# APA- Practical Work 2017-2018

## Albert Ribes

## Kerstin Winter

## January 16, 2018

## **Contents**

1	1.1 1.2 1.3	Desciption Desciption of the work and its goals Desciption of available data Instructions for running the code 1.3.1 Needed packages
2	Rela	ted Previous Work
3	3.1 3.2 3.3	Pre-processing 3.1.1 Treatment of missing values 3.1.2 Treatment of anomalous values 3.1.3 Treatment of incoherent values 3.1.4 Coding of non-continuous or non-ordered variables 3.1.5 Possible elimination of irrelevant variables 3.1.6 Creation of new useful variables (Feature extraction) 3.1.7 Normalization of the variables 3.1.8 Transformation of the variables Clustering Visualization Renaming of the atritutes
4	Res	ampling protocol
5	Metr	ric used to evaluate the models
6	6.1	Naive Bayes
7	7.1	ults obtained using non-linear methods Random Forest

	8.1 Estimation of the generalization error	8
9	Self-assessment of successes, failures and doubts	8
10	Scientific and personal conclusions	8
11	Possible extensions and known limitations	8

## **Todo list**

Tener todos los datasets listos para revista, preparados para poder meterlos en un modelo

## 1 Introduction

## 1.1 Desciption of the work and its goals

The goal of this project is to build a classification model to predict whether a lung cancer patient will die within one year after surgery or not. To do so we will study a dataset with real lung cancer patients.

As this is very sensitive information, our priority will be to minimize the amount of false negatives, i.e., avoid predicting a patient will not die within one year when it certainly does.

The data is taken from  $\verb|https://archive.ics.uci.edu/ml/datasets/Thoracic+Surgery Data\# [zieba2013boosted]$ 

## 1.2 Desciption of available data

The data we are working with is about patients who underwent major lung resections for primary lung cancer in the years from 2007 to 2011. For each patient we are given information about his diagnosis and effects produced by the cancer.

The dataset is very limited in the number of instances available: it only has 470. In addition, the distribution of the predicted class isn't quite balanced, since only 70 of the patients died in one year period. This may become a problem in some of the prediction models due to the fact that the results will be biased towards the biggest class. However, we can suppose that the data has been collected uniformly and that this proportion is similar to the real one.

For each patient we have 16 different atributes. 3 of them are numerical, and the rest are categorical. From those, 10 are binary. The response atribute is also binary.

## 1.3 Instructions for running the code

- Cual debe ser el working directory
- La variable que contiene el path que tiene que modificar
- Los resultados ya están calculados en un working space, para ahorrar tiempo.

#### 1.3.1 Needed packages

## 2 Related Previous Work

Nuestros datos son complicados, puesto que están muy balanceados y son muy pocos. El paper que referenciamos enfrente este tipo de problemas y propone usar SVM. Comentar que también usan este dataset.

Boosted SVM for extracting rules from imbalanced data in application to prediction of the post-operative life expectancy in the lung cancer patients

## 3 Data exploration process

## 3.1 Pre-processing

#### 3.1.1 Treatment of missing values

Our dataset do not have missing values, so there is no need to treat them.

#### 3.1.2 Treatment of anomalous values

The age of the patients is not well distributed. Most of them are over 60 years old, and only 4 of the patients are under 40. Due to this, it is very likely that the conclusions of our study will only be applicable to the elder people. However, we are reluctant to remove the younger patients.

#### 3.1.3 Treatment of incoherent values

The variable FEV1 which is the Forced Expiratory Volume in 1 second, shows a few anomalously high values. Depending on factors like age and sex of a patient the average value of the FEV1 should be around 3-6 litres, whereas the dataset shows values up to 86. As most of the values are within 0 and 10 we assume that the dataset contains the FEV1 in litres. To decide which values are to be determined as outliers we calculate the FEV1/FVC ratio which gives the percentage of the lung volume exhaled in the first second over the whole exhaled volume. All patients having a unrealistic ratio higher than 100%, which are 22 patients, are determined to be outliers and eliminated. We chose not to apply any stricter constraints because the dataset does not include the sex of the patients which influences the normal values of the FEV1 much.

