# Springboard Intensive Data Science Course Breast Cancer Classification Project Final Report 20 March 2017

Mentor: Pavitraa Parthasarathy

Student: Jing-Rui Li

Final project requirement:

Github code: https://github.com/jl3392/Capstone-Design/blob/master/final\_project\_code.ipynb

### 1.Introduction - Background

Breast cancer is one of the most dangerous form of cancers in the world. Annually, Breast caner kills an estimated 20,000 people in the US. If breast cancer is recognized and treated early, it is almost curable, with a 80 to 90% 5-year survival rate when treated with simple excision. Early diagnosis of the breast cancer will help save thousands of lives.

The Breast Cancer project aims to develop a system that takes breast features as inputs, and outputs the likelihood that the patient is malignant, in order to facilitate early diagnosis of breast cancer. Unique and relevant features such as statistical measures (mean, standard deviation and covariance) of the breast's shape information (area, diameter, compactness and asymmetry) was extracted from the pre-processed images. Texture information, obtained from a gray level covariance matrix was also included. A trained classifier will then compute the likelihood that the lesion is malignant and return this percentage to the user.

#### Link to the dataset:

http://scikit-learn.org/stable/modules/generated/sklearn.datasets.load breast cancer.html

#### 2. Dataset

This project uses the Breast Cancer Wisconsin (Diagnostic) Database to create a classifier that can help diagnose patients. Features are computed from a digitized image of a fine needle aspirate (FNA) of a breast mass. They describe characteristics of the cell nuclei present in the image. The dataset of the breast cancer information can be accessed from build-in python package sklearn.dataset.load\_breast\_cancer. (Link to the dataset:

http://scikit-learn.org/stable/modules/generated/sklearn.datasets.load breast cancer.html).

#### **Data Set Characteristics:**

:Number of Instances: 569

:Number of Attributes: 30 numeric, predictive attributes and the class

:Attribute Information:

- radius (mean of distances from center to points on the perimeter)
- texture (standard deviation of gray-scale values)
- perimeter
- area
- smoothness (local variation in radius lengths)
- compactness (perimeter^2 / area 1.0)
- concavity (severity of concave portions of the contour)
- concave points (number of concave portions of the contour)
- symmetry
- fractal dimension ("coastline approximation" 1)

The mean, standard error, and "worst" or largest (mean of the three largest values) of these features were computed for each image, resulting in 30 features. For instance, field 3 is Mean Radius, field 13 is Radius SE, field 23 is Worst Radius.

## - class:

- WDBC-Malignant
- WDBC-Benign

# :Summary Statistics:

Min Max

Min Max				
radius (mean):	6.981 28.11			
texture (mean):	9.71 39.28			
perimeter (mean):	43.79 188.5			
area (mean):	143.5 2501.0			
smoothness (mean):	0.053 0.163			
compactness (mean):	0.019 0.345			
concavity (mean):	0.0 0.427			
concave points (mean):	0.0 0.201			
symmetry (mean):	0.106 0.304			
fractal dimension (mean):	0.05 0.097			
radius (standard error):	0.112 2.873			
texture (standard error):	0.36 4.885			
perimeter (standard error):	0.757 21.98			
area (standard error):	6.802 542.2			
smoothness (standard error	c): 0.002 0.031			
compactness (standard erro	or): 0.002 0.135			
concavity (standard error):	0.0 0.396			
concave points (standard e	rror): 0.0 0.053			
symmetry (standard error)	0.008 0.079			
fractal dimension (standard	d error): 0.001 0.03			
radius (worst):	7.93 36.04			
texture (worst):	12.02 49.54			
perimeter (worst):	50.41 251.2			
area (worst):	185.2 4254.0			
smoothness (worst):	0.071 0.223			
compactness (worst):	0.027 1.058			
concavity (worst):	0.0 1.252			
concave points (worst):	0.0 0.291			
symmetry (worst):	0.156 0.664			
fractal dimension (worst):	0.055 0.208			

\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_

:Missing Attribute Values: None

:Class Distribution: 212 - Malignant, 357 - Benign

:Creator: Dr. William H. Wolberg, W. Nick Street, Olvi L. Mangasarian

:Donor: Nick Street

:Date: November, 1995

This is a copy of UCI ML Breast Cancer Wisconsin (Diagnostic) datasets. https://goo.gl/U2Uwz2

#### References

-----

- W.N. Street, W.H. Wolberg and O.L. Mangasarian. Nuclear feature extraction for breast tumor diagnosis. IS&T/SPIE 1993 International Symposium on Electronic Imaging: Science and Technology, volume 1905, pages 861-870, San Jose, CA, 1993.
- O.L. Mangasarian, W.N. Street and W.H. Wolberg. Breast cancer diagnosis and prognosis via linear programming. Operations Research, 43(4), pages 570-577, July-August 1995.
- W.H. Wolberg, W.N. Street, and O.L. Mangasarian. Machine learning techniques to diagnose breast cancer from fine-needle aspirates. Cancer Letters 77 (1994) 163-171.

