Capstone 2 Milestone Report: Breast Cancer in Wisconsin

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# Problem

There are numeric different characteristics of tumors that the breast cancer patients would have. The accuracy and efficiency of the diagnosis could significantly impact the patients’ recovery and potentially decrease the complexity of the treatment. Hence, it is important to help doctors to identify the characteristics of the tumors that patients have in a timely manner in order to provide necessary healthcare.

# Goal and Utility

The goal of this project is to analyze the symptoms and tumors characteristics, including radius, texture, perimeter, area, smoothness, compactness, concavity, concave points, symmetry, and fractal dimension that breast cancer could have at different stage, and provide prediction on whether the tumor is malignant or benign. The analysis and the prediction can quickly help doctor prescribe necessary treatment and healthcare.

# Data

The data set for this project contains 569 observations including patient ID number, and the features of the cell nucleus, such as radius, texture, perimeter, area, smoothness, compactness, concavity, concave points, symmetry, and fractal dimension.

The data set is available on UCI Machine Learning Repository. The page contains information about the data set and its source and relevant scientific publications:

<http://archive.ics.uci.edu/ml/datasets/Breast+Cancer+Wisconsin+%28Diagnostic%29>

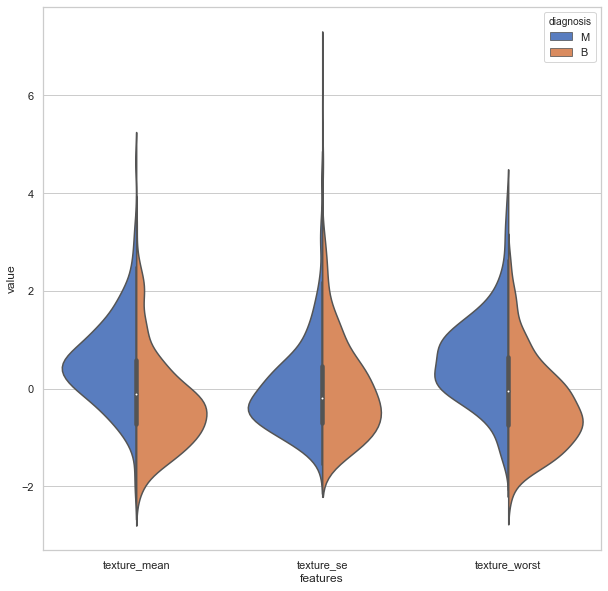
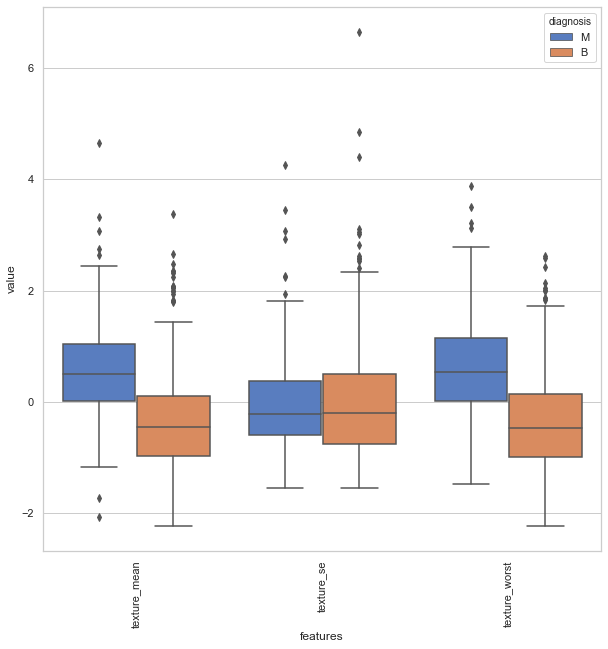
# Approach

