Project: Predicting Heart Disease with Classification Machine Learning Algorithms

Table of Contents

1. Introduction:

Scenario

Goal

Features & Predictor

- 2. Data Wrangling
- 3. Exploratory Data Analysis:

Correlations

Violin & Box Plots

Filtering data by positive & negative Heart Disease patient

4. Machine Learning + Predictive Analytics:

Prepare Data for Modeling

Modeling/Training

Making the Confusion Matrix

Feature Importance

Predictions

5. Conclusions

1. Introduction

Scenario:

You have just been hired at a Hospital with an alarming number of patients coming in reporting various cardiac symptoms. A cardioligist measures vitals & hands you this data to perform Data Analysis and predict whether certain patients have Heart Disease.

Goal:

```
-To predict whether a patient should be diagnosed with Heart Disea se. This is a binary outcome.
```

```
Positive (+) = 1, patient diagnosed with Heart Disease
```

Negative (-) = 0, patient not diagnosed with Heart Disease

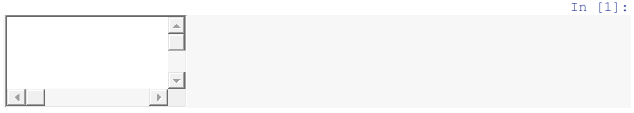
-To experiment with various Classification Models & see which yiel ds greatest accuracy.

- Examine trends & correlations within our data
- determine which features are important in determing Positive/Neg ative Heart Disease

Features & Predictor:

Our Predictor (Y, Positive or Negative diagnosis of Heart Disease) is determined by 13 features (X):

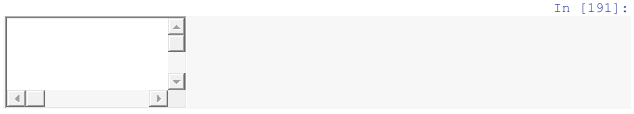
- 1. age (#)
- 2. sex: 1= Male, 0= Female (Binary)
- 3. (cp)chest pain type (4 values -Ordinal):Value 1: typical angina ,Value 2: atypical angina, Value 3: non-anginal pain , Value 4: asymptomatic (
- 4. (trestbps) resting blood pressure (#)
- 5. (chol) serum cholestoral in mg/dl (#)
- 6. (fbs)fasting blood sugar > 120 mg/dl(Binary)(1 = true; 0 = false)
- 7. (restecg) resting electrocardiographic results(values 0,1,2)
- 8. (thalach) maximum heart rate achieved (#)
- 9. (exang) exercise induced angina (binary) (1 = yes; 0 = no)
- 10. (oldpeak) = ST depression induced by exercise relative to rest (#)
- 11. (slope) of the peak exercise ST segment (Ordinal) (Value 1: upsloping , Value 2: flat , Value 3: downsloping)
- 12. (ca) number of major vessels (0-3, Ordinal) colored by fluoroscopy
- 13. (thal) maximum heart rate achieved (Ordinal): 3 = normal; 6 = fixed defect; 7 = reversable defect



import numpy as np

import pandas as pd
import matplotlib as plt
import seaborn as sns
import matplotlib.pyplot as plt

2. Data Wrangling



filePath = '/Users/jarar_zaidi/Downloads/datasets-33180-43520-heart.csv'

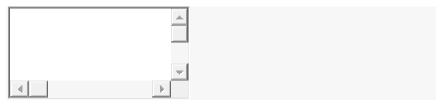
data = pd.read_csv(filePath)

data.head(5)

Out[191]:

In [194]:

	age	sex	сp	trestbps	chol	fbs	restecg	thalach	exang	oldpeak	slope	ca	thal	target
0	63	1	3	145	233	1	0	150	0	2.3	0	0	1	1
1	37	1	2	130	250	0	1	187	0	3.5	0	0	2	1
2	41	0	1	130	204	0	0	172	0	1.4	2	0	2	1
3	56	1	1	120	236	0	1	178	0	0.8	2	0	2	1
4	57	0	0	120	354	0	1	163	1	0.6	2	0	2	1



print("(Rows, columns): " + str(data.shape))

data.columns

```
(Rows, columns): (303, 14)
```



data.nunique(axis=0)# returns the number of unique values for each variable.