### 3. Data Visualization and Pre-processing

Firstly, Let's take a look at the number of Benign and Maglinant cases from the dataset. From the output shown below, there are 357 cases are benign while 212 cases are malignant.

print(df.groupby('diagnosis').size())

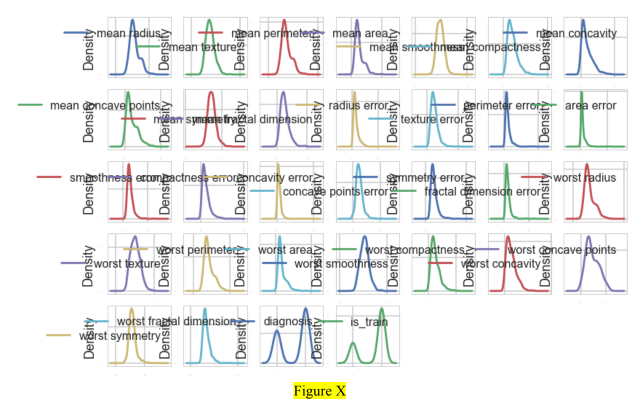
diagnosis

0 212

1 357

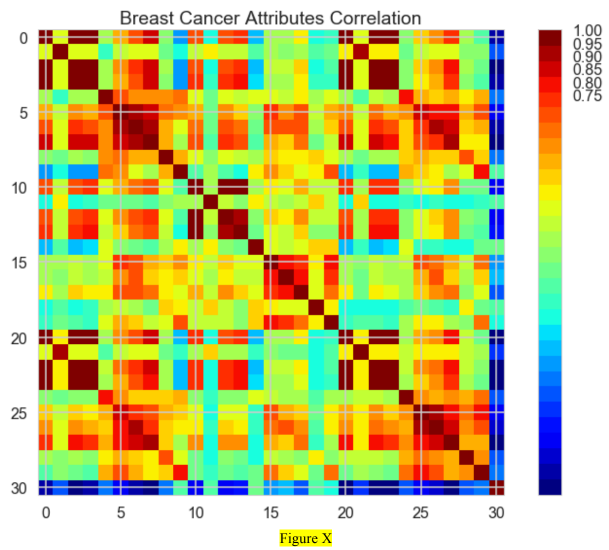
Next, I visualized the data using density plots to get a sense of the data distribution(Kernel density estimation). The kernel density estimate may be less familiar, but it can be a useful tool for plotting the shape of a distribution. Like the histogram, the KDE plots encodes the density of observations on one axis with height along the other axis.

.



From the outputs above, we can see the data shows a general Gaussian distribution.

Then, It is good to check the correlations between the attributes.



From the output graph below, The red around the diagonal suggests that attributes are correlated with each other. The yellow and green patches suggest some moderate correlation and the blue boxes show negative correlations.

#### 3. Baseline Algorithm Checking with cross validation

From the dataset, we will analysis and build a model to predict if a given set of symptoms lead to breast cancer. This is a binary classification problem, and a few algorithms are appropriate for use. Since we do not know which one will perform the best at the point, we will do a quick test on the few appropriate algorithms with default setting to get an early indication of how each of them perform. We will use 5 and 10 fold cross validation for each testing.

The following non-linear algorithms will be used, namely: Classification and Regression Trees (CART), Linear Support Vector Machines (SVM), Gaussian Naive Bayes (NB) and k-Nearest Neighbors (KNN).

