# Models Comparison for Liver Cirrhosis Survival Prediction

## Cleto Pellegrino

# August 11, 2024

## Contents

1	Introduction	2
2	Dataset	2
3	Data Visualization3.1 Preliminary Adjustments3.2 Plot Analysis	
4	Preprocessing	4
5	Models Definition and Evaluation         5.1 Validation of the Assumptions	<b>6</b> 7
6	Final Results	7
7	Appendix	8
8	References	9

#### 1 Introduction

The liver is the **second largest organ** in the human body and **one of the most important for human health**. Cirrhosis is a progressive condition that puts both a person's liver and life at risk. [1]

Different models can be compared to determine which one best predicts the likelihood of a patient to survive against the disease.

#### 2 Dataset

This study utilizes the Cirrhosis Prediction Dataset available on **Kaggle**[2], comprising data collected from the **Mayo Clinic trial in primary biliary cirrhosis** (*PBC*) of the liver conducted between 1974 and 1984.

The dataset includes various information such as Cholesterol, Triglycerides, and, crucially, the patient's **Status**. In total, there are 418 observations with 20 attributes.

#### 3 Data Visualization

#### 3.1 Preliminary Adjustments

Before commencing with a complete dataset visualization, several **adjustments** were made for some features:

- All features of type "object" were converted to "category" (in order to plot barplots and use the .describe function provided in pandas with specific characteristics for this type of features).
- The "ID" column was **dropped** since it's useless for the analysis, solely serving as an **identifier**.
- The "Stage" feature, originally represented as *float64*, was converted to **categorical**, since it can be considered the **name** of the current liver condition.
- Age, initially reported in days, was converted to years using the formula: round( $\frac{x}{365}$ , 0). This was made just for visualization reasons.
- Status C was converted to "High chance", Status CL was converted to "Transplant needed" and Status D was converted to "Low chance" (again, still just for visualization reasons).

#### 3.2 Plot Analysis

By looking at the **heatmap** created by calculating the **mutual information** between the features (Figure 1),

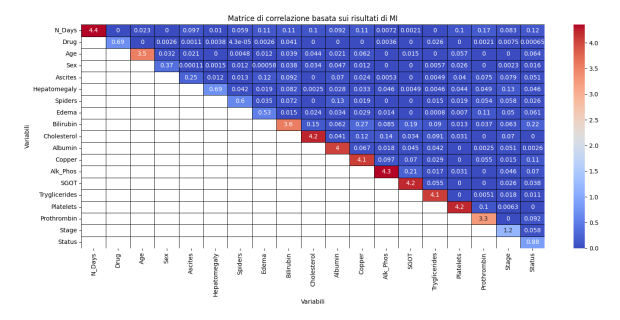


Figure 1: Heatmap of the mutual information between features.

one can see that there is **no strong positive/negative correlation** between them. For this reason, we cannot drop features based on their correlation, indeed no couples contain the **same type of information**.

Figure 2 demonstrates the imbalanced distribution of the dataset labels.

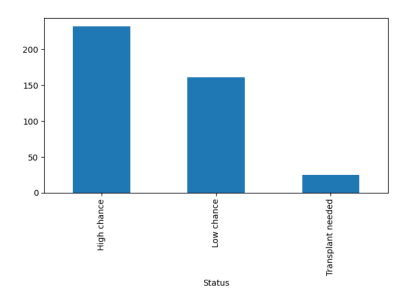


Figure 2: Distribution of patient statuses.

This is an important information not only for the **correct implementation of the pipelines** (imputing missing values, re-balancing), but also to decide **how we will evaluate their performances**, in order to select the best at predicting the survival chance of the patient.

# 4 Preprocessing

The Train set showed lots of *NaN* values, but due to the low total number of observations, removing all of them was not a possible option in order to maintain a substantial number of information.

To replace NaN values in numerical features with plausible values, a **CustomKNNImputer** with **different values of** k\_neighbours was used. The imputing was done also by dividing patients **by label**, in order to create new values only considering neighbours of the same class (this can lead to more separate classes and better classification. Indeed by doing this we assure that imputed values are **drawn from the same distribution of the other instances in the class**, reducing the risk of **bias** introduced by imputing based on the majority class).

For categorical features, in order to replace NaN values with plausible values, a **CustomSimpleImputer** with "mode" (reported in python as  $most\_frequent$ ) was used. Also in this case, patients where divided by label.