Out[195]:

age	41
sex	2
ср	4
trestbps	49
chol	152
fbs	2
restecg	3
thalach	91
exang	2
oldpeak	40
slope	3
ca	5
thal	4
target	2
dtype: int64	

In [7]:



#summarizes the count, mean, standard deviation, min, and max for numeric variables.

data.describe()

Out[7]:

	age	sex	ср	trest bps	chol	fbs	reste cg	thala ch	exan g	oldp eak	slope	ca	thal	targ et
co un t	303. 0000 00													
m ea n	54.3 6633 7	0.68 3168	0.96 6997	131. 6237 62	246. 2640 26	0.14 8515	0.52 8053	149. 6468 65	0.32 6733	1.03 9604	1.39 9340	0.72 9373	2.31 3531	0.54 4554
st d	9.08 2101	0.46 6011	1.03 2052	17.5 3814 3	51.8 3075 1	0.35 6198	0.52 5860	22.9 0516 1	0.46 9794	1.16 1075	0.61 6226	1.02 2606	0.61 2277	0.49 8835

	age	sex	ср	trest bps	chol	fbs	reste cg	thala ch	exan g	oldp eak	slope	ca	thal	targ et
mi n	29.0 0000 0	0.00	0.00	94.0 0000 0	126. 0000 00	0.00	0.00	71.0 0000 0	0.00	0.00	0.00 0000	0.00	0.00	0.00 0000
25 %	47.5 0000 0	0.00	0.00 0000	120. 0000 00	211. 0000 00	0.00	0.00	133. 5000 00	0.00	0.00	1.00 0000	0.00	2.00 0000	0.00 0000
50 %	55.0 0000 0	1.00 0000	1.00 0000	130. 0000 00	240. 0000 00	0.00 0000	1.00 0000	153. 0000 00	0.00 0000	0.80 0000	1.00 0000	0.00	2.00 0000	1.00 0000
75 %	61.0 0000 0	1.00 0000	2.00 0000	140. 0000 00	274. 5000 00	0.00	1.00 0000	166. 0000 00	1.00 0000	1.60 0000	2.00 0000	1.00 0000	3.00 0000	1.00 0000
m ax	77.0 0000 0	1.00 0000	3.00 0000	200. 0000 00	564. 0000 00	1.00 0000	2.00 0000	202. 0000 00	1.00 0000	6.20 0000	2.00 0000	4.00 0000	3.00 0000	1.00 0000

Luckily we have no missing data to handle!

In [199]:

Display the Missing Values

print(data.isna().sum())

0 age 0 sex 0 ср trestbps 0 chol fbs restecg 0 thalach exang 0 oldpeak slope 0 са 0 thal 0 target dtype: int64

Let's see if theirs a good proportion between our positive and negative results. It appears we have a good balance between the two.



data['target'].value_counts()

Out[9]:

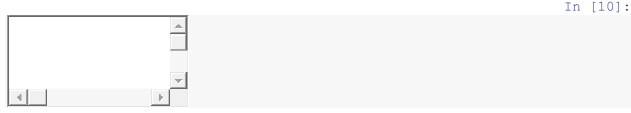
1 165
 138

Name: target, dtype: int64

3. Exploratory Data Analysis

Correlations

Correlation Matrix- let's you see correlations between all variables. Within seconds, you can see whether something is positivly or negativly correlated with our predictor (target)



calculate correlation matrix

```
corr = data.corr()
plt.subplots(figsize=(15,10))
sns.heatmap(corr, xticklabels=corr.columns, yticklabels=corr.columns, annot=True, cmap=sns.diverging_pal
ette(220, 20, as_cmap=True))
sns.heatmap(corr, xticklabels=corr.columns,
    yticklabels=corr.columns,
    annot=True,
    cmap=sns.diverging_palette(220, 20, as_cmap=True))
Out[10]:
<matplotlib.axes. subplots.AxesSubplot at 0x7f9c9bf367d0>
```

age -	1	-0.098	-0.069	0.28	0.21	0.12	-0.12	-0.4	0.097	0.21	-0.17	0.28	0.06
sex -	-0.098	1	-0.049	-0.057	-0.2	0.045	-0.058	-0.044	0.14	0.096	-0.031	0.12	0.2
cp -	-0.069	-0.049	1	0.048	-0.077	0.094	0.044	0.3	-0.39	-0.15	0.12	-0.18	-0.1
trestbps -	0.28	-0.057	0.048	1	0.12	0.18	-0.11	-0.047	0.068	0.19	-0.12	0.1	0.06
chol -	0.21	-0.2	-0.077	0.12	1	0.013	-0.15	-0.0099	0.067	0.054	-0.004	0.071	0.09
fbs -	0.12	0.045	0.094	0.18	0.013	1	-0.084	-0.0086	0.026	0.0057	-0.06	0.14	-0.03
restecg -	-0.12	-0.058	0.044	-0.11	-0.15	-0.084	1	0.044	-0.071	-0.059	0.093	-0.072	-0.01
thalach -	-0.4	-0.044	0.3	-0.047	-0.0099	-0.0086	0.044	1	-0.38	-0.34	0.39	-0.21	-0.09
exang -	0.097	0.14	-0.39	0.068	0.067	0.026	-0.071	-0.38	1	0.29	-0.26	0.12	0.23
oldpeak -	0.21	0.096	-0.15	0.19	0.054	0.0057	-0.059	-0.34	0.29	1	-0.58	0.22	0.2
slope -	-0.17	-0.031	0.12	-0.12	-0.004	-0.06	0.093	0.39	-0.26	-0.58	1	-0.08	-0.1
ca -	0.28	0.12	-0.18	0.1	0.071	0.14	-0.072	-0.21	0.12	0.22	-0.08	1	0.1
thal -	0.068	0.21	-0.16	0.062	0.099	-0.032	-0.012	-0.096	0.21	0.21	-0.1	0.15	1
target -	-0.23	-0.28	0.43	-0.14	-0.085	-0.028	0.14	0.42	-0.44	-0.43	0.35	-0.39	-0.3
'	age	sex	ф	restbps	chol	fbs	restecg	thalach	exang	oldpeak	slope	ca	tha

