calculate mean-RMSE

# and standard deviations for the dummy model.

# Pass dummy_r_negmse_cvscores to our function and capture the output dummy_r_rmse = negmse_to_rmse(dummy_r_negmse_cvscores)[0]
dummy_r_rmse_2std = negmse_to_rmse(dummy_r_negmse_cvscores)[1]

# Print out the mean RMSE and 2 standard variations for the dummy model
print('dummy mean cv RMSE: {:.2f} +- {:.2f}'.format(dummy_r_rmse,dummy_r_rmse_2std))
```

dummy mean cv RMSE: 2.28 +- 0.35

Part 1.2 Linear Regression and Residuals

```
In [8]: # 8. (4pts) Import the Linear Regression model and calculate mean RMSE using 5-fold Cross Validation

# Import the LinearRegression model from sklearn
from sklearn.linear_model import LinearRegression

# Generate 5-fold cv neg_mean_squared_error scores
# for the LinearRegression model with default settings
# on the training set.
lr_negmse_cvscores = cross_val_score(LinearRegression(), X_train_r, y_train_r, cv=5, scoring = 'neg_mean_squared_error')

# Use the function we wrote above to get mean RMSE and 2 standard deviations for Linear Regression.

lr_rmse, lr_rmse_2std = negmse_to_rmse(lr_negmse_cvscores)

# Print out the mean RMSE and 2 standard variations for LinearRegression
print('lr mean cv RMSE: {:.2f} + {:.2f}'.format(lr_rmse, lr_rmse_2std))
```

1r mean cv RMSE: 1.54 +- 0.20

```
In [9]: # 9. (6pts) Plot the residuals of a Linear Regression model

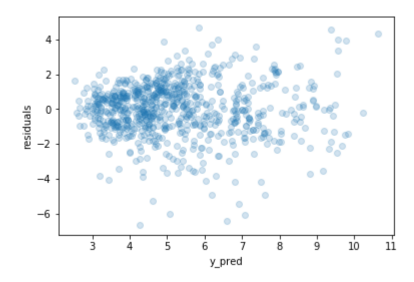
# Instantiate and retrain a linear regression model on the entire training set.
lr = LinearRegression(). fit(X_train_r, y_train_r)

# Generate predictions y_pred, again using the training set.
y_pred = lr. predict(X_train_r)

# Calculate residuals
# Recall: residual = y_pred - y
residuals = y_pred - y_train_r

# Plot predictions (x-axis) vs residuals (y-axis) using plt. scatter()
# In scatter set alpha=0.2 to make the markers somewhat transparent.
# Set axis/label names appropriately ('y_pred' and 'residuals')
# The residuals should appear fairly normal around 0 across the range of y_pred
plt. scatter(y_pred, residuals, alpha = 0.2)
plt. xlabel('y_pred')
plt. ylabel('residuals')
```

Out[9]: Text(0, 0.5, 'residuals')



Part 1.3 ElasticNet HyperParameter Tuning

```
[10]: # 10. (6pts) Use GridSearch to choose an optimal hyperparamter setting for ElasticNet
          # Import ElasticNet and GridSearchCV from sklearn
          from sklearn.linear model import ElasticNet
          from sklearn.model selection import GridSearchCV
           # Perform GridSearch over potential settings of the 11 ratio = [.1,.5,.9,1]
                 The only parameter in our search is the 11 ratio
                 Use 5-fold cross validation
          #
                 Ust
                Leave all other arguments as their defaults
                Fit on the training set
          params = {'11 ratio': [.1, .5, .9, 1]}
          gscv = GridSearchCV(ElasticNet(), params, cv=5)
          gscv. fit (X train r, y train r)
          # Print out the best parameter setting found using grid search and the best parameter s
          etting found
          print('gscv best params: {}'.format(gscv.best params))
          gscv best params: {'11_ratio': 0.1}
In [11]:
          # 11. (2pts) Calculate average RMSE for the ElasticNet model using 5-fold Cross Validat
           ion
          # Instantiate a new ElasticNet model with the optimal 11 ratio found above.
          en = ElasticNet(11 ratio = 0.1)
          # Generate 5-fold cv neg mean squared error scores
          # for the instantiated ElasticNet model on the training set.
          en negmse cyscores = cross val score(en, X train r, y train r,
                                                     cv = 5, scoring='neg mean squared error')
          # Use the function we wrote above to get mean RMSE and
          # 2 standard deviations scores.
          en rmse, en rmse 2std = negmse to rmse (en negmse cyscores)
          # Print out the mean RMSE and 2 standard variations for ElasticNet
          print ('en mean cv RMSE: {:.2f} + {:.2f}'. format (en_rmse, en_rmse_2std))
          en mean cv RMSE: 1.77 +- 0.26
```

Part 1.4 Evaluate on Test

Part 2: Classification

Here we build a model to classify low vs. high adjusted sales price.

Create Classification Target