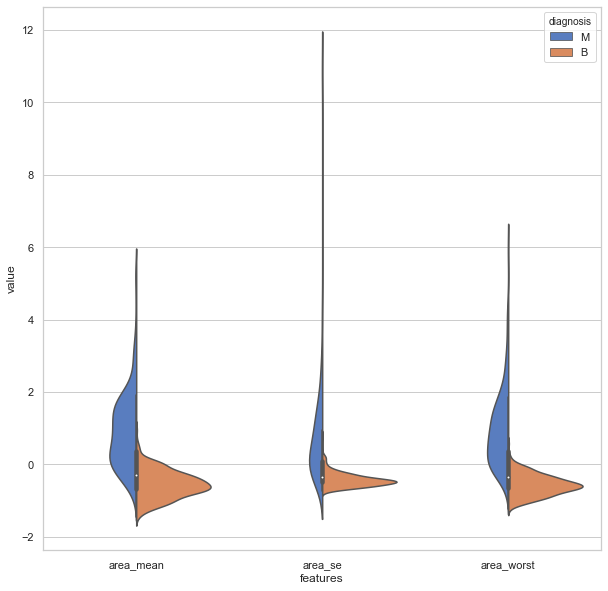
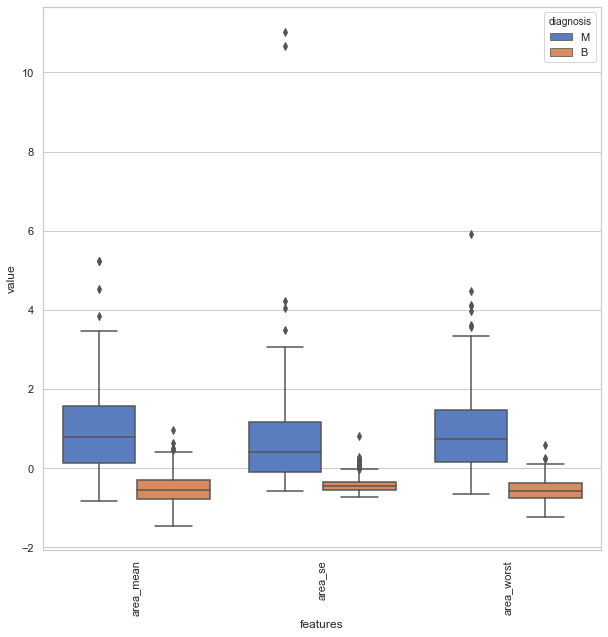
It is a supervised classification task. The diagnosis (whether the tumor is malignant or benign) will be our target variable, while the characteristics of the cell nucleus will be used as the predictive variables. Logistic Regression, K-Nearest Neighbor, Support Vector Machine, and Random Forest Classification are some of the machine learning models that could potentially be used in this project. Data exploratory analysis will provide deeper understandings of the data and address the problem more clearly, and cross-validation could ensure the outcomes of the model is reliable.