We can see there is a positive correlation between chest pain (cp) & target (our predictor). This makes sense since, The greater amount of chest pain results in a greater chance of having heart disease. Cp (chest pain), is a ordinal feature with 4 values: Value 1: typical angina ,Value 2: atypical angina, Value 3: non-anginal pain , Value 4: asymptomatic.

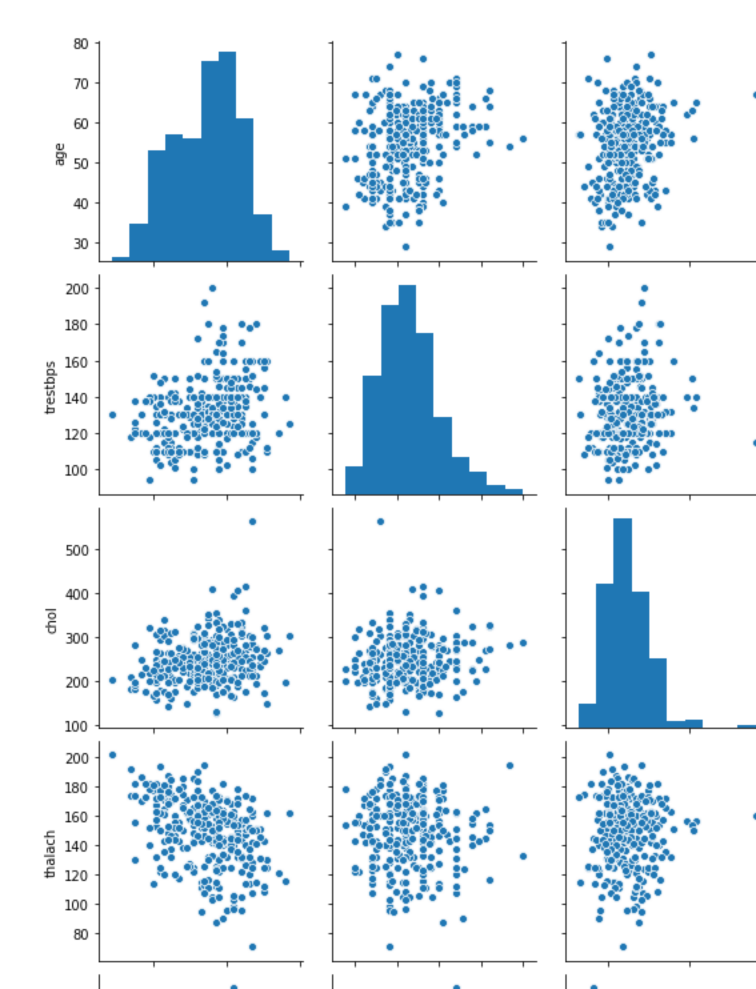
In addition, we see a negative correlation between exercise induced angina (exang) & our predictor. This makes sense because when you excercise, your heart requires more blood, but narrowed arteries slow down blood flow.

Pairplots are also a great way to immediatly see the correlations between all variables. But you will see me make it with only continous columns from our data, because with so many features, it can be difficult to see each one. So instead I will make a pairplot with only our continous features.



Out[11]:

<seaborn.axisgrid.PairGrid at 0x7f9c9bf1f410>



Chose to make a smaller pairplot with only the continus variables, to dive deeper into the relationships. Also a great way to see if theirs a positve or negative correlation!



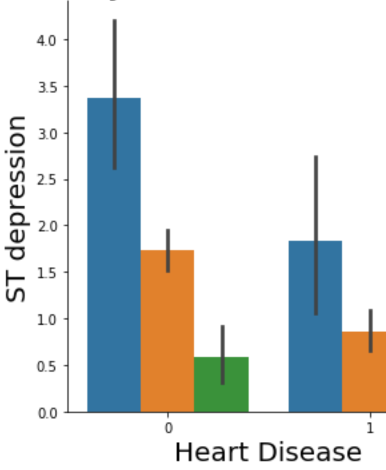
sns.catplot(x="target", y="oldpeak", hue="slope", kind="bar", data=data);

plt.title('ST depression (induced by exercise relative to rest) vs. Heart Disease', size=25) plt.xlabel('Heart Disease',size=20) plt.ylabel('ST depression',size=20)

Text(26.42645833333333343, 0.5, 'ST depression')

Out[12]:

ST depression (induced by exercise relative



ST segment depression occurs because when the ventricle is at rest and therefore repolarized. If the trace in the ST segment is abnormally low below the baseline, this can lead to this Heart Disease. This is supports the plot above because low ST Depression yields people at greater risk for heart disease. While a high ST depression is considered normal & healthy. The "slope" hue, refers to the

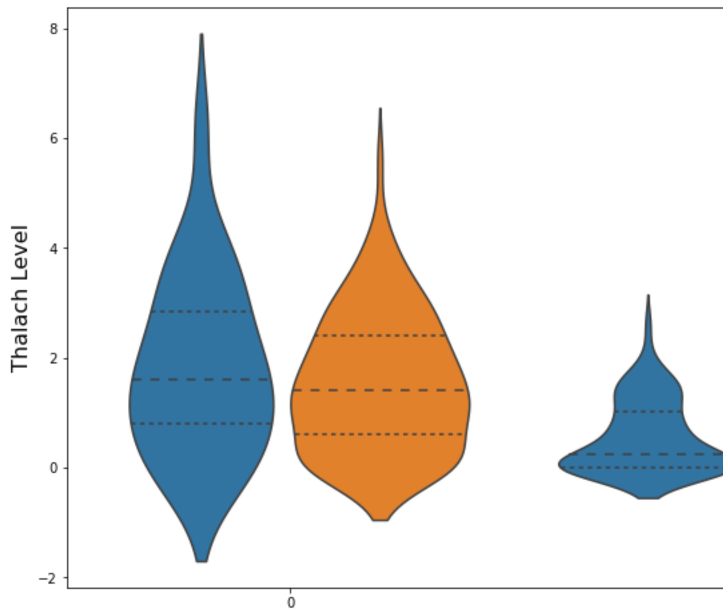
peak exercise ST segment, with values: 0: upsloping, 1: flat, 2: downsloping). Both positive & negative heart disease patients exhibit equal distributions of the 3 slope categories.

Violin & Box Plots

The advantages of showing the Box & Violin plots is that it showsthe basic statistics of the data, as well as its distribution. These plots are often used to compare the distribution of a given variable across some categories. It shows the median, IQR, & Tukey's fence. (minimum, first quartile (Q1), median, third quartile (Q3), and maximum). In addition it can provide us with outliers in our data.

```
plt.figure(figsize=(12,8))
sns.violinplot(x= 'target', y= 'oldpeak',hue="sex", inner='quartile',data= data)
plt.title("Thalach Level vs. Heart Disease",fontsize=20)
plt.xlabel("Heart Disease Target", fontsize=16)
plt.ylabel("Thalach Level", fontsize=16)
Out [156]:
Text(0, 0.5, 'Thalach Level')
```

Thalach Level vs. Heart Disease



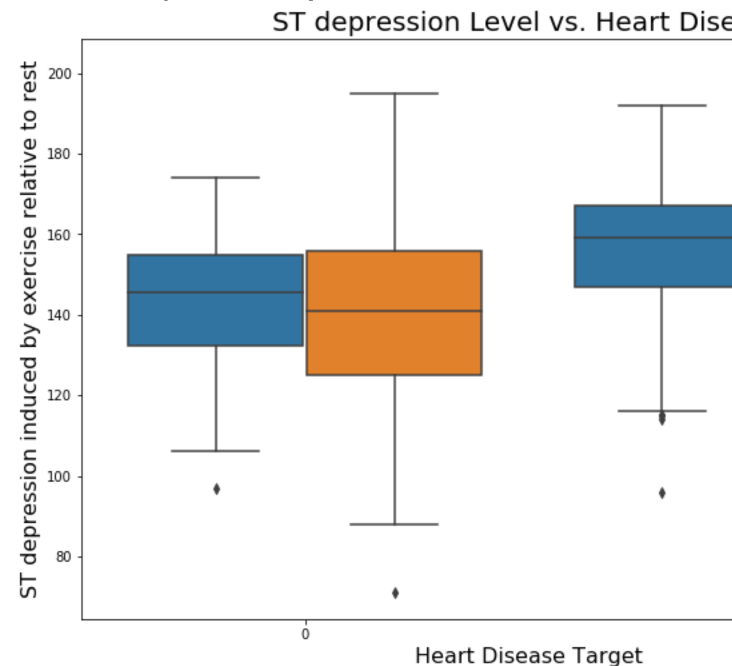
Heart Disease Target

We can see that the overall shape & distribution for negative & positive patients differ vastly. Positive patients exhibit a lower median for ST depression level & thus a great distribution of their data is between 0 & 2, while negative patients are between 1 & 3. In addition, we dont see many differences between male & female target outcomes.



```
plt.figure(figsize=(12,8))
sns.boxplot(x= 'target', y= 'thalach',hue="sex", data=data )
plt.title("ST depression Level vs. Heart Disease", fontsize=20)
plt.xlabel("Heart Disease Target",fontsize=16)
plt.ylabel("ST depression induced by exercise relative to rest", fontsize=16)
Out [14]:
```

Text(0, 0.5, 'ST depression induced by exercise relative to rest')



Positive patients exhibit a hightened median for ST depression level, while negative patients have lower levels. In addition, we dont see many differences between male & female target outcomes, expect for the fact that males have slightly larger ranges of ST Depression.

