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Introduction to ML via Sklearn

*Present x: The title slide and the Learning Objectives slide. An overview of what we will achieve in this course*

Lesson Objectives:

By the end of this lesson, you will be able to:

* Prepare data for different types of models
* Tune model hyperparameters
* Scale data and tune hyperparameters in a pipeline
* Extract feature importance from tuned model
* Evaluate performance of classification and regression models

Introduction

*Present x: Brief introduction to ML via Scikit-Learn*

Scikit-learn is a free, open-source library built for Python that contains an assortment of supervised and unsupervised machine learning algorithms. Additionally, scikit-learn provides functions for data preprocessing, hyperparameter tuning, and model evaluation. Scikit-learn streamlines the model building process and is easy to install on a wide variety of platforms. Started in 2007 as a Google Summer of Code project by David Corneapeau, and after a series of developments and releases, scikit-learn has evolved into one of the premier tools used by academics and professionals for general purpose machine learning. In this lesson,

First steps with Scikit learn

To get started with scikit-learn, we will begin by installing the library on our local machine. We will then cover how to preprocess our data’s features by creating dummy variables, scaling the features, and splitting them it into training and testing data sets.

Installation

Scikit-learn is natively installed in the Anaconda distribution of Python as well as some Docker images, but it can also be installed with pip. However, as a prerequisite for Scikit-learn’s installation, NumPy and SciPy must be installed. Once NumPy and SciPy have been installed, scikit-learn can be installed from the terminal using pip install scikit-learn.

Check to make sure scikit-learn is installed using:

import sklearn

print(sklearn.\_\_version\_\_)



Figure 2.x: Version of scikit-learn installed

Now that we have successfully installed scikit-learn on our machine, we can begin building machine learning models. In this chapter, we will be learning linear and logistic regression, support vector machines (SVM), decision trees, and random forests. First, we will cover linear and logistic regression.

Introduction to linear and logistic regression

*Present x: Introduction to linear and logistic regression*

In regression, a single dependent, or outcome, variable is predicted using 1 or more independent variables. Use-cases for regression include but are not limited to predicting:

* win percentage given a variety of team statistics
* the risk of heart disease given family history and a number of physical and psychological characteristics
* the likelihood of snowfall given a number of climate measurements

Linear and logistic regression are popular choices for predicting such aforementioned outcomes due to the ease and transparency of interpretability as well as the ability to extrapolate to values not seen in the training data.

The end goal of linear regression is to draw a straight line through the observations that minimizes the absolute distance between the line and the observations (i.e., the line of best fit). Therefore, in linear regression, it is assumed that the relationship between the feature(s) and the continuous dependent variable follows a straight, linear, line. Lines are defined in slope-intercept form (i.e., *y = a + bx*) whereas *a* is the intercept (i.e., the value of *y* when *x* is 0), *b* is the slope, and *x* is the independent variable. There are two types of linear regression: simple linear regression and multiple linear regression.

Simple linear regression

*Present x: Simple linear regression*

Simple linear regression models define the relationship between 1 feature and the continuous outcome variable using *y* = *α* + *βx*. This equation is similar to the slope-intercept form whereas *y* denotes the predicted value of the dependent variable, *α* denotes the intercept, *β* (beta) represents the slope, and *x* is the value of the independent variable. Given *x*, regression models compute the values for *α* and *β* that minimize the absolute difference between predicted y values (i.e., y-hat) and actual y values. For example, if we are predicting the weight of an individual in kg using height in meters as the lone predictor variable, and the simple linear regression model computes 1.5 as the value for *α* and 50 as the coefficient for *β*, this model can be interpreted as for every 1 m increase in height, weight increases by 50 kg. Thus, we can predict that the weight of an individual who is 1.8 m is 91.5 kg using *y* = 1.5 + (50 x 1.8). In the following exercises we will demonstrate conducting simple linear regression using scikit-learn.

Exercise 1: Preparing data for linear regression model

*Exercise 1: Preparing data for linear regression model*

To prepare our data for a simple linear regression model, we will use a random subset of the Weather in Szeged 2006-2016 dataset which consists of hourly weather measurements from April 1, 2006 to September 9, 2016 in Szeged, Hungary. The adapted data consists of 10000 observations of 8 variables:

* Temperature\_c: temperature in Celsius
* Humidity:
* Wind\_Speed\_kmh: wind speed in kilometers per hour
* Wind\_Bearing\_Degrees:
* Visibility\_km: visibility in kilometers
* Pressure\_millibars:
* Rain: rain = 1, snow = 0
* Description: warm, normal, or cold

1. Import the ‘weather.csv’ data set using the following code:

import pandas as pd

df = pd.read\_csv('weather.csv')

1. Explore the data using df.info()

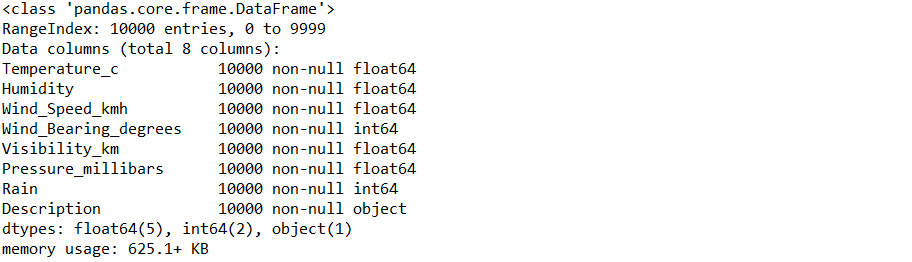


Figure 2.x: Information describing df

1. The ‘Description’ column is the lone categorical variable in df. Check the number of levels in ‘Description’ as follows:

levels = len(pd.value\_counts(df['Description']))

print('There are {} levels in the Description column'.format(levels))



Figure 2.x: Number of levels in the ‘Description’ column

1. The code below shows how to dummy code all categorical variables in 1 step:

import pandas as pd

df\_dummies = pd.get\_dummies(df, drop\_first=True)

1. The original data frame, df, consisted of 8 columns; one of which was nominal with 3 levels. In step 4, we transformed the nominal feature into 2 separate dummy variables and dropped the original feature, ‘Description’. Thus, df\_dummies should now contain 9 columns. Check this out using the following code:

print('There are {} columns in df\_dummies'.format(df\_dummies.shape[1]))



Figure 2.x: Number of columns after dummy coding

1. It is good practice to remove any possible order effects by shuffling the rows in the data before splitting the data into features (X) and outcome (y). To shuffle the rows in df\_dummies, refer to the code below:

from sklearn.utils import shuffle

df\_shuffled = shuffle(df\_dummies, random\_state=42)

1. Now that the data has been shuffled, we will split the rows in our data into features (X) and dependent variable (y). In this exercise, we will pretend that the column ‘Temperature\_c’ (temperature in Celsius) is the dependent variable and that we are preparing data to fit a linear regression model. First, we will split df\_shuffled into X and y as follows:

DV = ‘Temperature\_c’

X = df\_shuffled.drop(DV, axis=1) # get features (X)

y = df\_shuffled[DV]

1. Lastly, we will split X and y into testing and training data using the code below:

from sklearn.model\_selection import train\_test\_split

X\_train, X\_test, y\_train, y\_test = train\_test\_split(X, y, test\_size=0.33, random\_state=42)

Now that the data has been dummy coded, shuffled, split into X and y, and further divided into testing and training data sets it is ready to be used in a linear or logistic regression model.

