### Lab 3.6 - Student Notebook

## Importing the data and training the model

By running the following cells, the data will be imported and ready for use.

**Note:** The following cells represent the key steps in the previous labs.

```
In [1]:
       bucket='c169682a4380823l11215443t1w220560012833-labbucket-ujtpzd5oueum'
In [2]: import warnings, requests, zipfile, io
        warnings.simplefilter('ignore')
        import pandas as pd
        from scipy.io import arff
        import os
        import boto3
        import sagemaker
        import numpy as np
        from sklearn.metrics import roc_curve, auc
        import matplotlib.pyplot as plt
        from sagemaker.image_uris import retrieve
        from sklearn.model_selection import train_test_split
        sagemaker.config INFO - Not applying SDK defaults from location: /etc/xdg/sagem
        aker/config.yaml
        sagemaker.config INFO - Not applying SDK defaults from location: /home/ec2-use
        r/.config/sagemaker/config.yaml
        Matplotlib is building the font cache; this may take a moment.
In [3]: f_zip = 'http://archive.ics.uci.edu/ml/machine-learning-databases/00212/vertebra
        r = requests.get(f_zip, stream=True)
        Vertebral zip = zipfile.ZipFile(io.BytesIO(r.content))
        Vertebral zip.extractall()
        data = arff.loadarff('column 2C weka.arff')
        df = pd.DataFrame(data[0])
        class mapper = {b'Abnormal':1,b'Normal':0}
        df['class']=df['class'].replace(class mapper)
        cols = df.columns.tolist()
        cols = cols[-1:] + cols[:-1]
        df = df[cols]
        train, test_and_validate = train_test_split(df, test_size=0.2, random_state=42,
        test, validate = train test split(test and validate, test size=0.5, random state
        prefix='lab3'
        train file='vertebral train.csv'
        test file='vertebral test.csv'
```

```
validate_file='vertebral_validate.csv'
s3_resource = boto3.Session().resource('s3')
def upload_s3_csv(filename, folder, dataframe):
    csv_buffer = io.StringIO()
    dataframe.to csv(csv buffer, header=False, index=False )
    s3_resource.Bucket(bucket).Object(os.path.join(prefix, folder, filename)).pu
upload_s3_csv(train_file, 'train', train)
upload_s3_csv(test_file, 'test', test)
upload_s3_csv(validate_file, 'validate', validate)
container = retrieve('xgboost',boto3.Session().region_name,'1.0-1')
hyperparams={"num_round":"42",
             "eval_metric": "auc",
             "objective": "binary:logistic"}
s3 output location="s3://{}/{}/output/".format(bucket,prefix)
xgb_model=sagemaker.estimator.Estimator(container,
                                       sagemaker.get_execution_role(),
                                       instance_count=1,
                                       instance_type='ml.m4.xlarge',
                                       output_path=s3_output_location,
                                        hyperparameters=hyperparams,
                                        sagemaker_session=sagemaker.Session())
train_channel = sagemaker.inputs.TrainingInput(
    "s3://{}/{}/train/".format(bucket,prefix,train_file),
    content type='text/csv')
validate_channel = sagemaker.inputs.TrainingInput(
    "s3://{}/validate/".format(bucket,prefix,validate_file),
    content_type='text/csv')
data_channels = {'train': train_channel, 'validation': validate_channel}
xgb_model.fit(inputs=data_channels, logs=False)
batch_X = test.iloc[:,1:];
batch X file='batch-in.csv'
upload_s3_csv(batch_X_file, 'batch-in', batch_X)
batch_output = "s3://{}/batch-out/".format(bucket,prefix)
batch_input = "s3://{}/{}/batch-in/{}".format(bucket,prefix,batch_X_file)
xgb transformer = xgb model.transformer(instance count=1,
                                       instance_type='ml.m4.xlarge',
                                       strategy='MultiRecord',
                                       assemble_with='Line',
                                       output path=batch output)
xgb transformer.transform(data=batch input,
                         data_type='S3Prefix',
                         content_type='text/csv',
                         split_type='Line')
xgb_transformer.wait()
s3 = boto3.client('s3')
```

```
obj = s3.get_object(Bucket=bucket, Key="{}/batch-out/{}".format(prefix,'batch-in
target_predicted = pd.read_csv(io.BytesIO(obj['Body'].read()),names=['class'])

INFO:sagemaker:Creating training-job with name: sagemaker-xgboost-2025-08-14-12
-16-52-123

2025-08-14 12:16:57 Starting - Starting the training job..

2025-08-14 12:17:12 Starting - Preparing the instances for training....

2025-08-14 12:17:36 Downloading - Downloading input data.....

2025-08-14 12:18:06 Downloading - Downloading the training image.......

2025-08-14 12:19:02 Training - Training image download completed. Training in p
rogress.....

2025-08-14 12:19:28 Uploading - Uploading generated training model..

2025-08-14 12:19:41 Completed - Training job completed

INFO:sagemaker:Creating model with name: sagemaker-xgboost-2025-08-14-12-19-44-
389

INFO:sagemaker:Creating transform job with name: sagemaker-xgboost-2025-08-14-12-19-44-
389

INFO:sagemaker:Creating transform job with name: sagemaker-xgboost-2025-08-14-12-19-44-
389

INFO:sagemaker:Creating transform job with name: sagemaker-xgboost-2025-08-14-12-19-44-
389
```

# Step 1: Exploring the results

The output from the model will be a probablility. You must first convert that probability into one of the two classes, either 0 or 1. To do this, you can create a function to perform the conversion. Note the use of the threshold in the function.

```
In [4]: def binary_convert(x):
              threshold = 0.3
              if x > threshold:
                  return 1
              else:
                  return 0
         target predicted binary = target predicted['class'].apply(binary convert)
         print(target_predicted_binary.head(5))
         test.head(5)
         0
               1
         1
               1
         3
               1
         Name: class, dtype: int64
Out[4]:
                    pelvic_incidence pelvic_tilt lumbar_lordosis_angle sacral_slope pelvic_radius degre
         136
                          88.024499 39.844669
                                                          81.774473
                                                                       48.179830
                                                                                   116.601538
         230
                          65.611802 23.137919
                                                           62.582179
                                                                       42.473883
                                                                                   124.128001
                          52.204693 17.212673
         134
                                                          78.094969
                                                                       34.992020
                                                                                   136.972517
                                                           32.168463
                                                                                    99.712453
         130
                          50.066786
                                    9.120340
                                                                       40.946446
          47
                          41.352504 16.577364
                                                                                   113.266675
                                                           30.706191
                                                                       24.775141
```

Based on these results, you can see that the initial model might not be that good. It's difficult to tell by comparing a few values.

Next, you will generate some metrics to see how well the model performs.

## Step 2: Creating a confusion matrix

A *confusion matrix* is one of the key ways of measuring a classification model's performance. It's a table that maps out the correct and incorrect predictions. After you calculate a confusion matrix for your model, you can generate several other statistics. However, you will start by only creating the confusion matrix.

To create a confusion matrix, you need both the target values from your test data and the predicted value.

Get the targets from the test DataFrame.

```
In [5]: test_labels = test.iloc[:,0]
    test_labels.head()

Out[5]: 136     1
    230     0
    134     1
    130     1
    47     1
    Name: class, dtype: int64
```

Now, you can use the *scikit-learn* library, which contains a function to create a confusion matrix.

```
In [6]: from sklearn.metrics import confusion_matrix

matrix = confusion_matrix(test_labels, target_predicted_binary)

df_confusion = pd.DataFrame(matrix, index=['Nnormal','Abnormal'],columns=['Normal'])

df_confusion
```

```
        Normal
        Abnormal

        Nnormal
        7
        3

        Abnormal
        2
        19
```

You results will vary, but you should have results that are similiar to this example:

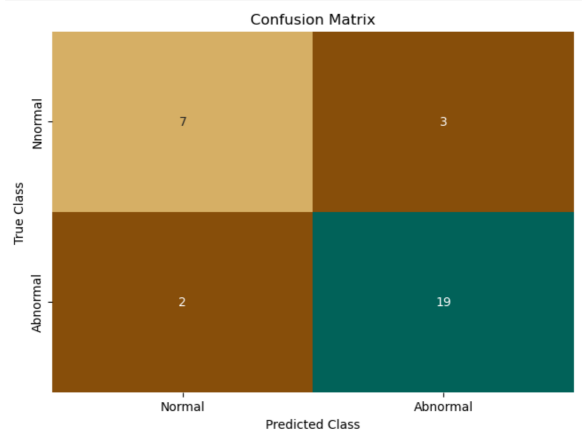
	Normal	Abnormal
Normal	7	3
Abnormal	3	18

The previous table shows that the model correctly predicted 7 Normal and 18 Abnormal values. However, it incorrectly predicted 3 Normal and 3 Abnormal values.