Filtering data by positive & negative Heart Disease patient

In [15]:

Filtering data by positive Heart Disease patient

pos_data = data[data['target']==1]
pos_data.describe()

													ouc []	. 0] •
	age	sex	ср	trest bps	chol	fbs	reste cg	thala ch	exan g	oldp eak	slope	ca	thal	ta rg et
co un t	165.0 0000 0	16 5. 0												
m ea n	52.49 6970	0.563 636	1.375 758	129.3 0303 0	242.2 3030 3	0.139 394	0.593 939	158.4 6666 7	0.139 394	0.583 030	1.593 939	0.363 636	2.121 212	1. 0
st d	9.550 651	0.497 444	0.952 222	16.16 9613	53.55 2872	0.347 412	0.504 818	19.17 4276	0.347 412	0.780 683	0.593 635	0.848 894	0.465 752	0. 0
mi n	29.00 0000	0.000	0.000	94.00 0000	126.0 0000 0	0.000	0.000	96.00 0000	0.000	0.000	0.000	0.000	0.000	1. 0
25 %	44.00 0000	0.000	1.000 000	120.0 0000 0	208.0 0000 0	0.000	0.000	149.0 0000 0	0.000	0.000	1.000	0.000	2.000 000	1. 0
50 %	52.00 0000	1.000 000	2.000 000	130.0 0000 0	234.0 0000 0	0.000	1.000 000	161.0 0000 0	0.000	0.200 000	2.000 000	0.000	2.000 000	1. 0
75 %	59.00 0000	1.000 000	2.000	140.0 0000 0	267.0 0000 0	0.000	1.000	172.0 0000 0	0.000	1.000 000	2.000	0.000	2.000 000	1. 0
m ax	76.00 0000	1.000 000	3.000 000	180.0 0000 0	564.0 0000 0	1.000 000	2.000 000	202.0 0000 0	1.000 000	4.200 000	2.000	4.000 000	3.000 000	1. 0

Filtering data by negative Heart Disease patient

In [16]:

Out[15]:



Filtering data by negative Heart Disease patient

neg_data = data[data['target']==0]
neg_data.describe()

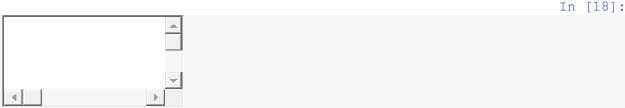
Out[16]:

In [17]:

	age	sex	ср	trest bps	chol	fbs	reste cg	thala ch	exan g	oldp eak	slope	ca	thal	ta rg et
co un t	138.0 0000 0	13 8. 0												
m ea n	56.60 1449	0.826 087	0.478 261	134.3 9855 1	251.0 8695 7	0.159 420	0.449 275	139.1 0144 9	0.550 725	1.585 507	1.166 667	1.166 667	2.543 478	0. 0
st d	7.962 082	0.380 416	0.905 920	18.72 9944	49.45 4614	0.367 401	0.541 321	22.59 8782	0.499 232	1.300 340	0.561 324	1.043 460	0.684 762	0. 0
mi n	35.00 0000	0.000	0.000	100.0 0000 0	131.0 0000 0	0.000	0.000	71.00 0000	0.000	0.000	0.000	0.000	0.000	0. 0
25 %	52.00 0000	1.000	0.000	120.0 0000 0	217.2 5000 0	0.000	0.000	125.0 0000 0	0.000	0.600 000	1.000	0.000	2.000	0. 0
50 %	58.00 0000	1.000 000	0.000	130.0 0000 0	249.0 0000 0	0.000	0.000	142.0 0000 0	1.000 000	1.400 000	1.000 000	1.000 000	3.000 000	0. 0
75 %	62.00 0000	1.000	0.000	144.7 5000 0	283.0 0000 0	0.000	1.000 000	156.0 0000 0	1.000 000	2.500 000	1.750 000	2.000	3.000 000	0. 0
m ax	77.00 0000	1.000	3.000 000	200.0 0000 0	409.0 0000 0	1.000	2.000	195.0 0000 0	1.000 000	6.200 000	2.000	4.000 000	3.000 000	0. 0



print("(Positive Patients ST depression): " + str(pos_data['oldpeak'].mean()))
print("(Negative Patients ST depression): " + str(neg_data['oldpeak'].mean()))



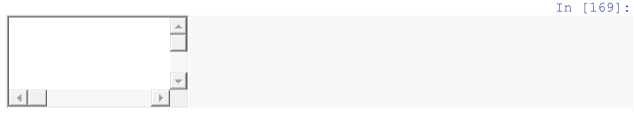
print("(Positive Patients thalach): " + str(pos_data['thalach'].mean()))
print("(Negative Patients thalach): " + str(neg_data['thalach'].mean()))
(Positive Patients thalach): 158.46666666666667
(Negative Patients thalach): 139.1014492753623

From comparing positive and negative patients we can see there are vast differenes in means for many of our Features. From examing the details, we can observe that positive patients experience heightened maximum heart rate achieved (thalach) average. In addition, positive patients exhibit about 1/3rd the amount of ST depression induced by exercise relative to rest (oldpeak).