Note:

It is acknowledged that we did not be scale our features when we prepared our data for linear and logistic regression analysis. However, later in the lesson when we learn about SVM, decision trees, and random forests we will be scaling our features in addition to the preprocessing steps described below.

**Discussion:** Why might we refrain from scaling our data before using it in a linear or logistic regression analysis?

**Answer**: A major reason for using linear or logistic regression is the ease of interpretability. Scaling the data creates an intercept and slope for a line of best fit that applies to the transformed features and not the original features, making interpretability much more convoluted.

Exercise 2: Fitting a simple linear regression model and determining the intercept and coefficient

*Exercise 2: Fitting a simple linear regression model*

In this exercise, we will continue using the data we prepared in Exercise 1 to fit a simple linear regression model to predict the temperature in Celsius from the humidity.

1. To instantiate a linear regression model refer to the code below:

from sklearn.linear\_model import LinearRegression

model = LinearRegression()

1. Fit the model to the ‘Humidity’ column in the training data using model.fit(X\_train[['Humidity']], y\_train).



Figure 2.x: Output from fitting the simple linear regression model

1. Extract the value for the intercept using the following code:

intercept = model.intercept\_

1. Extract the value for the coefficient as follows:

coefficient = model.coef\_

1. Now, we can print a message with the formula for predicting temperature in Celsius using the code below:

print('Temperature = {0:0.2f} + ({1:0.2f} x Humidity)'.format(intercept, coefficient[0]))



Figure 2.x: Formula to predict temperature in Celsius from humidity using simple linear regression

Great work! According to this simple linear regression model, a day with 0.78 humidity has a predicted temperature in Celsius of 10.56. Now that we are familiar with extracting the intercept and coefficients of our simple linear regression model, it is time to generate predictions and subsequently evaluate how the model performs on unseen, test data.

Exercise 3: Generating predictions and evaluating performance of simple linear regression model

*Exercise 3: Generating predictions and evaluating performance of simple linear regression model*

The very purpose of supervised learning is to use existing, labeled data to generate predictions. Thus, this exercise will demonstrate how to generate predictions on the test feature and generate model performance metrics by comparing the predictions to the actual values.

1. Generate predictions on the test data using the following:

predictions = model.predict(X\_test[['Humidity']])

1. A common way to evaluate model performance is to examine the correlation between the predicted and actual values using a scatterplot and a Pearson r correlation coefficient. Refer to the following code:

import matplotlib.pyplot as plt

from scipy.stats import pearsonr

plt.scatter(y\_test, predictions)

plt.xlabel('Y Test (True Values)')

plt.ylabel('Predicted Values')

plt.title('Predicted vs. Actual Values (r = {0:0.2f})'.format(pearsonr(y\_test, predictions)[0], 2))

plt.show()

Below is the resultant output:

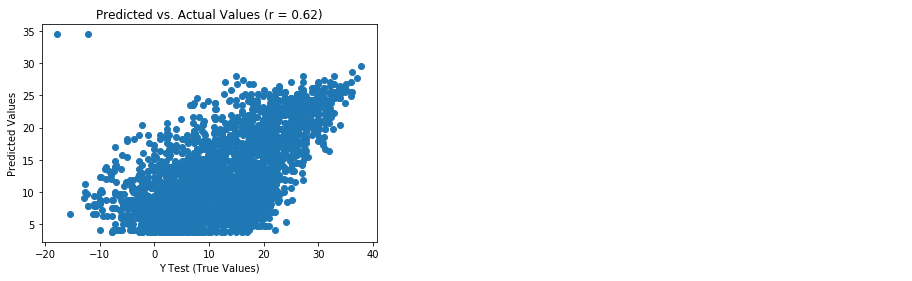


Figure 2.x: Predicted vs actual values from simple linear regression model

1. With a Pearson r value of 0.62, there is a moderate correlation between the predicted and actual values. A perfect model would have all points on the plot in a straight line and an r value of 1.0. Another indicator of model performance is the distribution of the residuals (i.e., the difference between the predicted and actual values). If the model fits the data well, the residuals will be normally distributed. To create a density plot of the residuals, refer to the following code:

import seaborn as sns

from scipy.stats import shapiro

sns.distplot((y\_test - predictions), bins = 50)

plt.xlabel('Residuals')

plt.ylabel('Density')

plt.title('Histogram of Residuals (Shapiro W p-value = {0:0.3f})'.format(shapiro(y\_test - predictions)[1]))

plt.show()

Refer to the resultant output below:

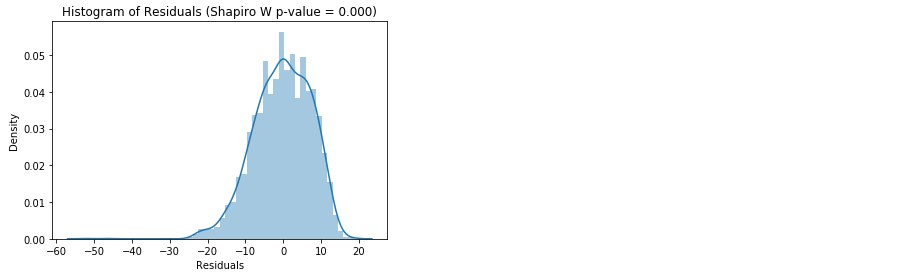


Figure 2.x: Histogram of residuals from simple linear regression model

1. The histogram shows us that the residuals are negatively skewed and the value of the Shapiro W in the title tells us that the distribution is not normal. This gives us further evidence that our model is not so good. Lastly, we will compute metrics for mean absolute error, mean squared error, root mean squared error, and R-squared and put them into a data frame using the code below:

from sklearn import metrics

import numpy as np

metrics\_df = pd.DataFrame({'Metric': ['MAE',

'MSE',

'RMSE',

'R-Squared'],

'Value': [metrics.mean\_absolute\_error(y\_test, predictions),

metrics.mean\_squared\_error(y\_test, predictions),

np.sqrt(metrics.mean\_squared\_error(y\_test, predictions)),

metrics.explained\_variance\_score(y\_test, predictions)]}).round(3)

print(metrics\_df)

Please refer to the resultant output:



Figure 2.x: Model evaluation metrics from simple linear regression model

There we have it. We have successfully used scikit-learn to fit and evaluate a simple linear regression model. This is the first step in a very exciting journey to becoming a machine learning guru. Next, we will continue expanding our knowledge of regression by exploring multiple linear regression.