```
In [17]: # We'll create a binary target by thresholding at the median of our AdjSalePrice
# High = 1, Low = 0
y_c = (df.AdjSalePrice_x100000 > df.AdjSalePrice_x100000.median()).astype(int)
```

Part 2.1 Create a Held-Aside Aet

Part 2.2 Measure baseline performance

baseline accuracy: 0.51

Part 2.3 Logistic Regression model

logr mean cv accuracy: 0.75 +- 0.06

```
# 16. (4pts) Perform 5-fold cross validated grid search over the number of trees and tr
ee depth.
# Import the RandomForestClassifier
from sklearn.ensemble import RandomForestClassifier
# Create the grid of parameters to evaluate
    using the settings n estimators: [5, 100, 200], max depth: [3, 5, 10].
params = {'n estimators': [5, 100, 200],
         'max depth': [3, 5, 10]}
# Instantiate and fit GridSearchCV on the classification training set
   using 5-folds, the RandomForestClassifier and default scoring.
   Make sure refit=True (default) so the model is retrained on the entire training se
gscv = GridSearchCV(RandomForestClassifier(), params, cv=5)
gscv.fit(X train c, y train c)
# Print out the best mean accuracy found and the best parameter setting found
print('rf best accuracy: {:.3f}'.format(gscv.best score ))
print('rf best params : {}'.format(gscv.best params ))
rf best accuracy: 0.797
rf best params : {'max depth': 10, 'n estimators': 200}
```

Part 2.4 Evaluate on Test

```
In [28]: # 17. (3pts) Evaluate the Random Forest Model on the test set

# Get the trained RandomForest model from gscv
# Note: there is no need to retrain here. See the documentation for clarification.

rf = gscv.best_estimator_

# Calculate accuracy on the test set using the trained model
test_acc = rf.score(X_test_c, y_test_c)

print('test acc : {:.2f}'.format(test_acc))

test acc : 0.79
```

Part 2.4 Plotting Precision-Recall curve for the Random Forest model

```
In [41]: # 18. (5pts) Get the retrained model from gscv and use it to
# import precision_recall_curve for the RandomForest model

# import precision_recall_curve from sklearn
from sklearn.metrics import precision_recall_curve

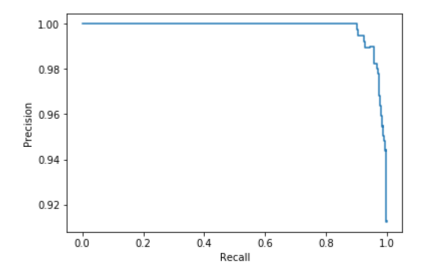
# Calculate P(y=1/x) for the training set using the trained RandomForest model
pypos_rf = rf.predict_proba(X_train_c)

# Calculate precision and recall using the y_train_c and pypos_rf
precision, recall, thresholds = precision_recall_curve(y_train_c, pypos_rf[:,1])

# Plot the curve using plt. step()
# Recall should be on the x-axis
# Label the x and y axes appropriately

plt. step(recall, precision)
plt.xlabel('Recall')
plt.ylabel('Precision')
```

Out[41]: Text(0, 0.5, 'Precision')



Part 2.6 Feature selection

kept columns: ['SqFtTotLiving_x1000' 'SqFtLot_x1000']