Before deciding for the best **scaler**, it's important to analyze the **boxplots** (and so the distribution) of our numerical features one by one:

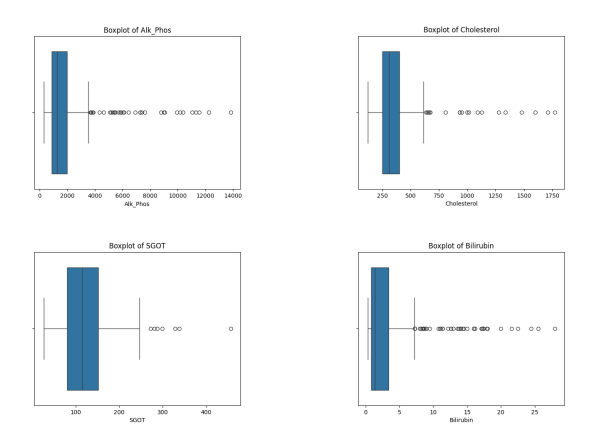


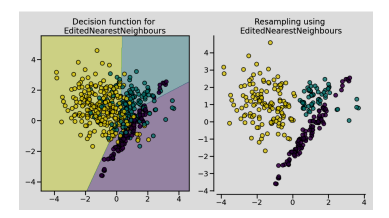
Figure 3: Boxplots of some numerical features as examples.

Given the relevant presence of outliers, the best choice is to apply a RobustScaler.

For the feature reduction, **normalized mutual information** was used, applied to a **SelectKBest** strategy with various values of K.

To balance instances, a double balancing technique was used. The undersampling technique was the **EditedNearestNeighbour**, with sampling strategy to decrease the number of instances in the "High chance" class.

This method was used since it can not only decrease the number of instances, but it's also capable of **separating** classes in the process, by deleting instances that may be close while being in different classes.

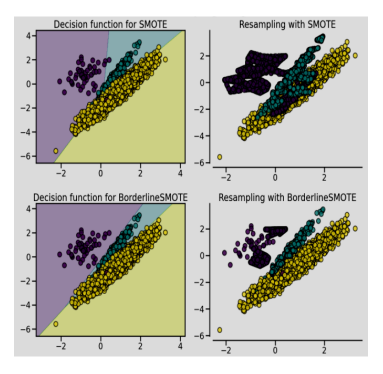


This was done with the goal of deleting instances of the High chance class that may make more difficult to classify the others.

The only problem with this approach is that there is no way to control **how many instances** are **deleted** (due to how it's implemented), differently from the other approach presented for the oversampling.

Oversampling was done in order to increase the number of instances in the Transplant needed class. The technique utilized was **SMOTE-ENC** [3](SMOTE for numerical and categorical) approach for a pipeline, where after the encoding there is the option to over sample with SMOTE or **BorderlineS-MOTE**, that can increase the chance of classifying instances which are considered hard to classify due to their position (close to the border of other classes).

SMOTE-ENC, differently from SMOTEN and SMOTE-NC, can handle continuous and categorical **simultaneously**, while SMOTEN and SMOTE-NC cannot.



All the methods were also used with different values of n-neighbours.

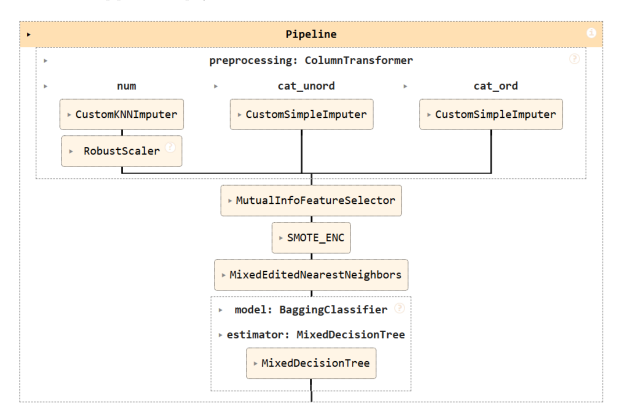
Overall, the goal of the whole sampling procedure was to decrease the number of instances of the majority class down to the closest class in terms of instances, then increasing the lowest up to the closest.

e.g.) Suppose:

- Starting labels: Low chance = 100, High chance = 200 and Transplant needed = 20;
- Final result: 100 for all the classes.