By using the *seaborn* and *matplotlib* Python libraries, you can plot these values in a chart to make them easier to read.

```
import seaborn as sns
import matplotlib.pyplot as plt

colormap = sns.color_palette("BrBG", 10)
sns.heatmap(df_confusion, annot=True, cbar=None, cmap=colormap)
plt.title("Confusion Matrix")
plt.tight_layout()
plt.ylabel("True Class")
plt.xlabel("Predicted Class")
plt.show()
```



**Tip:** If the chart doesn't display the first time, try running the cell again.

If these results are good enough for your application, then the model might be good enough. However, because there are consequences from incorrectly predicting *Normal* values -- that is, no abnormality was found when there actually was one -- the focus should be on reducing this result.

## Step 3: Calculating performance statistics

If you want to compare this model to the next model that you create, you need some metrics that you can record. For a binary classification problem, the confusion matrix data can be used to calculate various metrics.

To start, extract the values from the confusion matrix cells into variables.

```
In [8]: from sklearn.metrics import roc_auc_score, roc_curve, auc

TN, FP, FN, TP = confusion_matrix(test_labels, target_predicted_binary).ravel()

print(f"True Negative (TN) : {TN}")
print(f"False Positive (FP): {FP}")
print(f"False Negative (FN): {FN}")
print(f"True Positive (TP) : {TP}")

True Negative (TN) : 7
False Positive (FP): 3
False Negative (FN): 2
True Positive (TP) : 19
```

You can now calculate some statistics.

#### Sensitivity

Sensitivity is also known as hit rate, recall, or true positive rate (TPR). It measures the proportion of the actual positives that are correctly identified.

In this example, the sensitivity is the probablity of detecting an abnormality for patients with an abnormality.

```
In [9]: # Sensitivity, hit rate, recall, or true positive rate
Sensitivity = float(TP)/(TP+FN)*100
print(f"Sensitivity or TPR: {Sensitivity}%")
print(f"There is a {Sensitivity}% chance of detecting patients with an abnormali
```

Sensitivity or TPR: 90.47619047619048% There is a 90.47619047619048% chance of detecting patients with an abnormality have an abnormality

Question: Is the sensitivity good enough for this scenario?

#### **Specificity**

The next statistic is *specificity*, which is also known as the *true negative*. It measures the proportion of the actual negatives that are correctly identified.

In this example, the specificity is the probablity of detecting normal, for patients who are normal.

```
In [10]: # Specificity or true negative rate
Specificity = float(TN)/(TN+FP)*100
print(f"Specificity or TNR: {Specificity}%")
print(f"There is a {Specificity}% chance of detecting normal patients are normal
```

```
Specificity or TNR: 70.0%
There is a 70.0% chance of detecting normal patients are normal.
```

**Question:** Is this specificity too low, exactly right, or too high? What value would you want to see here, given the scenario?

#### Positive and negative predictive values

The precision, or positive predictive value, is the proportion of positive results.

In this example, the positive predictive value is the probability that subjects with a positive screening test truly have an abnormality.

The *negative predictive value* is the proportion of negative results.

In this example, the negative predictive value is the probability that subjects with a negative screening test truly have an abnormality.

```
In [12]: # Negative predictive value
    NPV = float(TN)/(TN+FN)*100
    print(f"Negative Predictive Value: {NPV}%")
    print(f"You don't have an abnormality, but there is a {NPV}% chance that is inco
    Negative Predictive Value: 77.777777777779%
    You don't have an abnormality, but there is a 77.77777777779% chance that is incorrect
```

Think about the impact of these values. If you were a patient, how worried should you be if the test for an abnormality was positive? On the opposite side, how reassured should you be if you tested negative?