Source of knowledge about FEV1 and FVC: https://www.nuvoair.com/blogs/blog/do-you-know-how-to-interpret-the-results-of-your-spirometry-test

El enlace antiguo ha caido, ahora es este:

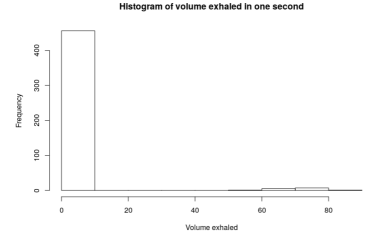


Figure 1: Show the ammount of people having each value

#### 3.1.4 Coding of non-continuous or non-ordered variables

As most of the dataset variables are logical, we have coded them as logical in R. We have converted the variable "DGN" to many binary variables, each one saying wether the patient showed that diagnosis or not.

Originally, the variables "PERFORMANCE" and "SIZE" were categorical, but as they seem to have some kind of order, we have coded them as numeric. The "PERFORMANCE" can have values 1,2,3 and the variable "SIZE" can have values 1,2,3,4. Both of them are then normalized.

"AGE" is also normalized in the range [0, 100]

#### 3.1.5 Possible elimination of irrelevant variables

Some of the variables of our dataset are not well represented. In particular:

DGN	There is just one patient with DGN = 1 and just 8 have DGN = 8
PAD	Just 8 patients have PAD = True
ASHTMA	Just 2 patients have ASHTMA = True
MI	Just 2 patients have MI = True

As we have very few instances we will train and run the models on two different datasets. One of them, thoraric.original, will contain all of the original attributes, and the other, thoraric.removed, will contain all attributes but "DGN.1", "DGN.8", "PAD", "ASHTMA" and "MI".

#### 3.1.6 Creation of new useful variables (Feature extraction)

We have not created any new variable as most of the ones we have are logical and it doesn't seem to be any relation among them.

#### 3.1.7 Normalization of the variables

We need to normalize only our numeric variables, which are the AGE, FVC and FEV1. As we have converted the variables "PERFORMANCE" and "SIZE" to numeric (as we will see in section 3.1.8) we also need to normalize them. To normalize the age we will only consider cases between 0 and 100 years old. For FVC and FEV1 the range will correspond to the maximum and minimum observed values with a margin of 10%.

#### 3.1.8 Transformation of the variables

Acording to the paper we found the accetable range for skewness in a numeric variable is (-2, +2). In the original dataset, the one which contained all the patients, the skewness of variables AGE, FVC and FEV were out of this range. But after eliminating 22 of the patients (in 3.1.3) all of them are inside the acceptable range, so there's no need to transform them.

## 3.2 Clustering

Quizá quitamos este apartado. Si nos da la vida, hacer muchos kmeans, cojer los mejores y cometnar alguna cosa

### 3.3 Visualization

Como nuestros datos son mixtos, es complicado encontrar un algoritmo de reducción de dimensión fiable que los pueda soportar. Buscamos alguna tabla un poco relevante que de una visión rápida de los datos, y ya

## 3.4 Renaming of the atritutes

Poner alguna tabla en la que se indique el cambio de nombres que se ha hecho desde los datos originales y los que nosotros hemos tocado

## 4 Resampling protocol

After the pre-processing we have 2 datasets with 448 instances. *thoraric.original* has 23 attributes and *thoraric.removed* has 19.~~0.85% of the patients have the target variable (DIED) equals FALSE.

The following lines explain the resampling protocol that we have used. All the numbers used can easily be tuned in the scripts provided.

To test our models we split our data into two different datasets, one for training and one for testing. The testing dataset will contain  $\frac{1}{3}$  of all the patients. As they are chosen

randomly, it is expected that the proportion of patients in each of the classes is kept. We will only use the testing dataset in the end, to test our models.