This decision was driven by the fact that only applying oversampling could have caused **over fitting**, since creating to much synthetic values may lead the model to learn the **noise in the data rather than the underlying patterns**.

The overall pipeline is displayed below.



#### 5 Models Definition and Evaluation

In order to conduct the experiments, three different models where used:

- Decision Tree with Bagging;
- KNNeighbour;
- AdaBoost with naive Bayes classifier.

All the model parameters where optimized using the Randomized Cross Validation method.

Also it was important to decide the scorer in order to decide the best classifier, and the decision was to choose the classifier that improved the  $f1\_macro\_average$  score. This was done to **penalize** classifiers that did not achieve good performances in classifying the Transplant needed class, which is actually the hardest given the very low number of instances, while also being crucial since a liver transplant is a very complex procedure, it is highly regulated, and only performed at designated transplant medical centers by highly trained transplant physicians and supporting medical team.

The hardest decision was made when choosing **how many folds** where needed in order to obtain a significant amount of information to perform statistical analysis (that could be considered **valid**) on the results. Indeed most of the tests may not be **efficient** when using few values, but on the contrary, a confusion matrix with not so many instances may be **useless to compare performances**.

The chosen method was the **Kruskall-Vallis** test (combined with the **Wilcoxon** test if statistically relevant differences were found between groups), which is a **non-parametric** test, that can be used to compare three or more independent distributions, and it has as an assumption, as mentioned in the sklearn page, to have **5 or more values**, together with independence and equal variances (in order to avoid staying "at the border" of the assumption, **10 folds** were selected).

#### 5.1 Validation of the Assumptions

Both the test mentioned before were used since the Q-Q plot Figure 4 of the scores showed that the distributions are **not normally distributed**, so non-parametric tests were selected.

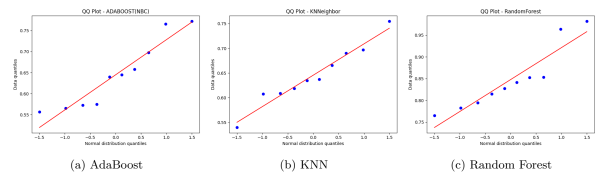


Figure 4: Q-Q plots for the different models.

After seeing this, the equal variances hypothesis can be proved by the Levene's test:

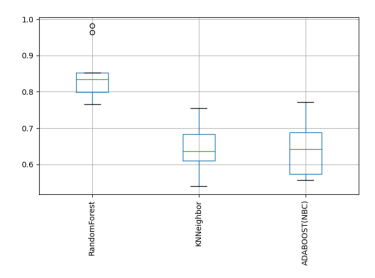
```
Levene's statistic: 0.46629856844442724 p value: 0.6322779360193294 There is no evidence to reject the null hypothesis of equal variances.
```

#### 6 Final Results

The results of the pipelines showed that overall the RandomForest approach **performed better** (all the results in the appendix).

The following are the best parameters for both the pipelines:

```
{
    'undersampler_k_neighbors': 9,
    'preprocessing_num_imputer_n_neighbors': 8,
    'oversampler_k_neighbors': 5,
    'oversampler_borderline': False,
    'model_estimator_max_depth': 5,
    'model_estimator_criterion': 'gini',
    'feature_selector_k': 12
}
```



# 7 Appendix

Kruskal-Wallis statistic: 18.59870967741935

p value: 9.148323388430848e-05

There is a statistically relevant difference between at least two groups.

Comparing RandomForest and KNNeighbor

Wilcoxon statistic: 0.0 p value: 0.001953125

There is a statistically relevant difference between RandomForest and KNNeighbor

----

Comparing RandomForest and ADABOOST(NBC)

Wilcoxon statistic: 0.0 p value: 0.001953125

There is a statistically relevant difference between RandomForest and ADABOOST(NBC)

\_\_\_\_

Comparing KNNeighbor and ADABOOST(NBC)

Wilcoxon statistic: 24.0 p value: 0.76953125

There is no statistically relevant difference between KNNeighbor and ADABOOST(NBC)

----

### 8 References

- 1. Medical News Today, "Cirrhosis of the liver: Life expectancy," https://www.medicalnewstoday.com/articles/cirrhosis-of-the-liver-life-expectancy
- 2. Cirrhosis Prediction Dataset https://www.kaggle.com/datasets/fedesoriano/cirrhosis-prediction-dataset
- 3. SMOTE-ENC algorithm https://arxiv.org/abs/2103.07612