#### False positive rate

The false positive rate (FPR) is the probability that a false alarm will be raised, or that a positive result will be given when the true value is negative.

```
In [13]: # Fall out or false positive rate
FPR = float(FP)/(FP+TN)*100
print( f"False Positive Rate: {FPR}%")
print( f"There is a {FPR}% chance that this positive result is incorrect.")
```

False Positive Rate: 30.0% There is a 30.0% chance that this positive result is incorrect.

#### False negative rate

The false negative rate -- or miss rate -- is the probability that a true positive will be missed by the test.

```
In [14]: # False negative rate
FNR = float(FN)/(TP+FN)*100
print(f"False Negative Rate: {FNR}%")
print(f"There is a {FNR}% chance that this negative result is incorrect.")
```

False Negative Rate: 9.523809523809524%
There is a 9.523809523809524% chance that this negative result is incorrect.

#### False discovery rate

In this example, the false discovery rate is the probability of predicting an abnormality when the patient doesn't have one.

```
In [15]: # False discovery rate
FDR = float(FP)/(TP+FP)*100
print(f"False Discovery Rate: {FDR}%" )
print(f"You have an abnormality, but there is a {FDR}% chance this is incorrect.
```

#### Overall accuracy

How accuracte is your model?

```
In [16]: # Overall accuracy
ACC = float(TP+TN)/(TP+FP+FN+TN)*100
print(f"Accuracy: {ACC}%")
```

Accuracy: 83.87096774193549%

In summary, you calculated the following metrics from your model:

```
In [17]: print(f"Sensitivity or TPR: {Sensitivity}%")
    print(f"Specificity or TNR: {Specificity}%")
    print(f"Precision: {Precision}%")
    print(f"Negative Predictive Value: {NPV}%")
    print( f"False Positive Rate: {FPR}%")
    print(f"False Negative Rate: {FNR}%")
    print(f"False Discovery Rate: {FDR}%")
    print(f"Accuracy: {ACC}%")
```

Sensitivity or TPR: 90.47619047619048%

Specificity or TNR: 70.0% Precision: 86.36363636363636%

Negative Predictive Value: 77.7777777779%

False Positive Rate: 30.0%

Accuracy: 83.87096774193549%

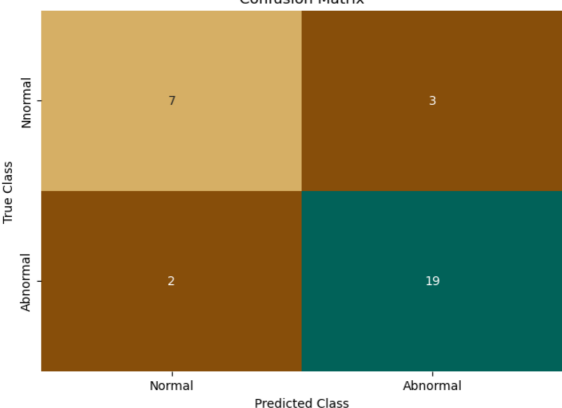
**Challenge task:** Record the previous values, then go back to step 1 and change the value used for the threshold. Values you should try are .25 and .75.

Did those threshold values make a difference?

thershold=0.25

```
In [25]:
          def binary convert(x):
              threshold = 0.25
              if x > threshold:
                   return 1
              else:
                   return 0
          target_predicted_binary = target_predicted['class'].apply(binary_convert)
          print(target_predicted_binary.head(5))
          test.head(5)
          0
               1
          1
               1
          2
               1
          3
               1
               1
          Name: class, dtype: int64
Out[25]:
               class pelvic_incidence pelvic_tilt lumbar_lordosis_angle sacral_slope pelvic_radius degre
          136
                  1
                          88.024499 39.844669
                                                         81.774473
                                                                     48.179830
                                                                                 116.601538
          230
                  0
                          65.611802 23.137919
                                                         62.582179
                                                                     42.473883
                                                                                 124.128001
          134
                  1
                          52.204693 17.212673
                                                         78.094969
                                                                     34.992020
                                                                                 136.972517
                                                                                  99.712453
          130
                           50.066786
                                    9.120340
                                                         32.168463
                                                                     40.946446
           47
                  1
                          41.352504 16.577364
                                                         30.706191
                                                                     24.775141
                                                                                 113.266675
          test_labels = test.iloc[:,0]
In [26]:
          test_labels.head()
Out[26]: 136
                 1
          230
          134
                  1
          130
                  1
          47
          Name: class, dtype: int64
In [27]:
          import seaborn as sns
          import matplotlib.pyplot as plt
          colormap = sns.color_palette("BrBG", 10)
          sns.heatmap(df_confusion, annot=True, cbar=None, cmap=colormap)
          plt.title("Confusion Matrix")
          plt.tight_layout()
          plt.ylabel("True Class")
          plt.xlabel("Predicted Class")
          plt.show()
```

