**SVM (Support Vector Machines)[¶](https://jupyterlab-0-labs-prod-jupyterlab-us-east-1.labs.cognitiveclass.ai/user/akulasamson/lab/tree/labs/coursera/ML0101EN/ML0101EN-Clas-SVM-cancer.ipynb" \l "SVM-(Support-Vector-Machines)" \t "_self)**

Estimated time needed: **15** minutes

**Objectives**

After completing this lab you will be able to:

* Use scikit-learn to Support Vector Machine to classify

In this notebook, you will use SVM (Support Vector Machines) to build and train a model using human cell records, and classify cells to whether the samples are benign or malignant.

SVM works by mapping data to a high-dimensional feature space so that data points can be categorized, even when the data are not otherwise linearly separable. A separator between the categories is found, then the data is transformed in such a way that the separator could be drawn as a hyperplane. Following this, characteristics of new data can be used to predict the group to which a new record should belong.

**Table of contents**

1. [Load the Cancer data](https://jupyterlab-0-labs-prod-jupyterlab-us-east-1.labs.cognitiveclass.ai/user/akulasamson/files/labs/coursera/ML0101EN/https%3A/%23load_dataset?_xsrf=2%7C85721345%7C294cb902c3ae357f3da8b1d9e974b2dc%7C1631229491)
2. [Modeling](https://jupyterlab-0-labs-prod-jupyterlab-us-east-1.labs.cognitiveclass.ai/user/akulasamson/files/labs/coursera/ML0101EN/https%3A/%23modeling?_xsrf=2%7C85721345%7C294cb902c3ae357f3da8b1d9e974b2dc%7C1631229491)
3. [Evaluation](https://jupyterlab-0-labs-prod-jupyterlab-us-east-1.labs.cognitiveclass.ai/user/akulasamson/files/labs/coursera/ML0101EN/https%3A/%23evaluation?_xsrf=2%7C85721345%7C294cb902c3ae357f3da8b1d9e974b2dc%7C1631229491)
4. [Practice](https://jupyterlab-0-labs-prod-jupyterlab-us-east-1.labs.cognitiveclass.ai/user/akulasamson/files/labs/coursera/ML0101EN/https%3A/%23practice?_xsrf=2%7C85721345%7C294cb902c3ae357f3da8b1d9e974b2dc%7C1631229491)

[ ]:



**!**pip install scikit**-**learn**==**0.23.1

[ ]:



**import** pandas **as** pd

**import** pylab **as** pl

**import** numpy **as** np

**import** scipy.optimize **as** opt

**from** sklearn **import** preprocessing

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

**%**matplotlib inline

**import** matplotlib.pyplot **as** plt

**Load the Cancer data**

The example is based on a dataset that is publicly available from the UCI Machine Learning Repository (Asuncion and Newman, 2007)[http://mlearn.ics.uci.edu/MLRepository.html]. The dataset consists of several hundred human cell sample records, each of which contains the values of a set of cell characteristics. The fields in each record are:

| **Field name** | **Description** |
| --- | --- |
| ID | Clump thickness |
| Clump | Clump thickness |
| UnifSize | Uniformity of cell size |
| UnifShape | Uniformity of cell shape |
| MargAdh | Marginal adhesion |
| SingEpiSize | Single epithelial cell size |
| BareNuc | Bare nuclei |
| BlandChrom | Bland chromatin |
| NormNucl | Normal nucleoli |
| Mit | Mitoses |
| Class | Benign or malignant |

For the purposes of this example, we're using a dataset that has a relatively small number of predictors in each record. To download the data, we will use !wget to download it from IBM Object Storage.