4. Machine Learning + Predictive Analytics

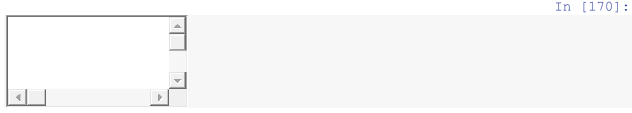
Prepare Data for Modeling

Assign the 13 features to X, & the last column to our classification predictor, y



X = data.iloc[:, :-1].values
y = data.iloc[:, -1].values

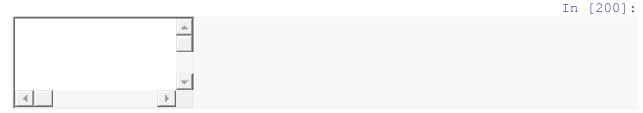
Split: the dataset into the Training set and Test set



from sklearn.model_selection import train_test_split

x_train, x_test, y_train, y_test = train_test_split(X,y,test_size = 0.2, random_state = 1)

Normalize: Standardizing the data will transform the data so that its distribution will have a mean of 0 and a standard deviation of 1.



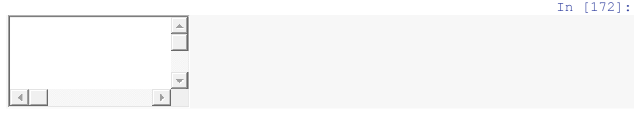
from sklearn.preprocessing import StandardScaler

sc = StandardScaler()
x_train = sc.fit_transform(x_train)
x_test = sc.transform(x_test)

Modeling / Training

We will now Train various Classification Models on the Training set & see which yields the highest accuracy. We will compare the accuracy of Logistic Regression, K-NN, SVM, Naives Bayes Classifier, Decision Trees, Random Forest, and XGBoost. Note: these are all supervised learning models.

Model 1: Logistic Regression



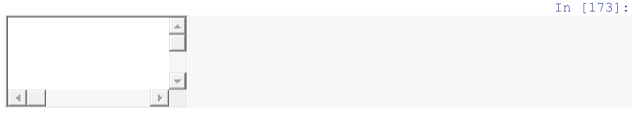
from sklearn.metrics import classification_report
from sklearn.linear_model import LogisticRegression

model1 = LogisticRegression(random_state=1) # get instance of model
model1.fit(x_train, y_train) # Train/Fit model

y_pred1 = model1.predict(x_test) # get y predictions
print(classification_report(y_test, y_pred1)) # output accuracy

	precision	recall	f1-score	support
0	0.77	0.67	0.71	30
1	0.71	0.81	0.76	31
accuracy			0.74	61
macro avg	0.74	0.74	0.74	61
weighted avg	0.74	0.74	0.74	61

Model 2: K-NN (K-Nearest Neighbors)



from sklearn.metrics **import** classification_report **from** sklearn.neighbors **import** KNeighborsClassifier

model2 = KNeighborsClassifier() # get instance of model
model2.fit(x_train, y_train) # Train/Fit model

y_pred2 = model2.predict(x_test) # get y predictions print(classification_report(y_test, y_pred2)) # output accuracy

support	f1-score	recall	precision	
30	0.74	0.70	0.78	0
31	0.77	0.81	0.74	1
61	0.75			accuracy
61	0.75	0.75	0.76	macro avg
61	0.75	0.75	0.76	weighted avg

Model 3: SVM (Support Vector Machine)



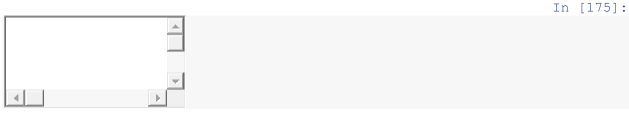
from sklearn.metrics import classification_report
from sklearn.svm import SVC

model3 = SVC(random_state=1) # get instance of model
model3.fit(x_train, y_train) # Train/Fit model

y_pred3 = model3.predict(x_test) # get y predictions
print(classification_report(y_test, y_pred3)) # output accuracy

	precision	recall	f1-score	support
0	0.80	0.67	0.73	30
1	0.72	0.84	0.78	31
accuracy			0.75	61
macro avg	0.76	0.75	0.75	61
weighted avg	0.76	0.75	0.75	61

Model 4: Naives Bayes Classifier



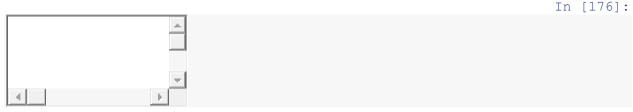
from sklearn.metrics **import** classification_report **from** sklearn.naive_bayes **import** GaussianNB

model4 = GaussianNB() # get instance of model
model4.fit(x_train, y_train) # Train/Fit model

y_pred4 = model4.predict(x_test) # get y predictions
print(classification_report(y_test, y_pred4)) # output accuracy

	precision	recall	f1-score	support
0	0.79	0.73	0.76	30
1	0.76	0.81	0.78	31
accuracy			0.77	61
macro avg	0.77	0.77	0.77	61
weighted avg	0.77	0.77	0.77	61

Model 5: Decision Trees



from sklearn.metrics import classification_report
from sklearn.tree import DecisionTreeClassifier

model5 = DecisionTreeClassifier(random_state=1) # get instance of model
model5.fit(x_train, y_train) # Train/Fit model

y_pred5 = model5.predict(x_test) # get y predictions
print(classification_report(y_test, y_pred5)) # output accuracy

	precision	recall	f1-score	support
0	0.68	0.70	0.69	30
1	0.70	0.68	0.69	31
accuracy			0.69	61
macro avg	0.69	0.69	0.69	61
weighted avg	0.69	0.69	0.69	61

Model 6: Random Forest



from sklearn.metrics import classification_report
from sklearn.ensemble import RandomForestClassifier

model6 = RandomForestClassifier(random_state=1)# get instance of model
model6.fit(x_train, y_train) # Train/Fit model

y_pred6 = model6.predict(x_test) # get y predictions
print(classification_report(y_test, y_pred6)) # output accuracy

	precision	recall	f1-score	support
0	0.88	0.70	0.78	30
1	0.76	0.90	0.82	31
accuracy			0.80	61
macro avg	0.82	0.80	0.80	61
weighted avg	0.81	0.80	0.80	61

Model 7: XGBoost

In [178]:

from xgboost import XGBClassifier

model7 = XGBClassifier(random_state=1)
model7.fit(x_train, y_train)
y_pred7 = model7.predict(x_test)
print(classification_report(y_test, y_pred7))

	preci	sion	recall	f1-sco	re suppo	support	
	0	0.69	0.67	0.	68	30	
	1	0.69	0.71	0.	70	31	
accurac	ЗУ			0.	69	61	
macro av	rg	0.69	0.69	0.	69	61	
weighted av	·g	0.69	0.69	0.	69	61	

From comparing the 7 models, we can conclude that Model 6: Random Forest yields the highest accuracy. With an accuracy of 80%.

We have precision, recall, f1-score and support:

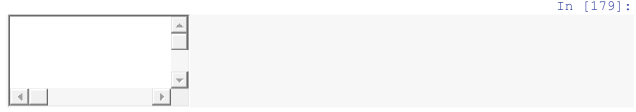
Precision: be "how many are correctly classified among that class"

Recall: "how many of this class you find over the whole number of element of this class"

F1-score: harmonic mean of precision and recall values. F1 score reaches its best value at 1 and worst value at 0. F1 Score = 2 x ((precision x recall) / (precision + recall))

Support: # of samples of the true response that lie in that class.

Making the Confusion Matrix



from sklearn.metrics import confusion_matrix, accuracy_score

cm = confusion_matrix(y_test, y_pred6)

print(cm)

accuracy_score(y_test, y_pred6)

```
[[21 9]
[ 3 28]]
```

Out[179]:

0.8032786885245902

21 is the amount of True Positives in our data, while 28 is the amount of True Negatives.

9 & 3 are the number of errors.

There are 9 type 1 error (False Positives)- You predicted positive and it's false.

There are 3 type 2 error (False Negatives)- You predicted negative and it's false.

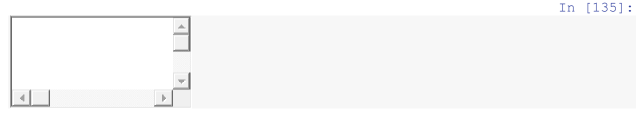
Hence if we calculate the accuracy its # Correct Predicted/ # Total. In other words, where TP, FN, FP and TN represent the number of true positives, false negatives, false positives and true negatives.

(TP + TN)/(TP + TN + FP + FN). (21+28)/(21+28+9+3) = 0.80 = 80% accuracy Note: A good rule of thumb is that any accuracy above 70% is considered good, but be careful because if your accuracy is extremly high, it may be too good to be true (an example of Overfitting). Thus, 80% is the ideal accuracy!

Feature Importance

Feature Importance provides a score that indicates how helpful each feature was in our model.