#### Confusion Matrix



```
In [28]: from sklearn.metrics import roc_auc_score, roc_curve, auc
         TN, FP, FN, TP = confusion_matrix(test_labels, target_predicted_binary).ravel()
         print(f"True Negative (TN) : {TN}")
         print(f"False Positive (FP): {FP}")
         print(f"False Negative (FN): {FN}")
         print(f"True Positive (TP) : {TP}")
         True Negative (TN): 6
         False Positive (FP): 4
         False Negative (FN): 2
         True Positive (TP): 19
In [29]: # Sensitivity, hit rate, recall, or true positive rate
         Sensitivity = float(TP)/(TP+FN)*100
         print(f"Sensitivity or TPR: {Sensitivity}%")
         print(f"There is a {Sensitivity}% chance of detecting patients with an abnormali
         Sensitivity or TPR: 90.47619047619048%
         There is a 90.47619047619048% chance of detecting patients with an abnormality
         have an abnormality
In [30]: # Specificity or true negative rate
         Specificity = float(TN)/(TN+FP)*100
         print(f"Specificity or TNR: {Specificity}%")
         print(f"There is a {Specificity}% chance of detecting normal patients are normal
         Specificity or TNR: 60.0%
         There is a 60.0% chance of detecting normal patients are normal.
In [31]: # Precision or positive predictive value
         Precision = float(TP)/(TP+FP)*100
```

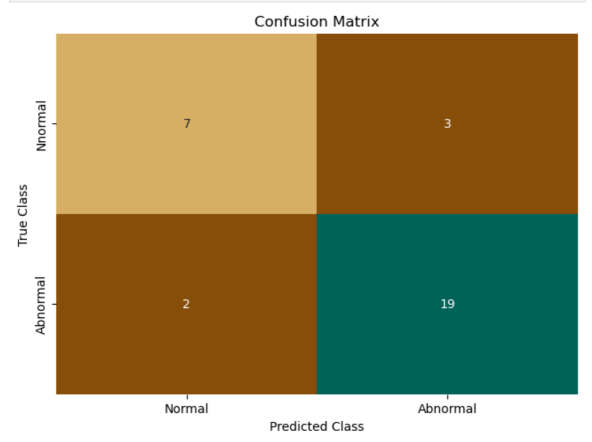
```
print(f"Precision: {Precision}%")
         print(f"You have an abnormality, and the probablity that is correct is {Precisio
         Precision: 82.6086956521739%
         You have an abnormality, and the probablity that is correct is 82.608695652173
In [32]: # Negative predictive value
         NPV = float(TN)/(TN+FN)*100
         print(f"Negative Predictive Value: {NPV}%")
         print(f"You don't have an abnormality, but there is a {NPV}% chance that is inco
         Negative Predictive Value: 75.0%
         You don't have an abnormality, but there is a 75.0% chance that is incorrect
        # Fall out or false positive rate
In [33]:
         FPR = float(FP)/(FP+TN)*100
         print( f"False Positive Rate: {FPR}%")
         print( f"There is a {FPR}% chance that this positive result is incorrect.")
         False Positive Rate: 40.0%
         There is a 40.0% chance that this positive result is incorrect.
In [34]: # False negative rate
         FNR = float(FN)/(TP+FN)*100
         print(f"False Negative Rate: {FNR}%")
         print(f"There is a {FNR}% chance that this negative result is incorrect.")
         False Negative Rate: 9.523809523809524%
         There is a 9.523809523809524% chance that this negative result is incorrect.
In [35]: # False discovery rate
         FDR = float(FP)/(TP+FP)*100
         print(f"False Discovery Rate: {FDR}%" )
         print(f"You have an abnormality, but there is a {FDR}% chance this is incorrect.
         False Discovery Rate: 17.391304347826086%
         You have an abnormality, but there is a 17.391304347826086% chance this is inco
         rrect.
In [36]: # Overall accuracy
         ACC = float(TP+TN)/(TP+FP+FN+TN)*100
         print(f"Accuracy: {ACC}%")
         Accuracy: 80.64516129032258%
In [37]:
        print(f"Sensitivity or TPR: {Sensitivity}%")
         print(f"Specificity or TNR: {Specificity}%")
         print(f"Precision: {Precision}%")
         print(f"Negative Predictive Value: {NPV}%")
         print( f"False Positive Rate: {FPR}%")
         print(f"False Negative Rate: {FNR}%")
         print(f"False Discovery Rate: {FDR}%" )
         print(f"Accuracy: {ACC}%")
```