Multiple linear regression

*Present x: Multiple linear regression*

Multiple linear regression models define the relationship between 2 or more features and the continuous outcome variable using *y* = *α* + *β1xi1 + β2xi2* + … + *βp-1xi,p-1.* Again, *α* represents the intercept and *β* denotes the slope for each feature (*x*) in the model. Thus, if we are predicting the weight of an individual in kg using height in m, total cholesterol in mg/dL, and minutes of cardiovascular exercise per day, and the multiple linear regression model computes 1.5 as the value for *α*, 50 as the coefficient for *β1*, 0.1 as the coefficient for *β2,* and -0.4 as the coefficient for *β3*, this model can be interpreted as for every 1 m increase in height, weight increases by 50 kg, controlling for all other features in the model. Additionally, for every 1 mg/dL increase in total cholesterol, weight increases by 0.1 kg, controlling for all other features in the model. Lastly, for every minute of cardiovascular exercise per day, weight decreases by 0.4 kg, controlling for all other features in the model. Thus, we can predict the weight of an individual who is 1.8 m tall, with total cholesterol of 200 mg/dL, and completes 30 minutes of cardiovascular exercise per day as 99.5 kg using y = 1.5 + (0.1 x 50) + (200 x 0.5) + (30 x -0.4). In the following exercise we will demonstrate conducting multiple linear regression using scikit-learn.

Exercise 4: Fitting a multiple linear regression model and determining the intercept and coefficient

*Exercise 4: Fitting a multiple linear regression model and determining the intercept and coefficient*

In this exercise, we will continue using the data we prepared in Exercise 1 to fit a multiple linear regression model to predict the temperature in Celsius from all of the features in the data.

1. To instantiate a linear regression model refer to the code below:

from sklearn.linear\_model import LinearRegression

model = LinearRegression()

1. Fit the model to the training data using model.fit(X\_train, y\_train).



Figure 2.x: Output from fitting the multiple linear regression model

1. Extract the value for the intercept using the following code:

intercept = model.intercept\_

1. Extract the value for the coefficients as follows:

coefficients = model.coef\_

1. Now, we can print a message with the formula for predicting temperature in Celsius using the code below:

print('Temperature = {0:0.2f} + ({1:0.2f} x Humidity) + ({2:0.2f} x Wind Speed) + ({3:0.2f} x Wind Bearing Degrees) + ({4:0.2f} x Visibility) + ({5:0.2f} x Pressure) + ({6:0.2f} x Rain) + ({7:0.2f} x Normal Weather) + ({8:0.2f} x Warm Weather)'.format(intercept, coefficients[0], coefficients[1], coefficients[2], coefficients[3], coefficients[4], coefficients[5], coefficients[6], coefficients[7]))



Figure 2.x: Formula to predict temperature in Celsius from humidity using multiple linear regression

Nice job! According to this multiple regression model, a day with 0.78 humidity, 5.0 wind speed, 81 wind bearing degrees, 3 km of visibility, 1000 millibars of pressure, no rain, and is described as normal has a predicted temperature in Celsius of 5.72 degrees. Now that we are familiar with extracting the intercept and coefficients of our multiple linear regression model, we can generate predictions and evaluate how the model performs on the test data.

**Discussion**: Why are some of the beta coefficients 0? By what value do I multiply the beta coefficient for ‘Rain’ if it is raining? And why is there no beta coefficient for a cold day?

**Answer**: Some of the beta coefficients are displayed as 0 because they were rounded to 2 decimal places. If it is raining, we would multiply the beta coefficient for ‘Rain’ by 1 (since it was not raining in our example, we multiplied it by 0). There is no beta coefficient for ‘Cold’ weather because there were 2 dummy variables created in place of the ‘Description’ column; one for ‘Normal’ and one for ‘Warm’.

Activity 1: Generating predictions and evaluating performance of multiple linear regression model

*Activity 1: Generating predictions and evaluating performance of multiple linear regression model*

In Exercise 3, we learned how to generate predictions and evaluate the performance of a simple linear regression model using a variety of methods. To reduce the code redundancy, we will evaluate the performance of our multiple linear regression model using the metrics in step 4 of Exercise 3 and we will determine if the multiple linear regression model performed better or worse in relation to the simple linear regression model.

1. Generate predictions on the test data using all of the features
2. Calculate the metrics for mean absolute error, mean squared error, root mean squared error, and R-squared and put them into a data frame
3. Determine if the multiple linear regression model performed better or worse in relation to the simple linear regression model

**Solution:**

1. Generate predictions on the test data using the following:

predictions = model.predict(X\_test)

1. Calculate the metrics for mean absolute error, mean squared error, root mean squared error, and R-squared and put them into a data frame as follows:

from sklearn import metrics

import numpy as np

metrics\_df = pd.DataFrame({'Metric': ['MAE',

'MSE',

'RMSE',

'R-Squared'],

'Value': [metrics.mean\_absolute\_error(y\_test, predictions),

metrics.mean\_squared\_error(y\_test, predictions),

np.sqrt(metrics.mean\_squared\_error(y\_test, predictions)),

metrics.explained\_variance\_score(y\_test, predictions)]}).round(3)

print(metrics\_df)

Please refer to the resultant output:



Figure 2.x: Model evaluation metrics from multiple linear regression model

1. The multiple linear regression model performed better in every metric relative to the simple linear regression model. Most notably, in the simple linear regression, only 38.9% of the variance in Temperature was described by the model. Whereas, in the multiple linear regression model, 86.6% of the variance in Temperature was explained by the combination of features.

The transparent nature of the intercept and beta coefficients make linear regression models very easy to interpret. In business, it is commonly requested that data scientists explain the effect of a certain feature on an outcome. Thus, linear regression provides metrics allowing a reasonable response to the business inquiry above. However, much of the time a problem requires the data scientist to predict an outcome measure that is not continuous, but categorical. For example, in insurance, given certain features of an insured what is the probability that this customer will not renew their policy? In this case, there is not a linear relationship between the features in the data and the outcome variable, so linear regression will falter. A viable option for conducting regression analysis on a categorical dependent variable is logistic regression.

**Discussion**: Is there something not discussed here that may be negatively affecting the performance of the multiple linear regression model? If so, what is it? And what can we do to prevent it?

**Answer**: Yes, there is something not discussed here that may be negatively affecting the performance of the multiple linear regression model. It is multicollinearity. Multicollinearity is a situation where features in the data are correlated to one another, making it difficult to discern the effects of those features on the outcome and ultimately weakening the statistical power of the model. Checking the correlation between predictors prior to including them in the model can help prevent multicollinearity. Or, performing supervised data compression using linear discriminant function analysis (LDA; Chapter 4) on the features prior to fitting a linear regression model will help us prevent multicollinearity.

Logistic regression

*Present x: Logistic regression*

Logistic regression uses categorical and continuous variables to predict a categorical outcome. When the dependent variable of choice has two categorical outcomes the analysis is termed binary logistic regression. However, if the outcome variable consists of more than 2 levels the analysis is referred to as multinomial logistic regression. For the purposes of this lesson we will focus our learning on the former.

When predicting a binary outcome, we do not have a linear relationship between the features and the outcome variable; an assumption of linear regression. Thus, to express a nonlinear relationship in a linear way we must transform the data using the logarithmic transformation. As a result, logistic regression allows us to predict the probability of the binary outcome occurring given the feature(s) in the model.