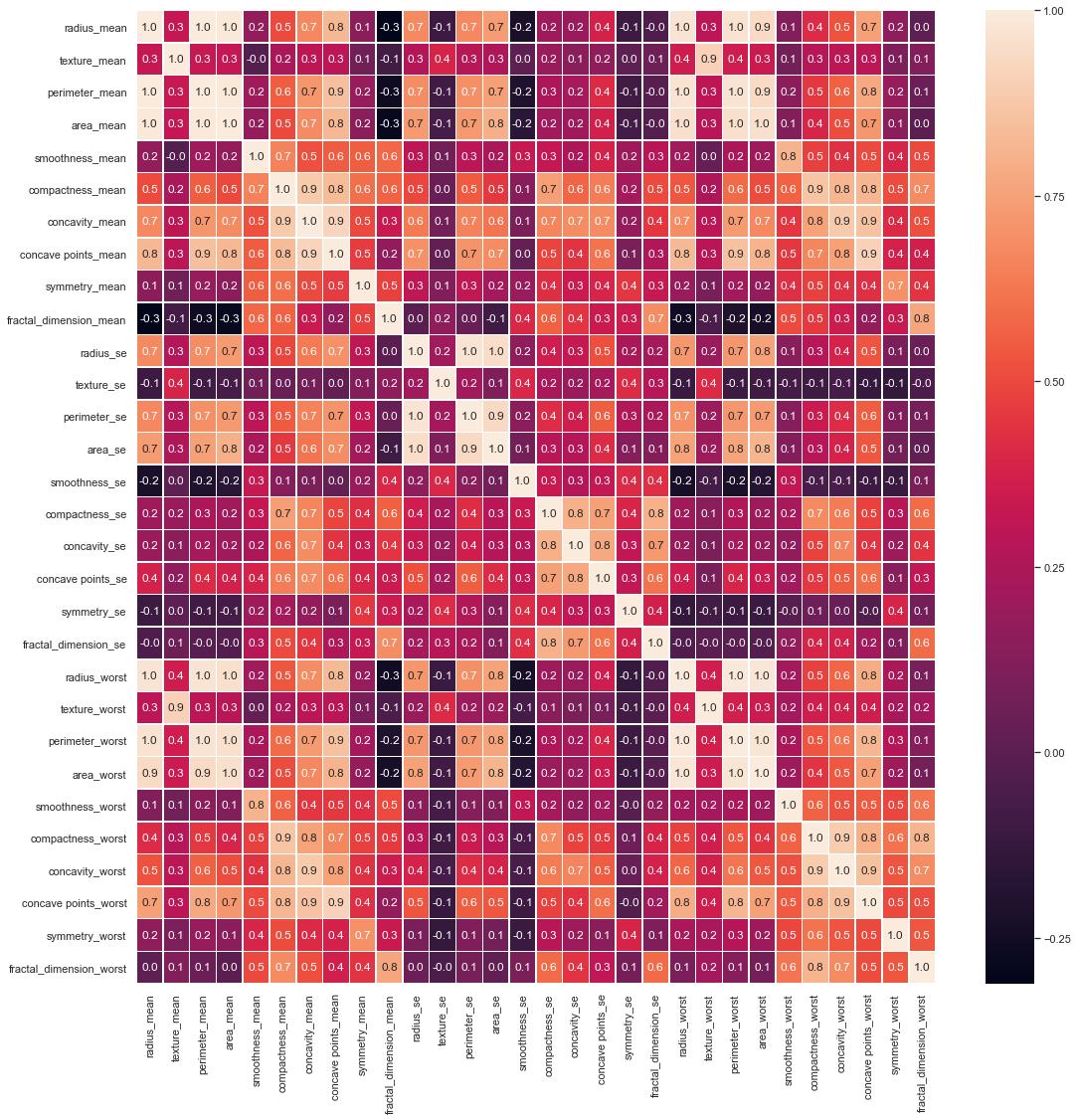
# Deliverables

The deliverables for the project will include the source code, the data set, and a paper addressing the purpose, approach, findings and results of the project.

# Exploratory Data Analysis

The data set contains 32 columns and 569 data points. There are no missing values (null) in any of the features, hence, it is not required to do much data imputation. In the univariate analysis, it is shown that there is no categorical variable, and 30 continuous variables in the data set. I excluded the feature “id” because the characteristics of id does not imply much information to the algorithm. Next, I am interested in knowing the distribution of each feature and identify if there is any outliers within the feature. The violin plots and box plots come in handy as it can quickly tell the distribution and outliers by feature and by the diagnosis (the target variable).

The above figures show the distributions of texture\_mean, texture\_se, and texture\_worst features after normalization. It shows that the two-diagnosis distributed quite equally in both cases, indicating that within these feature, it is difficult to tell whether the tumor is benign or malignant just by looking at the data. It also indicates that in the feature selection, we could possibly just include one of these features if they are highly correlated to each other. On the other hand, the figures below show a completely different story. It is easier to tell the two diagnosis just from looking at the data as the distributions of the two different groups are in different ranges. They are likely be selected as the features for the model.

Next, in the bivariate analysis, we can look at the correlation between each feature. The features that are more highly correlated could be “combined”, and just consider one of them to be the feature in the model. In this case, we can see that radius\_mean with concave points\_mean, area\_worst, radius\_se, perimeter\_se, area\_se, and concave points\_worst. The same ideas apply to all other features. Hence, in this stage, we can come up with a set of features that will be input into the model, which are radius mean, texture mean, smoothness mean, compactness mean, concavity mean, symmetry mean, fractal\_dimension\_mean, radius se, texture se, smoothness se, compactness se, concavity se, concave points se, symmetry se, fractal dimension se, smoothness worst, compactness worst, concavity worst, symmetry worst, and fractal dimension worst.