Sensitivity or TPR: 90.47619047619048%

Specificity or TNR: 60.0% Precision: 82.6086956521739% Negative Predictive Value: 75.0%

```
False Positive Rate: 40.0%
          False Negative Rate: 9.523809523809524%
          False Discovery Rate: 17.391304347826086%
          Accuracy: 80.64516129032258%
          Thershold=0.75
In [38]:
         def binary_convert(x):
              threshold = 0.75
              if x > threshold:
                  return 1
              else:
                  return 0
          target_predicted_binary = target_predicted['class'].apply(binary_convert)
          print(target_predicted_binary.head(5))
          test.head(5)
          0
               1
          1
               1
               1
          3
               1
          4
               1
          Name: class, dtype: int64
Out[38]:
               class pelvic_incidence pelvic_tilt lumbar_lordosis_angle sacral_slope pelvic_radius degre
          136
                          88.024499 39.844669
                                                                     48.179830
                  1
                                                         81.774473
                                                                                 116.601538
                          65.611802 23.137919
                                                         62.582179
                                                                     42.473883
                                                                                 124.128001
          230
          134
                  1
                          52.204693 17.212673
                                                         78.094969
                                                                     34.992020
                                                                                 136.972517
          130
                          50.066786
                                    9.120340
                                                         32.168463
                                                                     40.946446
                                                                                 99.712453
           47
                  1
                          41.352504 16.577364
                                                         30.706191
                                                                     24.775141
                                                                                 113.266675
In [39]: test_labels = test.iloc[:,0]
          test labels.head()
Out[39]: 136
                 1
          230
                 0
          134
                 1
          130
                 1
          47
          Name: class, dtype: int64
In [40]: import seaborn as sns
          import matplotlib.pyplot as plt
          colormap = sns.color_palette("BrBG", 10)
          sns.heatmap(df confusion, annot=True, cbar=None, cmap=colormap)
          plt.title("Confusion Matrix")
          plt.tight_layout()
          plt.ylabel("True Class")
```

```
plt.xlabel("Predicted Class")
plt.show()
```



```
In [41]: from sklearn.metrics import roc_auc_score, roc_curve, auc
         TN, FP, FN, TP = confusion_matrix(test_labels, target_predicted_binary).ravel()
         print(f"True Negative (TN) : {TN}")
         print(f"False Positive (FP): {FP}")
         print(f"False Negative (FN): {FN}")
         print(f"True Positive (TP) : {TP}")
         True Negative (TN): 7
         False Positive (FP): 3
         False Negative (FN): 5
         True Positive (TP): 16
In [42]: # Sensitivity, hit rate, recall, or true positive rate
         Sensitivity = float(TP)/(TP+FN)*100
         print(f"Sensitivity or TPR: {Sensitivity}%")
         print(f"There is a {Sensitivity}% chance of detecting patients with an abnormali
         Sensitivity or TPR: 76.19047619047619%
         There is a 76.19047619047619% chance of detecting patients with an abnormality
         have an abnormality
In [43]: # Specificity or true negative rate
         Specificity = float(TN)/(TN+FP)*100
         print(f"Specificity or TNR: {Specificity}%")
         print(f"There is a {Specificity}% chance of detecting normal patients are normal
         Specificity or TNR: 70.0%
         There is a 70.0% chance of detecting normal patients are normal.
```

```
In [44]: # Precision or positive predictive value
         Precision = float(TP)/(TP+FP)*100
         print(f"Precision: {Precision}%")
         print(f"You have an abnormality, and the probablity that is correct is {Precisio
         Precision: 84.21052631578947%
         You have an abnormality, and the probablity that is correct is 84.2105263157894
In [45]: # Negative predictive value
         NPV = float(TN)/(TN+FN)*100
         print(f"Negative Predictive Value: {NPV}%")
         print(f"You don't have an abnormality, but there is a {NPV}% chance that is inco
         Negative Predictive Value: 58.333333333333336%
         You don't have an abnormality, but there is a 58.33333333333336% chance that i
         s incorrect
In [ ]: # Fall out or false positive rate
         FPR = float(FP)/(FP+TN)*100
         print( f"False Positive Rate: {FPR}%")
         print( f"There is a {FPR}% chance that this positive result is incorrect.")
In [46]: #False negative rate
         FNR = float(FN)/(TP+FN)*100
         print(f"False Negative Rate: {FNR}%")
         print(f"There is a {FNR}% chance that this negative result is incorrect.")
         False Negative Rate: 23.809523809523807%
         There is a 23.809523809523807% chance that this negative result is incorrect.
In [47]: # False discovery rate
         FDR = float(FP)/(TP+FP)*100
         print(f"False Discovery Rate: {FDR}%" )
         print(f"You have an abnormality, but there is a {FDR}% chance this is incorrect.
         False Discovery Rate: 15.789473684210526%
         You have an abnormality, but there is a 15.789473684210526% chance this is inco
         rrect.
In [48]: # Overall accuracy
         ACC = float(TP+TN)/(TP+FP+FN+TN)*100
         print(f"Accuracy: {ACC}%")
         Accuracy: 74.19354838709677%
        print(f"Sensitivity or TPR: {Sensitivity}%")
In [49]:
         print(f"Specificity or TNR: {Specificity}%")
         print(f"Precision: {Precision}%")
         print(f"Negative Predictive Value: {NPV}%")
         print( f"False Positive Rate: {FPR}%")
         print(f"False Negative Rate: {FNR}%")
         print(f"False Discovery Rate: {FDR}%" )
         print(f"Accuracy: {ACC}%")
```