In logistic regression with 1 predictor, the logistic regression equation is shown below:

Equation 2.x: Logistic regression formula with 1 predictor

In equation 2.x, *P(Y)* is the probability of the outcome occurring, *e* is the base of natural logarithms, *α* is the intercept, *β* is the beta coefficient, and *x* is the value of the predictor. This equation can be extended to multiple predictors using the formula below:

Equation 2.x: Logistic regression formula with more than 1 predictor

Thus, using logistic regression to model the probability of an event occurring is the same as fitting a linear regression model except the continuous outcome variable has been replaced by the log odds of success for a binary outcome variable. In linear regression, we assumed a linear relationship between the predictor variable(s) and the outcome variable. Logistic regression, on the other hand, assumes a linear relationship between the predictor variable(s) and the natural log of *p*/(1-*p*), where *p* is the probability of the event occurring.

In the following exercise we will use the ‘weather.csv’ data set to demonstrate building a logistic regression model to predict the probability of rain.

Exercise 5: Fitting a logistic regression model and determining the intercept and coefficient

*Exercise 5: Fitting a logistic regression model and determining the intercept and coefficient*

To model the probability of rain (as opposed to snow), we will use the ‘weather.csv’ file as our data set, but instead of storing the continuous variable ‘Temperature\_c’ as the dependent variable, we will be storing the dichotomous variable ‘Rain’ as the outcome measure.

1. Import data using the following code:

import pandas as pd

df = pd.read\_csv('weather.csv')

Dummy code the ‘Summary’ variable as follows:

import pandas as pd

df\_dummies = pd.get\_dummies(df, drop\_first=True)

Shuffle df\_dummies using the code below:

from sklearn.utils import shuffle

df\_shuffled = shuffle(df\_dummies, random\_state=42)

Split the features and outcome into X and y, respectively as follows:

DV = 'Rain'

X = df\_shuffled.drop(DV, axis=1)

y = df\_shuffled[DV]

Split the features and outcome into training and testing data using the code below:

from sklearn.model\_selection import train\_test\_split

X\_train, X\_test, y\_train, y\_test = train\_test\_split(X, y, test\_size=0.33, random\_state=42)

1. Instantiate a logistic regression model using:

from sklearn.linear\_model import LogisticRegression

model = LogisticRegression()

1. Fit the logistic regression model to the training data using model.fit(X\_train, y\_train)

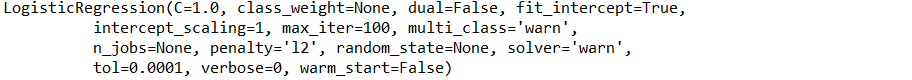


Figure 2.x: Output in console after fitting logistic regression model

1. Get the intercept using intercept = model.intercept\_
2. Extract the coefficients using coefficients = model.coef\_
3. Place the coefficients into a list using coef\_list = list(coefficients[0,:])
4. Match features to their coefficients, place them in a data frame, and print the data frame to the console as follows:

coef\_df = pd.DataFrame({'Feature': list(X\_train.columns),

'Coefficient': coef\_list})

print(coef\_df)

Refer to the resultant output below:



Figure 2.x: Features and their coefficients from logistic regression model

The coefficient for temperature can be interpreted as for every 1 degree increase in temperature, the log odds of rain increase by 5.69, controlling for all other features in the model. To generate predictions, we could convert the log odds to odds and odds to probability. However, scikit-learn has functionality to generate predicted probability as well as predicted classes.

Exercise 6: Generating predictions and evaluating performance of logistic regression model

*Exercise 6: Generating predictions and evaluating performance of multiple linear regression model*

In exercise 5 we learned how to fit a logistic regression model and extract the elements necessary to generate predictions. However, scikit-learn makes our lives much easier by providing us with functions to predict the probability of an outcome as well as the classes of an outcome. In this exercise, we will learn to generate predicted probabilities and classes as well as evaluating model performance using a confusion matrix and a classification report.

1. Generate predicted probabilities using predicted\_prob = model.predict\_proba(X\_test)[:,1]
2. Generate predicted classes using predicted\_class = model.predict(X\_test)
3. Evaluate a performance using a confusion matrix as follows:

from sklearn.metrics import confusion\_matrix

import numpy as np

cm = pd.DataFrame(confusion\_matrix(y\_test, predicted\_class))

cm['Total'] = np.sum(cm, axis=1)

cm = cm.append(np.sum(cm, axis=0), ignore\_index=True)

cm.columns = ['Predicted No', 'Predicted Yes', 'Total']

cm = cm.set\_index([['Actual No', 'Actual Yes', 'Total']])

print(cm)

Refer to the resultant output below:

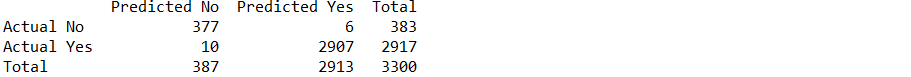


Figure 2.x: Confusion matrix from logistic regression model

From the confusion matrix, we can see that of the 383 observations that were not classified as rainy 377 of them were correctly classified and of the 2917 observations that were classified as rainy 2907 of them were correctly classified. To further inspect our model’s performance using metrics such as precision, recall, and f1-score we will generate a classification report.

1. Generate a classification report using the following code:

from sklearn.metrics import classification\_report

print(classification\_report(y\_test, predicted\_class))

Refer to the resultant output:

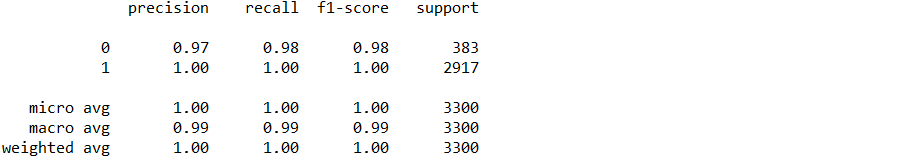


Figure 2.x: Classification report generated from logistic regression model

As we can see from our confusion matrix and classification report, our model is performing very well and may be difficult to improve upon. However, machine learning models including logistic regression consist of numerous hyperparameters that can be adjusted to further improve model performance. In the next exercise, we will learn to find the optimal combination of hyperparameters to maximize model performance.

Exercise 7: Tuning hyperparameters of logistic regression model

*Exercise 7: Tuning hyperparameters of multiple logistic regression model*

In step 3 of exercise 5, we fit a logistic regression model and the subsequent output from that model is displayed in Figure 2.x. Each of those arguments inside the LogisticRegression() function is set to a default hyperparameter. To tune the model, we will use scikit-learn’s grid search function which fits a model for every combination of possible hyperparameter values and sees the value for each hyperparameter resulting in the “best” model. In this exercise, we will learn how to use grid search to tune models.