#### Result for 5 fold cross validation

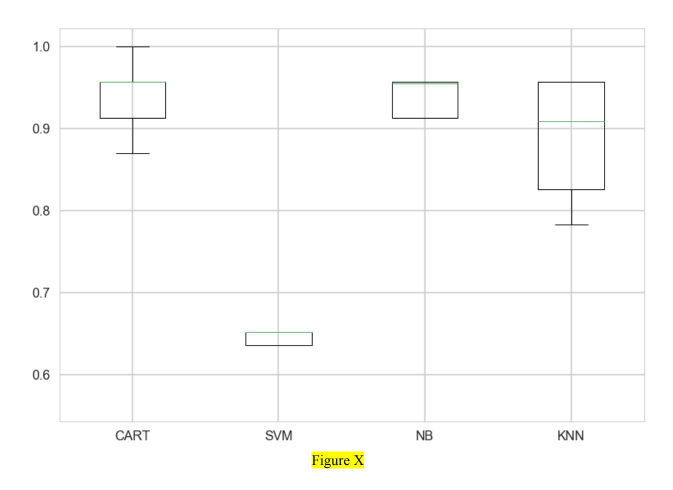
```
CART: 0.903557 (0.063812) (run time: 0.029673)

SVM: 0.657708 (0.070245) (run time: 0.039123)

NB: 0.921344 (0.050438) (run time: 0.012046)

KNN: 0.886166 (0.070369) (run time: 0.015470)
```

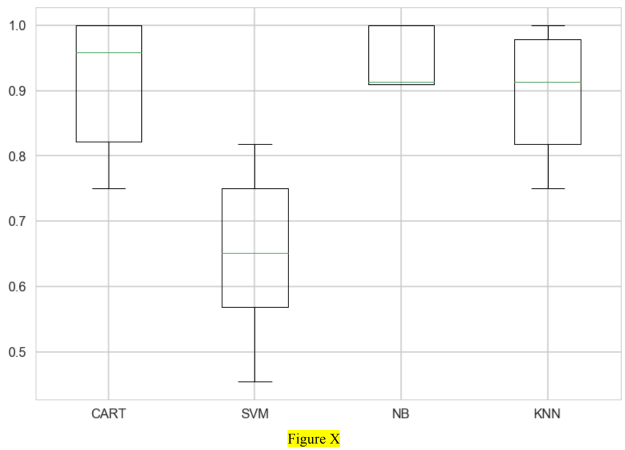
# Performance Comparison



# Result for 10 fold cross validation

CART: 0.913636 (0.094208) (run time: 0.026272)		
SVM: 0.655303 (0.108957) (run time: 0.034203)		
NB: 0.930303 (0.073434) (run time: 0.011217)		
KNN: 0.894697 (0.085405) (run time: 0.021682)		

# Performance Comparison

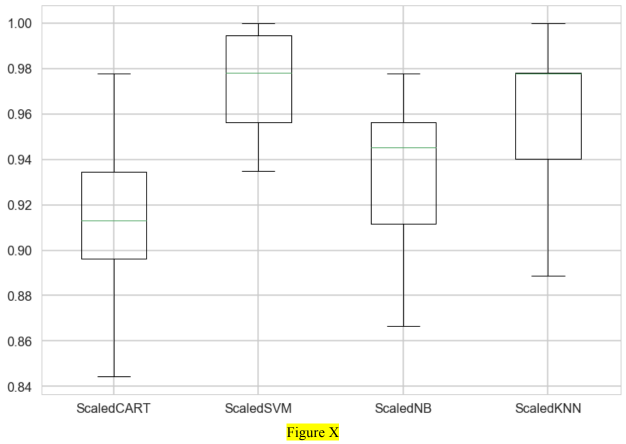


From the initial run, it looks like NB, KNN and CART performed the best given the dataset (all above 89% mean accuracy). Support Vector Machine has a surprisingly bad performance here. However, if we standardize the input dataset, it's performance should improve.

The performance of the few machine learning algorithm could be improved if a standardized dataset is being used. The improvement is likely for all the models. I will use pipelines that standardize the data and build the model for each fold in the cross-validation test harness. That way we can get a fair estimation of how each model with standardized data might perform on unseen data.

ScaledCART: 0.914396 (0.037331) (run time: 0.113696)		
ScaledSVM: 0.964879 (0.038621) (run time: 0.074927)		
ScaledNB: 0.931932 (0.038625) (run time: 0.033946)		
ScaledKNN: 0.958357 (0.038595) (run time: 0.053220)		

# Performance Comparison



From the performance figure above, we can see the SVM algorithm has a great improvement with the standardized data.