Sensitivity or TPR: 76.19047619047619%

Specificity or TNR: 70.0% Precision: 84.21052631578947%

Negative Predictive Value: 58.33333333333336%

False Positive Rate: 40.0%

False Negative Rate: 23.809523809523807% False Discovery Rate: 15.789473684210526%

Accuracy: 74.19354838709677%

## Step 4: Calculating the AUC-ROC Curve

The scikit-learn library has functions that can help you compute the *area under the* receiver operating characteristic curve (AUC-ROC).

- The ROC is a probability curve.
- The AUC tells you how well the model can distinguish between classes.

The AUC can be calculated. As you will see in the next lab, it can be used to measure the performance of the model.

In this example, the higher the AUC, the better the model is at distinguishing between abnormal and normal patients.

Depending on the value you set for the threshold, the AUC can change. You can plot the AUC by using the probability instead of your converted class.

```
In [18]: test_labels = test.iloc[:,0];
    print("Validation AUC", roc_auc_score(test_labels, target_predicted) )
        Validation AUC 0.8904761904761904
In []:
```

Typically, the ROC curve is plotted with the TPR against the FPR, where the TPR is on the y-axis and the FPR is on the x-axis.

scikit-learn has the **roc\_curve** function to help generate those values to plot.

```
In [19]: fpr, tpr, thresholds = roc_curve(test_labels, target_predicted)

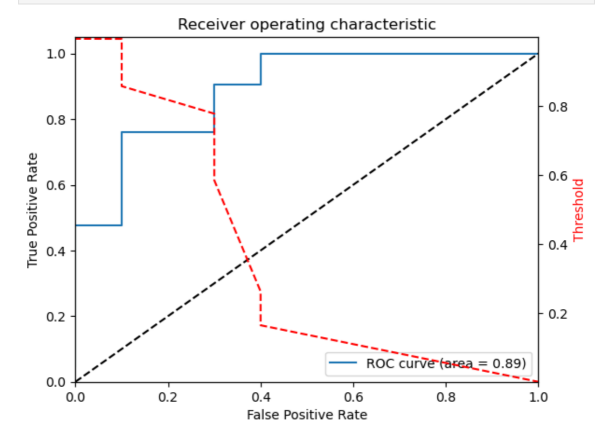
finite_indices = np.isfinite(thresholds)
fpr_finite = fpr[finite_indices]
tpr_finite = tpr[finite_indices]
thresholds_finite = thresholds[finite_indices]

plt.figure()
plt.plot(fpr_finite, tpr_finite, label='ROC curve (area = %0.2f)' % auc(fpr_finite)
plt.plot([0, 1], [0, 1], 'k--') # Dashed diagonal
plt.xlim([0.0, 1.0])
plt.ylim([0.0, 1.05])
plt.xlabel('False Positive Rate')
plt.ylabel('True Positive Rate')
plt.title('Receiver operating characteristic')