Continuing from Exercise 6:

1. The data has already been prepared for us (see Exercise 6), thus we can jump right into instantiating a grid of possible hyperparameter values as follows:

import numpy as np

grid = {'penalty': ['l1', 'l2'],

'C': np.linspace(1, 10, 10),

'solver': ['liblinear']}

1. Instantiate a grid search model as follows:

from sklearn.model\_selection import GridSearchCV

from sklearn.linear\_model import LogisticRegression

model = GridSearchCV(LogisticRegression(), grid, scoring='f1', cv=5)

1. Fit the model on the training using model.fit(X\_train, y\_train) (keep in mind, this may take a while) and find the resultant output below:

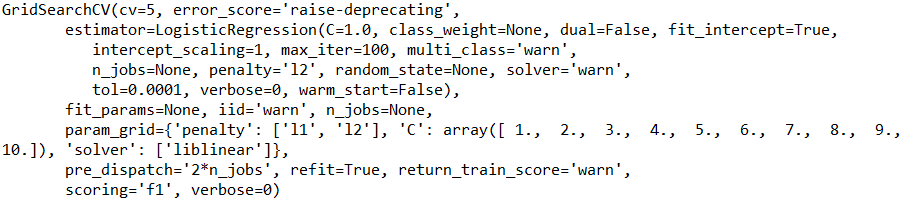


Figure 2.x: Output from grid search model

1. We can return the optimal combination of hyperparameters as a dictionary as follows:

best\_parameters = model.best\_params\_

print(best\_parameters)



Figure 2.x: Tuned hyperparameters from grid search model

We have found the combination of hyperparameters that maximize f1 score (i.e., the harmonic average of precision and recall). Remember, simply using the default hyperparamters in Exercise 5 resulted in a model that performed very well on the test data. Thus, in the following activity, we will evaluate how the model with tuned hyperparameters performed on the test data.

**Discussion**: Why is not every hyperparameter listed in the grid?

**Answer**: When not every hyperparameter is listed in the grid, it is because the user has decided to use the default hyperparameter in that case. Additionally, some hyperparameters will not work when other hyperparameters are selected simultaneously, so it is important to always refer and pay close attention to the algorithm’s documentation when setting up a grid for tuning.

Activity 2: Generating predictions and evaluating performance of tuned logistic regression model

*Activity 2: Evaluating tuned model performance*

Once the best combination of hyperparameters has been converged upon, we need to evaluate model performance much like we did in Exercise 5.

Continuing from Exercise 7:

1. Generate the predicted probabilities of rain
2. Generate the predicted class of rain
3. Evaluate performance with a confusion matrix and store it as a data frame
4. Print a classification report

**Solution:**

1. Generate predicted probabilities of rain using the following code:

predicted\_prob = model.predict\_proba(X\_test)[:,1]

1. Generate predicted class of rain using predicted\_class = model.predict(X\_test)
2. Evaluate performance using a confusion matrix and save it as a data frame using the following code:

from sklearn.metrics import confusion\_matrix

import numpy as np

cm = pd.DataFrame(confusion\_matrix(y\_test, predicted\_class))

cm['Total'] = np.sum(cm, axis=1)

cm = cm.append(np.sum(cm, axis=0), ignore\_index=True)

cm.columns = ['Predicted No', 'Predicted Yes', 'Total']

cm = cm.set\_index([['Actual No', 'Actual Yes', 'Total']])

print(cm)

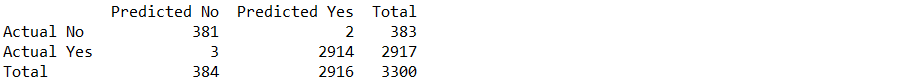


Figure 2.x: Confusion matrix from tuned logistic regression model

1. Nice! We have decreased our number of false positives from 6 to 2. Additionally, our false negatives were lowered from 10 to 3 (see Exercise 6). For further evaluation, print a classification report as follows:

from sklearn.metrics import classification\_report

print(classification\_report(y\_test, predicted\_class))

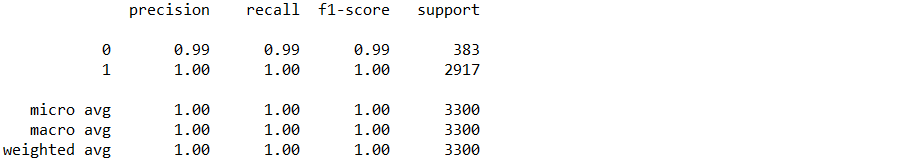


Figure 2.x: Classification report from tuned logistic regression model

By tuning the hyperparameters of the logistic regression model, we were able to improve upon a logistic regression model that was already performing very well. We will continue to expand upon tuning models in the following exercises and activities.

Classification using support vector machines (SVM)

*Present x: Max-Margin classification using support vector machines*

EXPLAIN SVM HERE…

Exercise 8: Preparing data for support vector classifier

*Exercise 8: Preparing data for support vector classifier*

1. Import ‘weather.csv’ using the following code:

import pandas as pd

df = pd.read\_csv('weather.csv')

1. Dummy code the categorical feature, 'Summary' as follows:

import pandas as pd

df\_dummies = pd.get\_dummies(df, drop\_first=True)

1. Shuffle df\_dummies to remove any ordering effects using the code below:

from sklearn.utils import shuffle

df\_shuffled = shuffle(df\_dummies, random\_state=42)

1. Slit df\_shuffled into X and y using the following code:

DV = 'Rain'

X = df\_shuffled.drop(DV, axis=1)

y = df\_shuffled[DV] # get DV (y)

1. Scale the features, X, by transforming them to z-scores as follows:

from sklearn.preprocessing import StandardScaler

model = StandardScaler()

X\_scaled = model.fit\_transform(X)

1. Split X\_scaled and y into testing and training data using the code below:

from sklearn.model\_selection import train\_test\_split

X\_train, X\_test, y\_train, y\_test = train\_test\_split(X\_scaled, y, test\_size=0.33, random\_state=42)

INSERT TRANSITION HERE…

Exercise 9: Tuning SVC model using grid search

*Exercise 9: Tuning SVC model using grid search*

INSERT INTRO HERE…

1. Instantiate grid for which to search using the following code:

import numpy as np

grid = {'C': np.linspace(1, 10, 10),

'kernel': ['linear', 'poly', 'rbf', 'sigmoid'],

'shrinking': [True, False],

'tol': np.linspace(0.001, 1, 10)}

1. Instantiate GridSearchCV model with the gamma hyperparameter set to ‘auto’ to avoid warnings and probability to ‘True’ so we can extract probability of rain as follows:

from sklearn.model\_selection import GridSearchCV

from sklearn.svm import SVC

model = GridSearchCV(SVC(gamma='auto', probability=True), grid, scoring='f1', cv=5)

1. Fit the gridsearch model using model.fit(X\_train, y\_train)

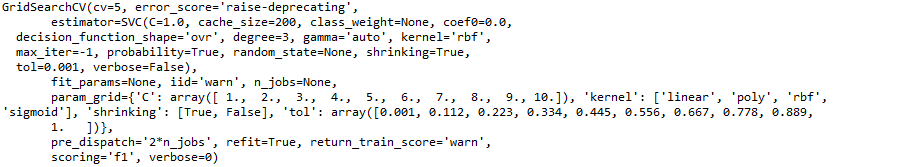


Figure 2.x: Output from fitting grid search model

1. Print the best parameters using the code below:

best\_parameters = model.best\_params\_

print(best\_parameters)

See the resultant output below:



Figure 2.x: Tuned hyperparameters for SVC model

INSERT TRANSITION HERE…

Activity 3: Generating predictions and evaluating performance of grid search SVC model

*Activity 3: Generating predictions and evaluating performance of grid search SVC model*