**Did you know?** When it comes to Machine Learning, you will likely be working with large datasets. As a business, where can you host your data? IBM is offering a unique opportunity for businesses, with 10 Tb of IBM Cloud Object Storage: [Sign up now for free](http://cocl.us/ML0101EN-IBM-Offer-CC)

[ ]:



*#Click here and press Shift+Enter*

**!**wget **-**O cell\_samples.csv https:**//**cf**-**courses**-**data.s3.us.cloud**-**object**-**storage.appdomain.cloud**/**IBMDeveloperSkillsNetwork**-**ML0101EN**-**SkillsNetwork**/**labs**/**Module**%**203**/**data**/**cell\_samples.csv

**Load Data From CSV File**

[ ]:



cell\_df **=** pd.read\_csv("cell\_samples.csv")

cell\_df.head()

The ID field contains the patient identifiers. The characteristics of the cell samples from each patient are contained in fields Clump to Mit. The values are graded from 1 to 10, with 1 being the closest to benign.

The Class field contains the diagnosis, as confirmed by separate medical procedures, as to whether the samples are benign (value = 2) or malignant (value = 4).

Let's look at the distribution of the classes based on Clump thickness and Uniformity of cell size:

[ ]:



ax **=** cell\_df[cell\_df['Class'] **==** 4][0:50].plot(kind**=**'scatter', x**=**'Clump', y**=**'UnifSize', color**=**'DarkBlue', label**=**'malignant');

cell\_df[cell\_df['Class'] **==** 2][0:50].plot(kind**=**'scatter', x**=**'Clump', y**=**'UnifSize', color**=**'Yellow', label**=**'benign', ax**=**ax);

plt.show()

**Data pre-processing and selection**

Let's first look at columns data types:

[ ]:



cell\_df.dtypes

It looks like the **BareNuc** column includes some values that are not numerical. We can drop those rows:

[ ]:



cell\_df **=** cell\_df[pd.to\_numeric(cell\_df['BareNuc'], errors**=**'coerce').notnull()]

cell\_df['BareNuc'] **=** cell\_df['BareNuc'].astype('int')

cell\_df.dtypes

[ ]:



feature\_df **=** cell\_df[['Clump', 'UnifSize', 'UnifShape', 'MargAdh', 'SingEpiSize', 'BareNuc', 'BlandChrom', 'NormNucl', 'Mit']]

X **=** np.asarray(feature\_df)

X[0:5]

We want the model to predict the value of Class (that is, benign (=2) or malignant (=4)). As this field can have one of only two possible values, we need to change its measurement level to reflect this.

[ ]:



cell\_df['Class'] **=** cell\_df['Class'].astype('int')

y **=** np.asarray(cell\_df['Class'])

y [0:5]

**Train/Test dataset**

We split our dataset into train and test set:

[ ]:



X\_train, X\_test, y\_train, y\_test **=** train\_test\_split( X, y, test\_size**=**0.2, random\_state**=**4)

print ('Train set:', X\_train.shape, y\_train.shape)

print ('Test set:', X\_test.shape, y\_test.shape)

**Modeling (SVM with Scikit-learn)**

**Did you know? IBM Watson Studio lets you build and deploy an AI solution, using the best of open source and IBM software and giving your team a single environment to work in.**[**Learn more here.**](https://cocl.us/ibm_watson_studio_infobox)

The SVM algorithm offers a choice of kernel functions for performing its processing. Basically, mapping data into a higher dimensional space is called kernelling. The mathematical function used for the transformation is known as the kernel function, and can be of different types, such as:

1.Linear

2.Polynomial

3.Radial basis function (RBF)

4.Sigmoid

Each of these functions has its characteristics, its pros and cons, and its equation, but as there's no easy way of knowing which function performs best with any given dataset. We usually choose different functions in turn and compare the results. Let's just use the default, RBF (Radial Basis Function) for this lab.

[ ]:



**from** sklearn **import** svm

clf **=** svm.SVC(kernel**=**'rbf')

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

After being fitted, the model can then be used to predict new values:

[ ]:



yhat **=** clf.predict(X\_test)

yhat [0:5]

**Evaluation**

[ ]:



**from** sklearn.metrics **import** classification\_report, confusion\_matrix

**import** itertools

[ ]:



**def** plot\_confusion\_matrix(cm, classes,

normalize**=False**,

title**=**'Confusion matrix',

cmap**=**plt.cm.Blues):

"""

This function prints and plots the confusion matrix.