plt.legend(loc="lower right")
```

```
roc_auc = auc(fpr, tpr)

if thresholds_finite.size > 0:
    ax2 = plt.gca().twinx()
    ax2.plot(fpr_finite, thresholds_finite, markeredgecolor='r', linestyle='dash ax2.set_ylabel('Threshold', color='r')
    ax2.set_ylim([thresholds_finite[-1], thresholds_finite[0]])
    ax2.set_xlim([fpr_finite[0], fpr_finite[-1]])

plt.show()
```



**Challenge task:** Update the previous code to use *target\_predicted\_binary* instead of *target\_predicted*. How does that change the graph? Which is the most useful?

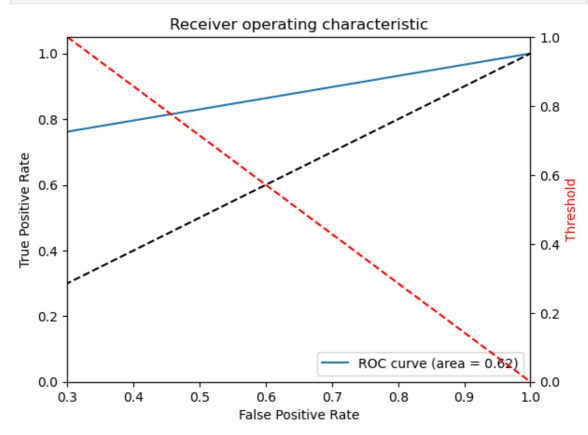
```
In [50]:
         test_labels = test.iloc[:,0];
         print("Validation AUC", roc_auc_score(test_labels, target_predicted_binary) )
         Validation AUC 0.7309523809523808
        fpr, tpr, thresholds = roc_curve(test_labels, target_predicted_binary)
In [51]:
         finite indices = np.isfinite(thresholds)
         fpr finite = fpr[finite indices]
         tpr_finite = tpr[finite_indices]
         thresholds_finite = thresholds[finite_indices]
         plt.figure()
         plt.plot(fpr_finite, tpr_finite, label='ROC curve (area = %0.2f)' % auc(fpr_finite)
         plt.plot([0, 1], [0, 1], 'k--') # Dashed diagonal
         plt.xlim([0.0, 1.0])
         plt.ylim([0.0, 1.05])
         plt.xlabel('False Positive Rate')
         plt.ylabel('True Positive Rate')
         plt.title('Receiver operating characteristic')
```

```
plt.legend(loc="lower right")

roc_auc = auc(fpr, tpr)

if thresholds_finite.size > 0:
    ax2 = plt.gca().twinx()
    ax2.plot(fpr_finite, thresholds_finite, markeredgecolor='r', linestyle='dash ax2.set_ylabel('Threshold', color='r')
    ax2.set_ylim([thresholds_finite[-1], thresholds_finite[0]])
    ax2.set_xlim([fpr_finite[0], fpr_finite[-1]])

plt.show()
```



# Congratulations!

You have completed this lab, and you can now end the lab by following the lab guide instructions.