Thus, I decide to focus on SVM for the algorithm tuning. We can tune two key parameter of the SVM algorithm - the value of C and the type of kernel. The default C for SVM is 1.0 and the kernel is Radial Basis Function (RBF). We will use the grid search method using 10-fold cross-validation with a standardized copy of the sample training dataset. We will try over a combination of C values and the following kernel types 'linear', 'poly', 'rbf' and 'sigmoid'.

```
Best: 0.969231 using {'C': 2.0, 'kernel': 'rbf'}
0.964835 (0.026196) with: {'C': 0.1, 'kernel':
                                                'linear'}
0.826374 (0.058723) with: {'C': 0.1, 'kernel':
                                               'poly'}
0.940659 (0.038201) with: {'C': 0.1,
                                     'kernel':
0.949451 (0.032769) with: {'C': 0.1,
                                     'kernel':
                                     'kernel':
0.962637 (0.029474) with: {'C': 0.3,
                                                'linear'}
                                     'kernel':
0.868132 (0.051148) with: {'C': 0.3,
                                                'poly'}
0.958242 (0.031970) with: {'C': 0.3,
                                     'kernel':
0.953846 (0.043579) with: {'C': 0.3, 'kernel': 'sigmoid'}
0.956044 (0.030933) with: {'C': 0.5, 'kernel': 'linear'}
0.881319 (0.050677) with: {'C': 0.5, 'kernel': 'poly'}
0.964835 (0.029906) with: {'C': 0.5, 'kernel':
```

```
0.953846 (0.026785) with: {'C': 0.5, 'kernel': 'sigmoid'}
0.953846 (0.031587) with: {'C': 0.7,
                                     'kernel':
                                                'linear'}
0.885714 (0.038199) with: {'C': 0.7,
                                     'kernel': 'poly'}
0.967033 (0.037271) with: {'C': 0.7, 'kernel': 'rbf'}
0.953846 (0.028513) with: {'C': 0.7,
                                     'kernel': 'sigmoid'}
0.951648 (0.028834) with: {'C': 0.9,
                                     'kernel': 'linear'}
0.887912 (0.038950) with: {'C': 0.9,
                                     'kernel':
0.967033 (0.037271) with: {'C': 0.9,
                                     'kernel': 'rbf'}
0.958242 (0.033272) with: {'C': 0.9,
                                     'kernel': 'sigmoid'}
0.953846 (0.026546) with: {'C': 1.0,
                                     'kernel': 'linear'}
0.890110 (0.038311) with: {'C': 1.0,
                                     'kernel':
                                               'poly'}
                                     'kernel':
0.967033 (0.033027) with: {'C': 1.0,
                                                'rbf'}
0.964835 (0.031496) with: {'C': 1.0,
                                     'kernel': 'sigmoid'}
0.956044 (0.025765) with: {'C': 1.3, 'kernel': 'linear'}
0.894505 (0.039427) with: {'C': 1.3,
                                     'kernel': 'poly'}
0.967033 (0.028188) with: {'C': 1.3,
                                     'kernel': 'rbf'}
0.958242 (0.033224) with: {'C': 1.3,
                                     'kernel':
                                                'sigmoid'}
0.958242 (0.024765) with: {'C': 1.5,
                                     'kernel': 'linear'}
0.896703 (0.039791) with: {'C': 1.5,
                                     'kernel': 'poly'}
0.967033 (0.028188) with: {'C': 1.5,
                                     'kernel': 'rbf'}
0.945055 (0.034215) with: {'C': 1.5,
                                     'kernel': 'sigmoid'}
0.956044 (0.021766) with: {'C': 1.7,
                                     'kernel':
                                               'linear'}
                                     'kernel': 'poly'}
0.903297 (0.033409) with: {'C': 1.7,
0.967033 (0.024479) with: {'C': 1.7,
                                     'kernel': 'rbf'}
0.956044 (0.035354) with: {'C': 1.7,
                                     'kernel': 'sigmoid'}
0.956044 (0.021766) with: {'C': 2.0,
                                     'kernel': 'linear'}
                                     'kernel':
0.909890 (0.033680) with: {'C': 2.0,
0.969231 (0.022370) with: {'C': 2.0, 'kernel': 'rbf'}
0.947253 (0.028116) with: {'C': 2.0, 'kernel': 'sigmoid'}
```

We can see the most accurate configuration was SVM with an RBF kernel and C=2.0, with the accuracy of 96.92%.