In the training dataset we have 299 patients, and it is expected that over 254 of them will be DIED = FALSE and 45 will be DIED = TRUE. As most of the models are sensitive to non-balanced datasets, we need to do something to balance our training dataset. We use the *bagging* algorithm to generate 51 balanced datasets, and each one of them will be used to train 51 models of the same type. That number could be different, but it is good for it to be odd, to avoid ties. Then, to generate a prediction, each of the models will be used to make a *hard* vote, and the class predicted by the majority will be the answer. This way we have constructed what we have called a *super-model* built with simpler models.

Each of the *bags* will contain the same amount of instances. As we want them to be balanced, all of them will contain every TRUE instance in the training dataset, and a random sample of the same size of the FALSE instances. Thus, each bag will have over  $2\times45=90$  instances.

## 5 Metric used to evaluate the models

Explicar porque usamos esta métrica, qué queremos incentivar, etc

## 6 Results obtained using linear/quadratic methods

## 6.1 Naive Bayes

- Puede ser adecuado porque admite tipos de datos distintos sin ningún problema
- No necesita ningún hiperparámetro
- Puede ser más sensible a haber quitado algunos atributos, pues afecta a la suposición Naive de independencia
- Mostrar la confussion matrix
- Comentar si muestra buenos o malos resutlados
- Comentar diferencias entre cada uno de los dos dataset, y explicar porq qué
- Identificar por qué suponemos que los resultados son buenos o malos.

### 6.2 KNN

- Por qué decidimos usar distancia manhattan en vez de euclidea
- Ver, en promedio, cuantos vecinos se han considerado

- Comentar que nosotros ya hemos normalizado los datos y que hemos puesto de forma adecuada las variables categóricas
- Qué podría afectar a este modelo y cómo nos afectará a nosotros.
- Mostrar los resultado y la confussion matrix
- Comentar los resultados
- Comentar las diferencias entre cada uno de los dos datasets
- Comentar por qué creemos que los resultados han sido buenos o malos

#### 6.3 General Linear Model

- Los datos se considera todos como reales, y no entiende de booleanos o categóricas
- No hemos usado funciones de base especiales, solamente la identidad con cada uno de los atributos.
- Ver en promedio cual ha sido la regularización
- Qué puede afectar a este modelo y cómo nos va a afectar a nosotros
- Mostrar los resultados y la confussion matrix
- Comentar los resultados
- Cometnar las diferencias entre cada uno de los datasets
- Comentar por qué creemos que los resultados han sido buenos o malos

## 7 Results obtained using non-linear methods

### 7.1 Random Forest

- Puesto que random forest ya hace algún tipo de resampling, quizá era mejor probarlo con el dataset preprocesado original, y no con los 51 bags que hemos creado.
   Comentar por qué hemos hecho esto y lo que podría pasar
- Comentar los tipos de bosques que han quedado después del crossvalidation: la cantidad de árboles que tiene cada uno, el mínimo de instancias en cada nodo, la cantidad de atributos que se consideran, etc.
- Qué puede afectar a este modelo y cómo nos va a afectar a nosotros
- Mostrar los resultados y la confussion matrix

- Comentar los resultados
- Comentar las diferencias entre cada uno de los dos datasets
- Comentar por qué creemos que los resultados han sido buenos o malos

#### 7.2 Neural Network

- Ver en promedio cuantas neuronas había en cada modelo y cuanta regularización se ha usado, después de los datos de crossvalidation
- Indicar que no se han hecho skips, y que únicamente hay una capa oculta...
- No hace ningún tipo de distinción entre los tipos de datos de entrada. Para él todo son reales, los booleanos también
- Qué puede afectar a este modelo y cómo nos puede afectar a nosotros
- Mostrar los resultados y la confussion matrix
- Comentar los resultados
- Comentar las diferencias entre cada uno de los dos datasets
- Comentar por qué creemos que los resultados han sido buenos o malos
- 8 Desciption and justification of the final model chosen
- 8.1 Estimation of the generalization error
- 9 Self-assessment of successes, failures and doubts
- 10 Scientific and personal conclusions
- 11 Possible extensions and known limitations