Normalization can be applied by setting `normalize=True`.

"""

**if** normalize:

cm **=** cm.astype('float') **/** cm.sum(axis**=**1)[:, np.newaxis]

print("Normalized confusion matrix")

**else**:

print('Confusion matrix, without normalization')

​

print(cm)

​

plt.imshow(cm, interpolation**=**'nearest', cmap**=**cmap)

plt.title(title)

plt.colorbar()

tick\_marks **=** np.arange(len(classes))

plt.xticks(tick\_marks, classes, rotation**=**45)

plt.yticks(tick\_marks, classes)

​

fmt **=** '.2f' **if** normalize **else** 'd'

thresh **=** cm.max() **/** 2.

**for** i, j **in** itertools.product(range(cm.shape[0]), range(cm.shape[1])):

plt.text(j, i, format(cm[i, j], fmt),

horizontalalignment**=**"center",

color**=**"white" **if** cm[i, j] **>** thresh **else** "black")

​

plt.tight\_layout()

plt.ylabel('True label')

plt.xlabel('Predicted label')

[ ]:



*# Compute confusion matrix*

cnf\_matrix **=** confusion\_matrix(y\_test, yhat, labels**=**[2,4])

np.set\_printoptions(precision**=**2)

​

print (classification\_report(y\_test, yhat))

​

*# Plot non-normalized confusion matrix*

plt.figure()

plot\_confusion\_matrix(cnf\_matrix, classes**=**['Benign(2)','Malignant(4)'],normalize**=** **False**, title**=**'Confusion matrix')

You can also easily use the **f1\_score** from sklearn library:

[ ]:



**from** sklearn.metrics **import** f1\_score

f1\_score(y\_test, yhat, average**=**'weighted')

Let's try the jaccard index for accuracy:

[ ]:



**from** sklearn.metrics **import** jaccard\_score

jaccard\_score(y\_test, yhat,pos\_label**=**2)

**Practice**

Can you rebuild the model, but this time with a \_\_linear\_\_ kernel? You can use \_\_kernel='linear'\_\_ option, when you define the svm. How the accuracy changes with the new kernel function?

[ ]:



*# write your code here*

​

Click here for the solution

clf2 **=** svm.SVC(kernel**=**'linear')

clf2.fit(X\_train, y\_train)

yhat2 **=** clf2.predict(X\_test)

print("Avg F1-score: %.4f" **%** f1\_score(y\_test, yhat2, average**=**'weighted'))

print("Jaccard score: %.4f" **%** jaccard\_score(y\_test, yhat2,pos\_label**=**2))

**Want to learn more?**

IBM SPSS Modeler is a comprehensive analytics platform that has many machine learning algorithms. It has been designed to bring predictive intelligence to decisions made by individuals, by groups, by systems – by your enterprise as a whole. A free trial is available through this course, available here: [SPSS Modeler](https://www.ibm.com/analytics/spss-statistics-software?utm_medium=Exinfluencer&utm_source=Exinfluencer&utm_content=000026UJ&utm_term=10006555&utm_id=NA-SkillsNetwork-Channel-SkillsNetworkCoursesIBMDeveloperSkillsNetworkML0101ENSkillsNetwork20718538-2021-01-01)

Also, you can use Watson Studio to run these notebooks faster with bigger datasets. Watson Studio is IBM's leading cloud solution for data scientists, built by data scientists. With Jupyter notebooks, RStudio, Apache Spark and popular libraries pre-packaged in the cloud, Watson Studio enables data scientists to collaborate on their projects without having to install anything. Join the fast-growing community of Watson Studio users today with a free account at [Watson Studio](https://www.ibm.com/cloud/watson-studio?utm_medium=Exinfluencer&utm_source=Exinfluencer&utm_content=000026UJ&utm_term=10006555&utm_id=NA-SkillsNetwork-Channel-SkillsNetworkCoursesIBMDeveloperSkillsNetworkML0101ENSkillsNetwork20718538-2021-01-01)