INSERT INTRO HERE…

1. Generate predicted probabilities
2. Extract predicted classes
3. Create and print a confusion matrix
4. Generate and print a classification report

**Solution:**

1. Generate predicted probabilities of rain using predicted\_prob = model.predict\_proba(X\_test)[:,1]
2. Extract predicted classes of rain using predicted\_class = model.predict(X\_test)
3. Create and print a confusion matrix using the code below:

from sklearn.metrics import confusion\_matrix

import numpy as np

cm = pd.DataFrame(confusion\_matrix(y\_test, predicted\_class))

cm['Total'] = np.sum(cm, axis=1)

cm = cm.append(np.sum(cm, axis=0), ignore\_index=True)

cm.columns = ['Predicted No', 'Predicted Yes', 'Total']

cm = cm.set\_index([['Actual No', 'Actual Yes', 'Total']])

print(cm)

See the resultant output below:



Figure 2.x: Confusion matrix from tuned SVC model

1. Generate and print a classification report as follows:

from sklearn.metrics import classification\_report

print(classification\_report(y\_test, predicted\_class))

See the resultant output below:



Figure 2.x: Classification report from tuned SVC model

INSERT TRANSITION HERE…

Decision trees

*Present x: Decision trees*

EXPLAIN DECISION TREES HERE…

Activity 4: Prepare data for decision tree classifier pipeline

*Activity 4: Prepare data for decision tree classifier pipeline*

1. Import ‘weather.csv’ and store it as a data frame using the following:

import pandas as pd

df = pd.read\_csv('weather.csv')

1. Dummy code the 'Summary' column as follows:

import pandas as pd

df\_dummies = pd.get\_dummies(df, drop\_first=True)

1. Shuffle df\_dummies using the code below:

from sklearn.utils import shuffle

df\_shuffled = shuffle(df\_dummies, random\_state=42)

1. Split df\_shuffled into X and y as follows:

DV = 'Rain'

X = df\_shuffled.drop(DV, axis=1)

y = df\_shuffled[DV]

1. Split X and y into testing and training data

from sklearn.model\_selection import train\_test\_split

X\_train, X\_test, y\_train, y\_test = train\_test\_split(X, y, test\_size=0.33, random\_state=42)

INSERT TRANSITION HERE…

Exercise 10: Tuning decision tree classifier using grid search in pipeline:

*Exercise 10: Tuning decision tree model*

INSERT INTRO HERE…

1. Set up the steps for a pipeline

from sklearn.preprocessing import StandardScaler

from sklearn.tree import DecisionTreeClassifier

steps = [('scaler', StandardScaler()), ('Tree', DecisionTreeClassifier())]

1. Setup the pipeline

from sklearn.pipeline import Pipeline

pipeline = Pipeline(steps)

1. Specify the hyperparameter space

import numpy as np

parameters = {'Tree\_\_criterion': ['gini', 'entropy'],

'Tree\_\_min\_weight\_fraction\_leaf': np.linspace(0.0, 0.5, 10),

'Tree\_\_min\_impurity\_decrease': np.linspace(0.0, 1.0, 10),

'Tree\_\_class\_weight': [None, 'balanced'],

'Tree\_\_presort': [True, False]}

1. Instantiate the GridSearchCV model

from sklearn.model\_selection import GridSearchCV

model = GridSearchCV(pipeline, parameters, scoring='f1', cv=5)

1. Fit to the training set

model.fit(X\_train, y\_train)

See the resultant output displayed below:

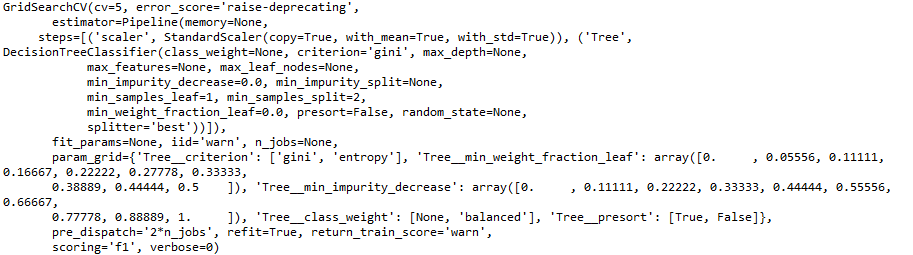


Figure 2.x: Output from fitting a grid search model using a pipeline

1. Print the tuned parameters

best\_parameters = model.best\_params\_

print(best\_parameters)

See the resultant output below:



Figure 2.x: Tuned hyperparameters for decision tree classifier

Remember, in practice, it is common for decision makers to ask how various features are affecting the predictions. In linear and logistic regression, the intercept and coefficient(s) make model predictions very transparent. However, many machine learning models are a black box where the effects of features on predictions is not made explicit. However, in tree-based models, scikit-learn provides an attribute termed ‘feature\_importances\_’ which returns an array containing values of relative feature importance for each feature. Conversely, this attribute is not available from a grid search model. In the next exercise, we will learn to programmatically extract values from the best\_parameters dictionary and re-fit the tuned decision tree model; allowing us to access the attributes provided by the decision tree classifier function.

Exercise 11: Programmatically extracting tuned hyperparameters from decision tree classifier grid search model

*Exercise 11: Programmatically extracting tuned hyperparameters*

In Exercise 10, we saved the tuned hyperparameters as key value pairs in the best\_parameters dictionary. This allows us to programmatically access the values and assign them to the appropriate hyperparameters of a decision tree classifier model. By fitting the tuned decision tree model, we will be able to access the attributes made available from the scikit-learn decision tree classifier function.

1. Print the best\_parameters dictionary using print(best\_parameters)

See the resultant output below:



Figure 2.x: best\_parameters dictionary

1. Prove that we can access the value for ‘Tree\_criterion’ using print(best\_parameters['Tree\_\_criterion'])

See the resultant output below:



Figure 2.x: Value assigned to the ‘Tree\_criterion’ key in the best\_parameters dictionary

1. Instantiate decision tree classifier model and assign the values to the corresponding hyperparameters as follows:

from sklearn.tree import DecisionTreeClassifier

model = DecisionTreeClassifier(class\_weight=best\_parameters['Tree\_\_class\_weight'],

criterion=best\_parameters['Tree\_\_criterion'],

min\_impurity\_decrease=best\_parameters['Tree\_\_min\_impurity\_decrease'],

min\_weight\_fraction\_leaf=best\_parameters['Tree\_\_min\_weight\_fraction\_leaf'],

presort=best\_parameters['Tree\_\_presort'])

1. Fit the model to the training data using model.fit(X\_train, y\_train)

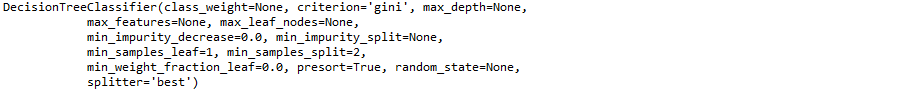


Figure 2.x: Output from fitting the decision tree classifier model with tuned hyperparameters

1. Extract feature\_importances attribute using print(model.feature\_importances\_)

The resultant output is shown below:



Figure 2.x: Array of feature importance from tuned decision tree classifier model

1. From the array in Figure 2.x, we can see that the first feature completely dominated the other variables in terms of feature importance. Visualize this using the code below:

import pandas as pd

import matplotlib.pyplot as plt

df\_imp = pd.DataFrame({'Importance': list(model.feature\_importances\_)}, index=X.columns)

df\_imp\_sorted = df\_imp.sort\_values(by=('Importance'), ascending=True)

df\_imp\_sorted.plot.barh(figsize=(5,5))

plt.title('Relative Feature Importance')

plt.xlabel('Relative Importance')

plt.ylabel('Variable')

plt.legend(loc=4)

plt.show()

See the resultant output below:

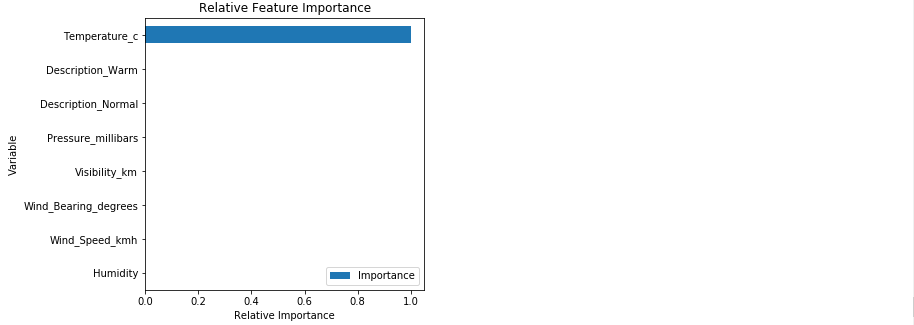


Figure 2.x: Feature importance from tuned decision tree classifier model

It looks like temperature in Celsius was the sole driver in this classification problem. With the outcome measure being rain (‘Rain’=1) or snow (‘Rain’=0) and the way in which decision trees make value threshold decisions, it makes sense that the algorithm used temperature to determine if there was rainfall or snowfall at the time of measurement. In the upcoming activity, we will evaluate how the model performed.

Activity 5: Generating predictions and evaluating performance of decision tree classifier model

*Activity 5: Generating predictions and evaluating performance of decision tree classifier model*

We have generated predictions and evaluated model performance in previous exercises and activities. We will be taking the same approach in this activity to evaluate the performance of our tuned decision tree classifier model.

1. Generate predicted probabilities of rain
2. Generate predicted classes of rain
3. Generate and print a confusion matrix
4. Print a classification report

**Solution:**

1. Generate predicted probabilities of rain using predicted\_prob = model.predict\_proba(X\_test)[:,1]
2. Generate predicted classes of rain using predicted\_class = model.predict(X\_test)
3. Generate and print a confusion matrix with the code below:

from sklearn.metrics import confusion\_matrix

import numpy as np

cm = pd.DataFrame(confusion\_matrix(y\_test, predicted\_class))

cm['Total'] = np.sum(cm, axis=1)

cm = cm.append(np.sum(cm, axis=0), ignore\_index=True)

cm.columns = ['Predicted No', 'Predicted Yes', 'Total']

cm = cm.set\_index([['Actual No', 'Actual Yes', 'Total']])

print(cm)

Refer to the resultant output below:



Figure 2.x: Confusion matrix from tuned decision tree classifier model

1. Print a classification report as follows:

from sklearn.metrics import classification\_report

print(classification\_report(y\_test, predicted\_class))

Refer to the resultant output below:



Figure 2.x: Classification report from tuned decision tree classifier model

By tuning a decision tree classifier model on our ‘weather.csv’ dataset, we were able to predict rain (or snow) almost perfectly. We are able to see that the sole driving feature was temperature in celsius. This makes sense due to the way in which decision trees use recursive partitioning to make predictions.

Sometimes, after evaluation, a single model is a weak learner and does not perform well. However, by combining weak learners, we create a stronger learner. The approach of combining numerous weak learners to create a stronger learner is termed ensemble. Random forest models combine numerous decision tree models to create a stronger, ensemble model. Decision trees can be used for classification or regression problems, thus random forests can be used for classification or regression problems. In the upcoming exercises and activities, we will tune and evaluate a random forest regressor in a pipeline to predict temperature in Celsius from the features in ‘weather.csv’.

Random forests

*Present x: Random forests*

DESCRIBE RANDOM FORESTS HERE…

Exercise 12: Preparing data for random forest regressor pipeline

*Exercise 12: Preparing data for random forest regressor pipeline*

INSERT INTRO HERE…

1. Import ‘weather.csv’ and save it as df using the following code:

import pandas as pd

df = pd.read\_csv('weather.csv')

1. Dummy code the categorical variables as follows:

import pandas as pd

df\_dummies = pd.get\_dummies(df, drop\_first=True)

1. Remove any possible ordering effects by shuffling df\_dummies using the code below:

from sklearn.utils import shuffle

df\_shuffled = shuffle(df\_dummies, random\_state=42)

1. Split df\_shuffled into X and y using the following code:

DV = 'Temperature\_c'

X = df\_shuffled.drop(DV, axis=1)

y = df\_shuffled[DV]

1. Split X and y into testing and training data as follows:

from sklearn.model\_selection import train\_test\_split

X\_train, X\_test, y\_train, y\_test = train\_test\_split(X, y, test\_size=0.33, random\_state=42)

Now that we have imported, shuffled, separated our data into features (X) and dependent variable (y), and split X and y into testing and training data we will scale X and tune a random forest regressor model in a pipeline.

Activity 6: Scaling features and tuning a random forest regressor in a pipeline

*Activity 6: Scaling features and tuning a random forest regressor in a pipeline*

INSERT INTRO HERE…

1. Set up the steps for a pipeline
2. Setup the pipeline
3. Specify the hyperparameter space
4. Instantiate the GridSearchCV model optimizing the explained variance
5. Fit the grid search model to the training set
6. Print the tuned parameters

**Solution:**

1. Set up the steps for a pipeline using the following code:

from sklearn.preprocessing import StandardScaler

from sklearn.ensemble import RandomForestRegressor

steps = [('scaler', StandardScaler()), ('Forest', RandomForestRegressor(n\_estimators=10))]

1. Setup the pipeline using the code below:

from sklearn.pipeline import Pipeline

pipeline = Pipeline(steps)

1. Specify the hyperparameter space as follows:

import numpy as np

parameters = {'Forest\_\_criterion': ['mse','mae'],

'Forest\_\_max\_features': ['auto', 'sqrt', 'log2', None],

'Forest\_\_min\_impurity\_decrease': np.linspace(0.0, 1.0, 10),

'Forest\_\_bootstrap': [True, False],

'Forest\_\_warm\_start': [True, False]}

1. Instantiate the GridSearchCV model optimizing the explained variance using the following code:

from sklearn.model\_selection import GridSearchCV

model = GridSearchCV(pipeline, parameters, scoring='explained\_variance', cv=5)

1. Fit the grid search model to the training set using model.fit(X\_train, y\_train)
2. Print the tuned parameters as follows:

best\_parameters = model.best\_params\_

print(best\_parameters)

See the resultant output below:



Figure 2.x: Tuned hyperparameters from random forest regressor grid search model

INSERT TRANSITION HERE…

Exercise 13: Programmatically extracting tuned hyperparameters and determining feature importance from random forest regressor grid search model

*Exercise 13: Programmatically extracting tuned hyperparameters from random forest regressor grid search model*

INSERT INTRO HERE…

1. Instantiate a random forest regressor model with the values for each key from the best\_parameters disctionary assigned to the corresponding hyperparameter.

from sklearn.ensemble import RandomForestRegressor

model = RandomForestRegressor(criterion=best\_parameters['Forest\_\_criterion'],

max\_features=best\_parameters['Forest\_\_max\_features'],

min\_impurity\_decrease=best\_parameters['Forest\_\_min\_impurity\_decrease'],

bootstrap=best\_parameters['Forest\_\_bootstrap'],

warm\_start=best\_parameters['Forest\_\_warm\_start'])

1. Fit the model on the training data using model.fit(X\_train, y\_train)

Find the resultant output below:

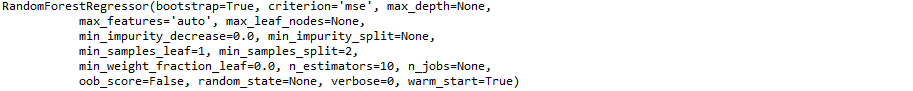


Figure 2.x: Output from fitting the random forest regressor model with tuned hyperparameters

1. Plot feature importance in descending order using the following code:

import pandas as pd

import matplotlib.pyplot as plt

df\_imp = pd.DataFrame({'Importance': list(model.feature\_importances\_)}, index=X.columns)

df\_imp\_sorted = df\_imp.sort\_values(by=('Importance'), ascending=True)

df\_imp\_sorted.plot.barh(figsize=(5,5))

plt.title('Relative Feature Importance')

plt.xlabel('Relative Importance')

plt.ylabel('Variable')

plt.legend(loc=4)

plt.show()

See the resultant output below:

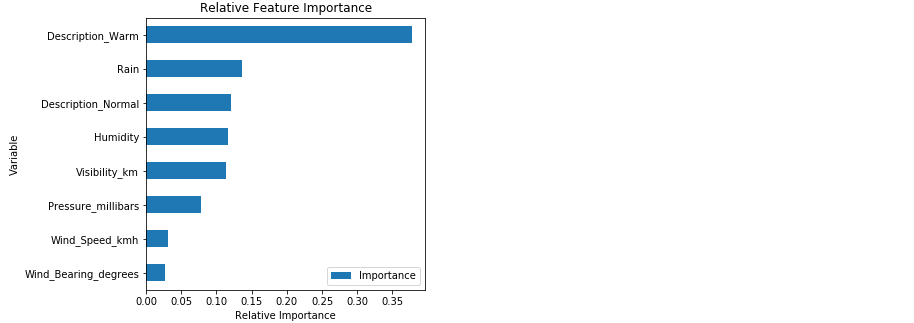


Figure 2.x: Feature importance from tuned random forest regressor model

INSERT TRANSITION HERE…

Activity 7: Generating predictions and evaluating performance of tuned random forest regressor model

*Present x: Generating predictions and evaluating performance of tuned random forest regressor model*

INSERT INTRO HERE…

Continuing from Exercise 13:

1. Generate predictions on the test data
2. Plot correlation of predicted and actual values
3. Plot distribution of residuals
4. Compute metrics, place them in a dataframe, and print it

**Solution:**

1. Generate predictions on the test data using predictions = model.predict(X\_test)
2. Plot correlation of predicted and actual values using the following code:

import matplotlib.pyplot as plt

from scipy.stats import pearsonr

plt.scatter(y\_test, predictions)

plt.xlabel('Y Test (True Values)')

plt.ylabel('Predicted Values')

plt.title('Predicted vs. Actual Values (r = {0:0.2f})'.format(pearsonr(y\_test, predictions)[0], 2))

plt.show()

Refer to the resultant output below:

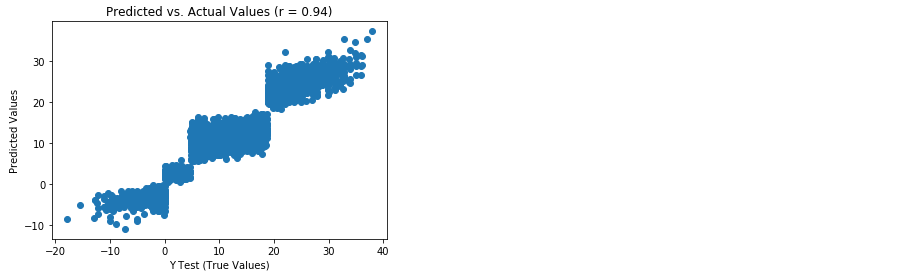


Figure 2.x: Scatterplot of predicted and actual values from tuned random forest regression model

1. Plot distribution of residuals as follows:

import seaborn as sns

from scipy.stats import shapiro

sns.distplot((y\_test - predictions), bins = 50)

plt.xlabel('Residuals')

plt.ylabel('Density')

plt.title('Histogram of Residuals (Shapiro W p-value = {0:0.3f})'.format(shapiro(y\_test - predictions)[1]))

plt.show()

Refer to the resultant output below:

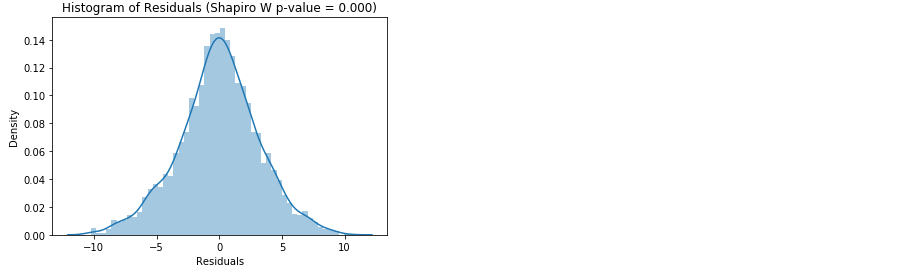


Figure 2.x: Histogram of residuals from tuned random forest regression model

1. Compute metrics, place them in a dataframe, and print it using the code below:

from sklearn import metrics

import numpy as np

metrics\_df = pd.DataFrame({'Metric': ['MAE',

'MSE',

'RMSE',

'R-Squared'],

'Value': [metrics.mean\_absolute\_error(y\_test, predictions),

metrics.mean\_squared\_error(y\_test, predictions),

np.sqrt(metrics.mean\_squared\_error(y\_test, predictions)),

metrics.explained\_variance\_score(y\_test, predictions)]}).round(3)

print(metrics\_df)

Find the resultant output below:



Figure 2.x: Model evaluation metrics from tuned random forest regression model

The random forest regressor model seems to outperform the multiple linear regression as evidenced by lower MAE, MSE, and RMSE as well as greater explained variance. ELABORATE ON THIS…

Summary

*Present x: Summary*

INSERT SUMMARY HERE…

Assessment Questions:

INSERT ASSESSMENT QUESTIONS HERE…