## 4. Applying various forecasting models

#### 4.1 SVM Analysis

From the baseline processing, I selected SVM out of four algorithms as the most accurate model for forecasting the breast cancer data. Using the SVM tuning method, I further selected the parameters with RBF kernel and C=2.0. Now, I would like to use the selected parameters to estimate the accuracy of the test dataset.

# **SVM Test Dataset Result**

# Accuracy score

```
print("Accuracy score %f" % accuracy score(Y test, predictions))
print(classification report(Y test, predictions))
Accuracy score 0.991228
precision
             recall f1-score
                                 support
                  0.97
                                       0.99
                                                    39
          0
                             1.00
          1
                  1.00
                             0.99
                                       0.99
                                                    75
```

```
avg / total 0.99 0.99 0.99 114
```

#### Confusion Matrix

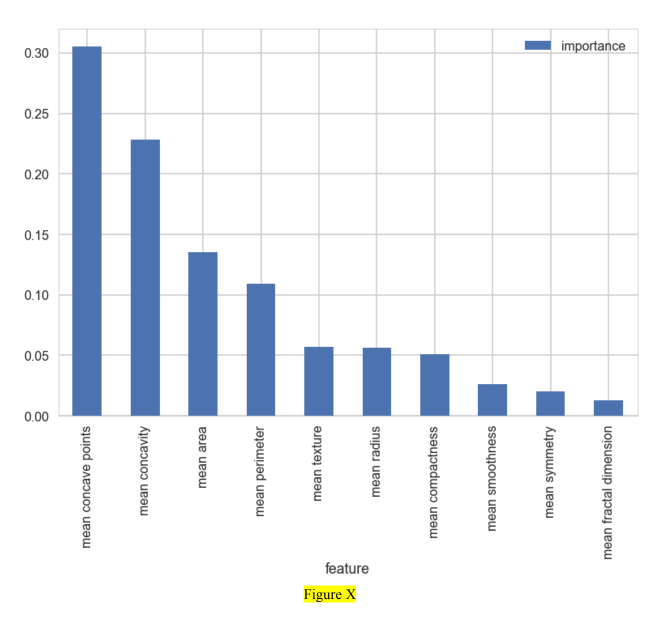
```
print(confusion_matrix(Y_test, predictions))
[[39  0]
  [ 1 74]]
```

We can see that we achieve an accuracy of 99.1% on the held-out test dataset. From the confusion matrix, there is only 1 case of mis-classification. The performance of this algorithm is expected to be high given the symptoms for breast cancer should exhibit certain clear patterns.

# 4.2 Random Forest Analysis

Although we found that SVM is one of the best model for the dataset, I would like to further explore the dataset and rank the importance of each feature using another algorithm-random forest analysis. Likewise, the data was split into test and train datasets for the testing.

impor featu	ctance	
		0 205
	concave points	0.305
mean	concavity	0.228
mean	area	0.135
mean	perimeter	0.109
mean	texture	0.057
mean	radius	0.056
mean	compactness	0.051
mean	smoothness	0.026
mean	symmetry	0.020
mean	fractal dimension	0.013



Secondly, I did confusion matrix analysis. The columns are the diagnosis we predicted for the test data and the rows are the actual result for the test data.

Predicted Dignosis	benign	malignant			
Actual Diagnosis					
0	4	52			
1	95	3			

# 5. Thoughts and Conclusion

In this project, I compared prediction accuracy among four different algorithms. Through the comparison, I found that SVM did not perform well before standardize the data, however, it turned out to be the best estimation algorithm with cross validation score of 96.4% after standardization. It reminded me that when doing analysis on an unfamiliar dataset, it is critical to standardize the data before making any comparison or further analysis.

Later through the project, I did tuning SVM and select RBF and C=2.0 as the best classifiers with accuracy of 96.92%. The tuning process helped me to find the best parameters for the dataset, which help to improve the accuracy score in the test data. This reminds me the importance of selecting the appropriate classifiers before doing the prediction.

Finally, the feature ranking helps me get a better understanding of the importance of each feature. This will help improve the model in the further to make predictions when some features are missing(modify the weight of each feature).

#### Citation

https://pandas.pydata.org/pandas-docs/version/0.18.1/visualization.html

http://scikit-

learn.org/stable/modules/generated/sklearn.ensemble. Random Forest Classifier.html

 $\frac{https://stackoverflow.com/questions/41458834/how-is-scikit-learn-cross-val-predict-accuracy-score-calculated}{}$ 

https://www.programcreek.com/python/example/91159/sklearn.model\_selection.cross\_val\_predict\_