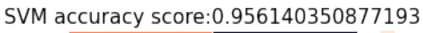
## 10\_Support\_vector\_Machines

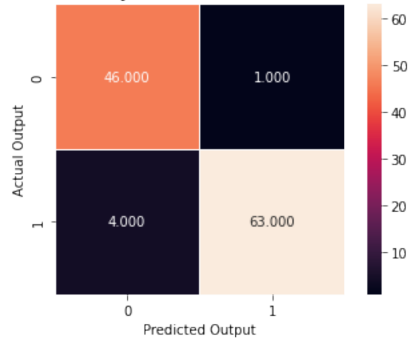
June 18, 2023

```
[]: # importing sklearn dataset library
     from sklearn import datasets
     # Load dataset
     cancer = datasets.load_breast_cancer()
[]: # print the name of 30 features
     print("Features:", cancer.feature_names)
     # print the label type of cancer
     print("labels:",cancer.target_names)
    Features: ['mean radius' 'mean texture' 'mean perimeter' 'mean area'
     'mean smoothness' 'mean compactness' 'mean concavity'
     'mean concave points' 'mean symmetry' 'mean fractal dimension'
     'radius error' 'texture error' 'perimeter error' 'area error'
     'smoothness error' 'compactness error' 'concavity error'
     'concave points error' 'symmetry error' 'fractal dimension error'
     'worst radius' 'worst texture' 'worst perimeter' 'worst area'
     'worst smoothness' 'worst compactness' 'worst concavity'
     'worst concave points' 'worst symmetry' 'worst fractal dimension']
    labels: ['malignant' 'benign']
[]: # print data(feature) shape
     cancer.data.shape
[]: (569, 30)
[]: # PRINT THE cancer data feature(top 5 records)
     print(cancer.data[0:5])
    [[1.799e+01 1.038e+01 1.228e+02 1.001e+03 1.184e-01 2.776e-01 3.001e-01
      1.471e-01 2.419e-01 7.871e-02 1.095e+00 9.053e-01 8.589e+00 1.534e+02
      6.399e-03 4.904e-02 5.373e-02 1.587e-02 3.003e-02 6.193e-03 2.538e+01
      1.733e+01 1.846e+02 2.019e+03 1.622e-01 6.656e-01 7.119e-01 2.654e-01
      4.601e-01 1.189e-01]
     [2.057e+01 1.777e+01 1.329e+02 1.326e+03 8.474e-02 7.864e-02 8.690e-02
      7.017e-02 1.812e-01 5.667e-02 5.435e-01 7.339e-01 3.398e+00 7.408e+01
      5.225e-03 1.308e-02 1.860e-02 1.340e-02 1.389e-02 3.532e-03 2.499e+01
      2.341e+01 1.588e+02 1.956e+03 1.238e-01 1.866e-01 2.416e-01 1.860e-01
      2.750e-01 8.902e-02]
```

```
[1.969e+01 2.125e+01 1.300e+02 1.203e+03 1.096e-01 1.599e-01 1.974e-01
    1.279e-01 2.069e-01 5.999e-02 7.456e-01 7.869e-01 4.585e+00 9.403e+01
    6.150e-03 4.006e-02 3.832e-02 2.058e-02 2.250e-02 4.571e-03 2.357e+01
    2.553e+01 1.525e+02 1.709e+03 1.444e-01 4.245e-01 4.504e-01 2.430e-01
    3.613e-01 8.758e-02]
   [1.142e+01 2.038e+01 7.758e+01 3.861e+02 1.425e-01 2.839e-01 2.414e-01
    1.052e-01 2.597e-01 9.744e-02 4.956e-01 1.156e+00 3.445e+00 2.723e+01
   9.110e-03 7.458e-02 5.661e-02 1.867e-02 5.963e-02 9.208e-03 1.491e+01
   2.650e+01 9.887e+01 5.677e+02 2.098e-01 8.663e-01 6.869e-01 2.575e-01
    6.638e-01 1.730e-01]
   [2.029e+01 1.434e+01 1.351e+02 1.297e+03 1.003e-01 1.328e-01 1.980e-01
    1.043e-01 1.809e-01 5.883e-02 7.572e-01 7.813e-01 5.438e+00 9.444e+01
    1.149e-02 2.461e-02 5.688e-02 1.885e-02 1.756e-02 5.115e-03 2.254e+01
    1.667e+01 1.522e+02 1.575e+03 1.374e-01 2.050e-01 4.000e-01 1.625e-01
    2.364e-01 7.678e-02]]
[]: # print the cancer labels O:malingnant 1: benign
   print(cancer.target)
  [0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;1\;1\;1\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0\;0
   1 0 1 0 0 1 1 1 0 0 1 0 0 0 1 1 1 0 1 1 0 0 1 1 1 0 0 1 1 1 1 0 1 1 0 1 1
   1 1 1 1 1 1 1 0 0 0 0 0 0 1
[]: # import train test split function
   from sklearn.model_selection import train_test_split
   X_train, X_test, y_train, y_test = train_test_split(cancer.data,cancer.
   →target,test size=0.2,random state=0)
[]: # import sum model
   from sklearn import svm
   # Create a SVM Classifier
   clf = svm.SVC(kernel='linear') # linear kernal
   # train the model using training set
   clf.fit(X_train, y_train)
```

```
# predict the response of the test dataset
     y_pred = clf.predict(X_test)
[]: # import sklearn metrics module for accuracy calculation
     from sklearn import metrics
     score = metrics.accuracy_score(y_test, y_pred)
     print("Accuracy:", score)
    Accuracy: 0.956140350877193
[]: # Model Precision: what percentage of positive tuples are labeled as such?
     print("Precision:", metrics.precision_score(y_test,y_pred))
     # Model Recall: what percentage of positive tuples are labeled as such?
     print("Recall:", metrics.recall_score(y_test,y_pred))
    Precision: 0.984375
    Recall: 0.9402985074626866
[]: # confusion matrix
     from sklearn import metrics
     cm = metrics.confusion_matrix(y_test, y_pred)
     print(cm)
    [[46 1]
     [ 4 63]]
[]: import seaborn as sns
     import matplotlib.pyplot as plt
     plt.figure("figsize"== (12,12))
     sns.heatmap(cm, annot=True, fmt=".3f", linewidths=.5, square = True)
     plt.xlabel('Predicted Output')
     plt.ylabel('Actual Output')
     all_sample_title = 'SVM accuracy score:{0}'.format(score)
     plt.title(all_sample_title, size = 15)
[]: Text(0.5, 1.0, 'SVM accuracy score:0.956140350877193')
```





[]: