# Breast Cancer Classification

Breast cancer (BC) is one of the most common cancers among women worldwide, representing most new cancer cases and cancer-related deaths according to global statistics, making it a significant public health problem in today’s society.

**Objective**The early diagnosis of BC can improve the prognosis and chance of survival significantly, as it can promote timely clinical treatment to patients. Further accurate classification of benign tumors can prevent patients undergoing unnecessary treatments. Thus, the correct diagnosis of BC and classification of patients into malignant or benign groups is the subject of much research. Because of its unique advantages in critical features detection from complex BC datasets, machine learning (ML) is widely recognized as the methodology of choice in BC pattern classification and forecast modelling.

**The Data :** UCI Machine Learning Repository for BC [dataset](http://archive.ics.uci.edu/ml/datasets/breast+cancer+wisconsin+%28diagnostic%29)The dataset used in this story is publicly available and was created by Dr. William H. Wolberg, physician at the University of Wisconsin Hospital at Madison, Wisconsin, USA. To create the dataset Dr. Wolberg used fluid samples, taken from patients with solid breast masses and an easy-to-use graphical computer program called Xcyt, which is capable of perform the analysis of cytological features based on a digital scan. The program uses a curve-fitting algorithm, to compute ten features from each one of the cells in the sample, then it calculates the mean value, extreme value and standard error of each feature for the image, returning a 30 real-valuated vector

## **Data Wrangling**

We conduct the following steps to clean up data and pick useful features.

1. Search for missing values:

* **Unnamed: 32** feature includes NaN, so we do not need it.

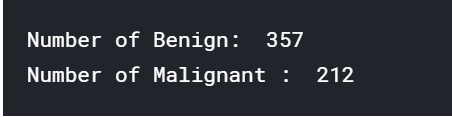
1. Drop duplicates:

* Didn’t find any duplicates.

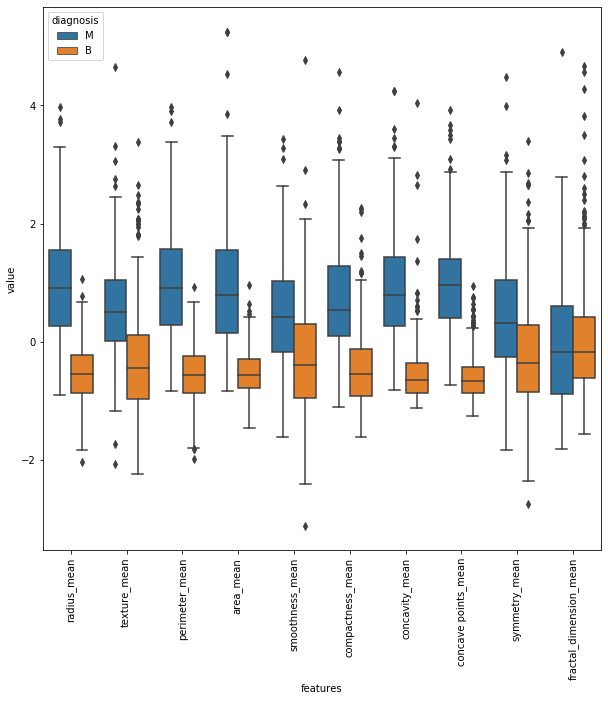
1. **Diagnosis** is our class label.
2. There is an **id** that can not be used for analysis and we drop it.
3. Because differences between values of features are very high to observe on plot (like the **area\_mean** feature's max value is 2500 and **smoothness\_mean** features' max 0.16340.), so we need **standardisation before visualization, feature selection or classification.**

At this point, we do not have any idea about other feature names.

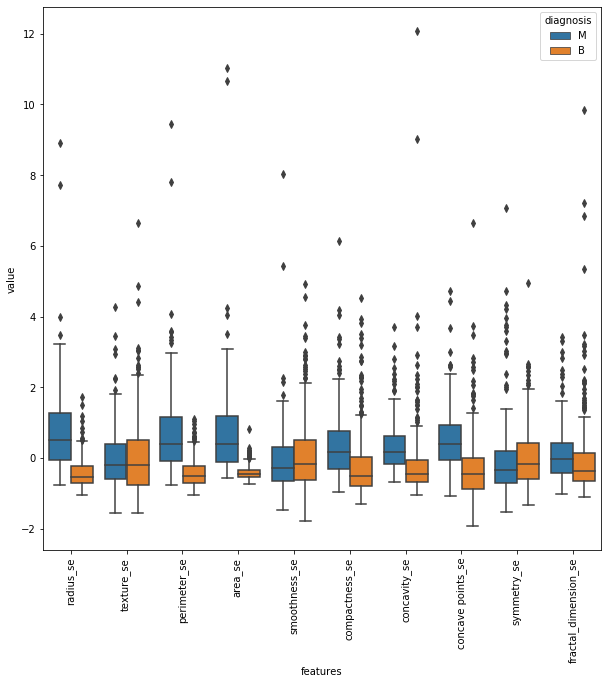
## **Data Exploration**



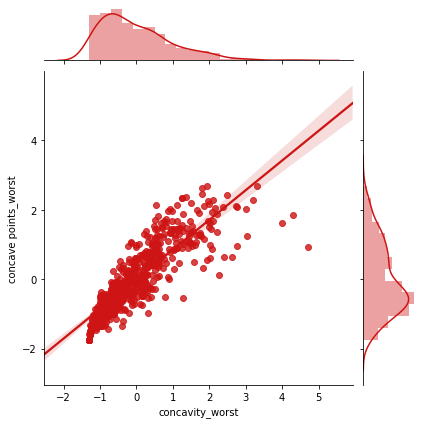
 In Box Plot, we plot features in 3 group and each group includes 10 features to observe better.

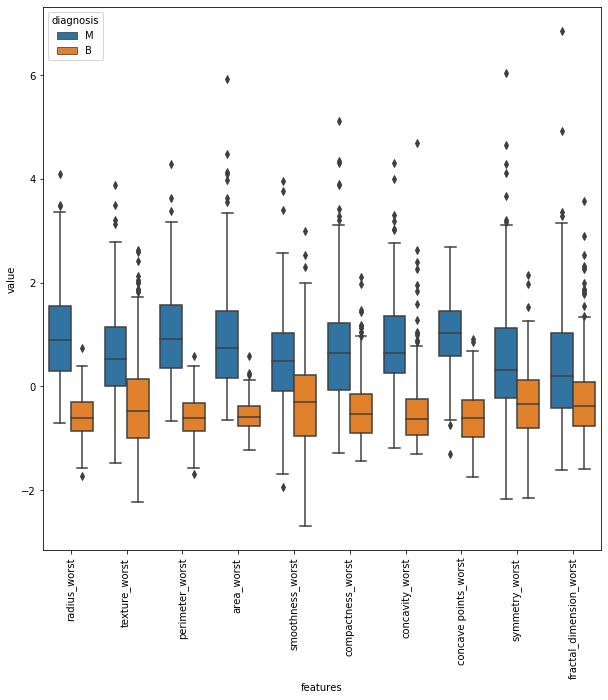


in **texture\_mean** feature, median of the Malignant and Benign looks like separated so it can be good for classification. However, in **fractal\_dimension\_mean** feature, median of the Malignant and Benign does not looks like separated so it does not give good information for classification.

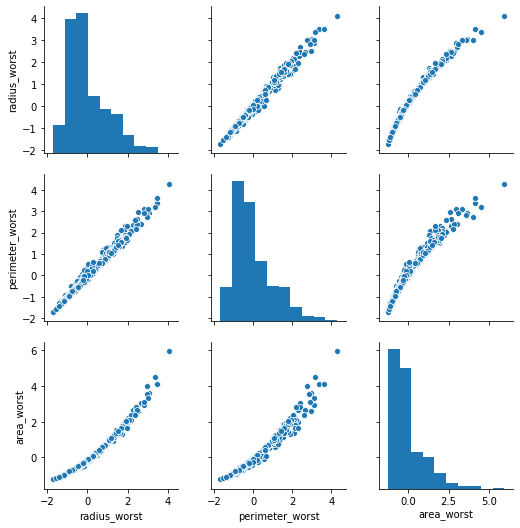


in **concave points\_se and concavity\_se** features, median of the Malignant and Benign looks like separated so it can be good for classification. However, in **texture\_se** feature, median of the Malignant and Benign does not looks like separated so it does not give good information for classification. Also looks like there is correlation between **concavity\_worst and concave point\_worst.**

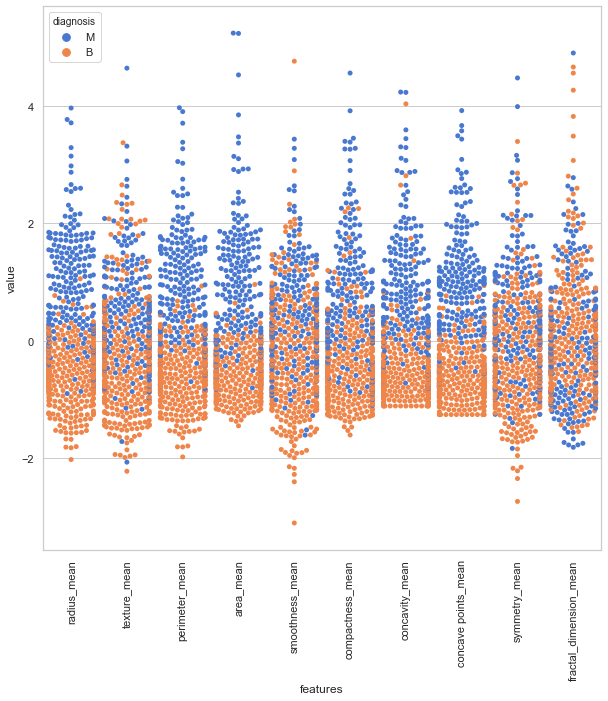
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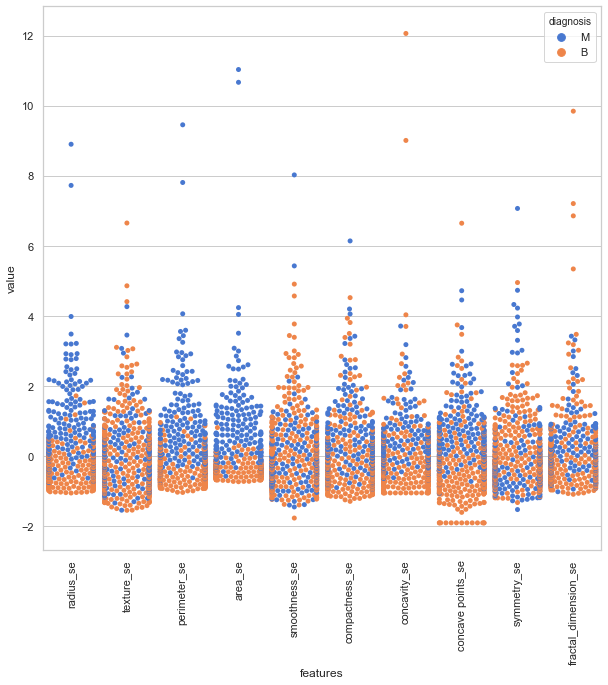
 Also it seems **radius\_worst**, **perimeter\_worst** and **area\_worst** are correlated as it can be seen pair plot. We use these for feature selection.



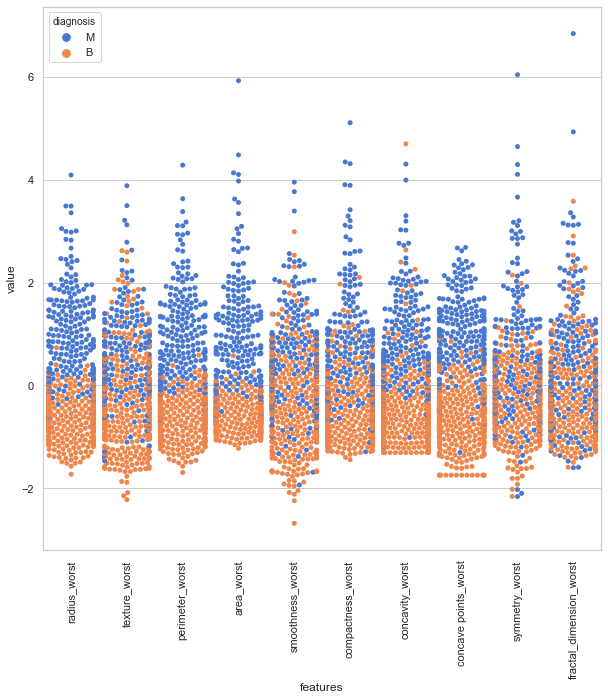
Now in swarm Plot, we plot features in 3 group and each group includes 10 features to observe better.



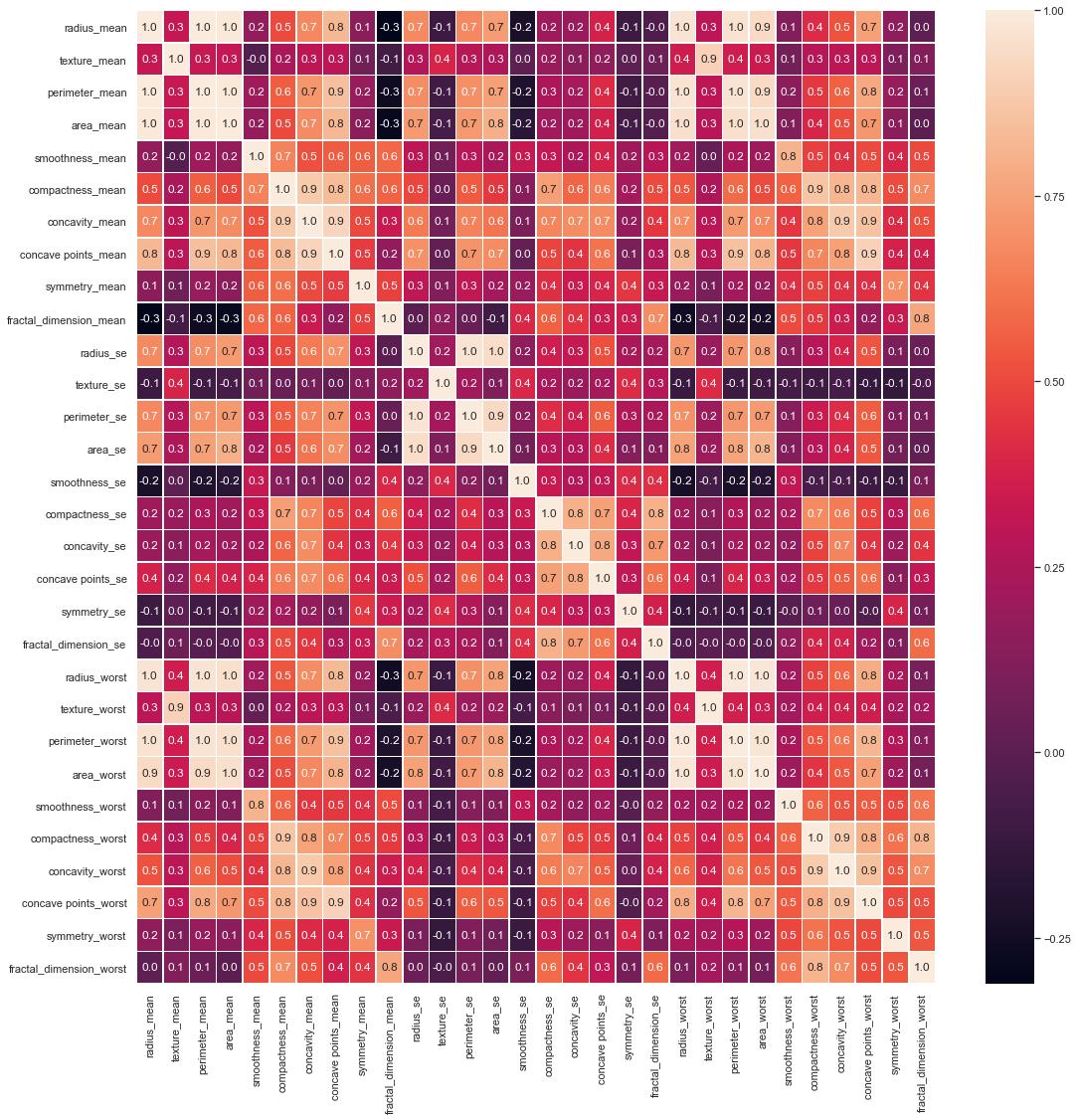
I think **perimeter\_mean** and **area\_mean** looks like malignant and benign are seprated not totaly but mostly.



I think **area\_se** looks like malignant and benign are seprated not totaly but mostly. Hovewer, **smoothness\_se** looks like malignant and benign are mixed so it is hard to classfy while using this feature.



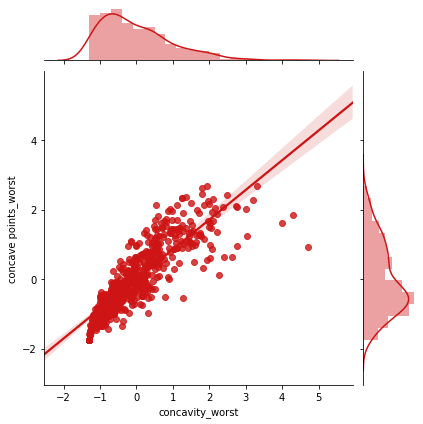
I think area\_worst and perimeter\_worst looks like malignant and benign are seprated not totaly but mostly. Hovewer, texture\_worst looks like malignant and benign are mixed so it is hard to classfy while using this feature.



## **EDA – Inferential Statistics**

**In the following EDA, I try to perfom analyze correlation between variable of concavity\_worst and concave point\_worst.**

# ****Initial observations for Correlation between concavity\_worst and concave point\_worst****



**Pearson correlation coefficient is 0.86, which means there is strong correlation between these two variables.**

# Setup hypothesis test

**Ho : The variable of concavity\_worst and concave point\_worst are independent**

**Ha : The variable of concavity\_worst and concave point\_worst are correlated**

**Statistical significance for 𝛼 = 0.01**

**Test Statistic: Pearson correlation coefficient**

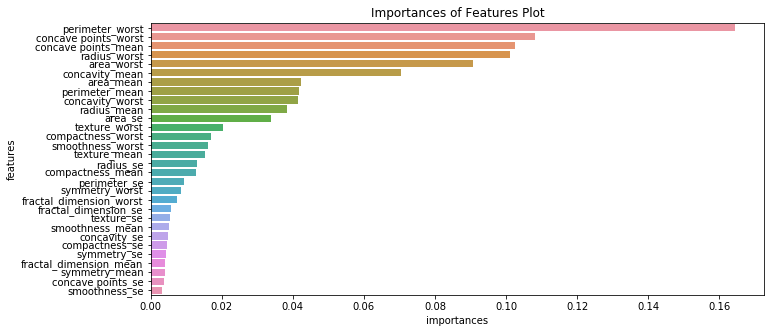
To do so, permute the concavity\_worst but leave the concave point\_worst values fixed. This simulates the hypothesis that they are totally independent of each other. For each permutation, compute the Pearson correlation coefficient and assess how many of your permutation replicates have a Pearson correlation coefficient greater than the observed one.**As p-value less than**α**, we can reject the null hypothesis and accept alternative hypothesis which means that there is a correlation between variable of concavity\_worst and concave point\_worst**

## **Machine Learning: Overview**

* Random Forest (RF) was used to find out importance of features.
* Boruta and Variance Inflation Factor (VIF) used for features selection.
* XGBoost was used for prediction.
* Overall strategy:
  1. Use Random Forest to find out importance of components
  2. Reduce features with VIF compare with Boruta has more accuracy in this case
  3. Use XGBoost on important components of all data.
  4. Split the data into training and test datasets
  5. Train various learning models, with cross-validated hyperparameter tuning
  6. Apply the models to the test dataset to get measures of performance

## **Random Forest**

I visualize the importance of features using random forest model.



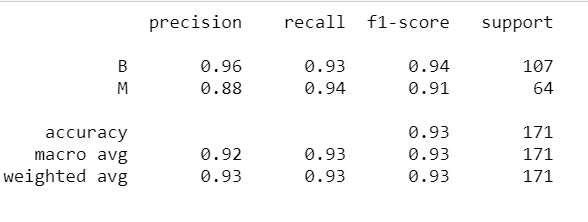
## **Feature Selection with Bruta and VIF**

With using Bruta, features has been reduced to 23 features and with using Variance Inflation Factor (VIF) features has been reduced to 13 features.

## **XGBoost**

* The best estimator was selected by Randomized Grid Search with Stratified Shuffle Split Cross Validation.
* The best hyperparameters included a tree count of 250, max depth of 10, and allowing trees to consider all features

# The results of the XGBoost algorithm with Boruta feature selection





# The results of the XGBoost algorithm with Boruta feature selection