The higher the Feature Score, the more that feature is used to make key decisions & thus the more important it is.



get importance

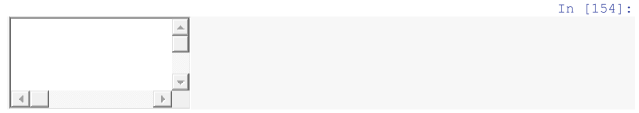
importance = model6.feature_importances_

summarize feature importance

for i,v in enumerate(importance):

```
print('Feature: %0d, Score: %.5f' % (i,v))
Feature: 0, Score: 0.07814
```

```
Feature: 1, Score: 0.04206
Feature: 2, Score: 0.16580
Feature: 3, Score: 0.07477
Feature: 4, Score: 0.07587
Feature: 5, Score: 0.00828
Feature: 6, Score: 0.02014
Feature: 7, Score: 0.12772
Feature: 8, Score: 0.06950
Feature: 9, Score: 0.09957
Feature: 10, Score: 0.04677
Feature: 11, Score: 0.11667
Feature: 12, Score: 0.07473
```



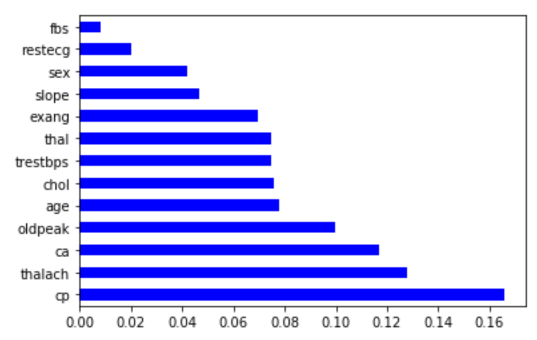
index= data.columns[:-1]

 $importance = pd. Series (model 6. feature_importances_, index=index)$

importance.nlargest(13).plot(kind='barh', colormap='winter')

Out[154]:

<matplotlib.axes._subplots.AxesSubplot at 0x7f9ca1545b90>



From the Feature Importance graph above, we can conclude that the top 4 significant features were chest pain type (cp), maximum heart rate achieved (thalach), number of major vessels (ca), and ST depression induced by exercise relative to rest (oldpeak).

Predictions

Scenario: A patient develops cardiac symptoms & you input his vitals into the Machine Learning Algorithm.

He is a 20 year old male, with a chest pain value of 2 (atypical angina), with resting blood pressure of 110.

In addition he has a serum cholestoral of 230 mg/dl.

He is fasting blood sugar > 120 mg/dl.

He has a resting electrocardiographic result of 1.

The patients maximum heart rate achieved is 140.

Also, he was exercise induced angina.

His ST depression induced by exercise relative to rest value was 2.2.

The slope of the peak exercise ST segment is flat.

He has no major vessels colored by fluoroscopy, and in addition his maximum heart rate achieved is a reversable defect.

Based on this information, can you classify this patient with Heart Disease?



print(model6.predict(sc.transform([[20,1,2,110,230,1,1,140,1,2.2,2,0,2]])))

Γ1

Yes! Our machine learning algorithm has classified this patient with Heart Disease. Now we can properly diagnose him, & get him the help he needs to recover. By diagnosing him early, we may prevent worse symtoms from arising later.

Predicting the Test set results:

First value represents our predicted value, Second value represents our actual value.

If the values match, then we predicted correctly. We can see that our results are very accurate!

In [185]: 4 y_pred = model6.predict(x_test) print(np.concatenate((y_pred.reshape(len(y_pred),1), y_test.reshape(len(y_test),1)),1)) [[0 0]] [1 1] [0 0] [0 0] [0 0] [0 0] [0 0] [1 1] [0 0] [1 1] [1 1] [0 0] [1 0] [0 0] [0 0] [1 0] [1 1] [0 0] [1 1] [1 0] [1 1] [0 0] [1 1]

[1 1] [1 1] [1 1] [0 0] [1 1] [1 1] [1 1] [1 1] [1 1] [1 1] [1 1] [0 0] [1 1] [0 1] [0 0] [1 0] [0 1] [1 1] [0 0] [0 1] [0 0] [1 0] [1 0] [0 0] [1 1] [1 0] [1 1] [1 1] [1 0] [0 0] [1 1] [1 1] [1 1] [1 1] [0 0] [1 0]

Conclusions

[0 0] [1 1]]

- 1. Our Random Forest algorithm yields the highest accuracy, 80%. Any accuracy above 70% is considered good, but be careful because if your accuracy is extremly high, it may be too good to be true (an example of Overfitting). Thus, 80% is the ideal accuracy!
- 2. Out of the 13 features we examined, the top 4 significant features that helped us classify between a positive & negative Diagnosis were chest pain type (cp), maximum heart rate

- achieved (thalach), number of major vessels (ca), and ST depression induced by exercise relative to rest (oldpeak).
- 3. Our machine learning algorithm can now classify patients with Heart Disease. Now we can properly diagnose patients, & get them the help they needs to recover. By diagnosing detecting these features early, we may prevent worse symtoms from arising later.

