Liver Disease Classification

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1 Introduction

This report is part of the 'HarvardX: PH125.9x Data Science: Capstone' course. In this report, we chose a dataset of our choice and apply various machine learning techniques to perform binary classification to diagnose liver disease.

1.1 Background

The liver plays an important role in keeping us healthy. The main job of liver is to filter the blood coming from the digestive tract, before passing it to the rest of the body. The liver also turns nutrients into chemicals our body needs, turns food into energy, and filters out poisons. So, malfunctioning of liver affects the whole body.

The classification techniques are used in various automatic medical diagnoses tools[1]. The problems with liver patients are not easily discovered in an early stage. An early diagnosis of liver problems will help in increasing the survival rate of patiets. We can detect the liver disease by analyzing the levels of enzymes in the blood [2, 3]. A classification algorithm capable of automatically detecting the liver disease can assisst the doctors.

1.2 Aim of Project

The patients with liver disease are on the rise because of excessive consumption of alcohol, inhale of harmful gases, intake of contaminated food, pickles and drugs. The aim of this proect is to develop a bianry classifier, which can use blood enzymes information to diagnose liver disease.

2 Dataset and Evaluation Metrics

We use the Liver Patient Records, which are collected from North East of Andhra Pradesh, India. The data set contains:

1. 416 liver patient records and 167 non-liver patient records.

2.1 Download Data

The dataset is publically available online both at Kaggle and UCI repository. We download data from the website. Then, we split data into a training and validation sets.

• 10% of the data is used for validation, and 90

```
# Install packages (if not installed)
####################################
# Note: this process could take a couple of minutes
repos_path<- "http://cran.us.r-project.org"</pre>
if(!require(tidyverse)) install.packages("tidyverse", repos =repos_path)
## Loading required package: tidyverse
## -- Attaching packages ------
## v ggplot2 3.2.1
                  v purrr
                            0.3.3
## v tibble 2.1.3 v dplyr
                            0.8.3
         1.0.0
## v tidyr
                   v stringr 1.4.0
## v readr
         1.3.1
                   v forcats 0.4.0
## -- Conflicts ------ tid
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                 masks stats::lag()
if(!require(caret)) install.packages("caret", repos = repos_path)
## Loading required package: caret
## Warning: package 'caret' was built under R version 3.6.3
## Loading required package: lattice
##
## Attaching package: 'caret'
## The following object is masked from 'package:purrr':
##
##
     lift
```

```
if(!require(data.table)) install.packages("data.table", repos =repos_path)
## Loading required package: data.table
## Warning: package 'data.table' was built under R version 3.6.3
##
## Attaching package: 'data.table'
## The following objects are masked from 'package:dplyr':
##
##
       between, first, last
## The following object is masked from 'package:purrr':
##
##
       transpose
if(!require(lubridate)) install.packages("lubridate", repos = repos_path)
## Loading required package: lubridate
##
## Attaching package: 'lubridate'
## The following objects are masked from 'package:data.table':
##
##
       hour, isoweek, mday, minute, month, quarter, second, wday,
##
       week, yday, year
## The following object is masked from 'package:base':
##
##
       date
if(!require(dplyr)) install.packages("dplyr", repos = repos_path)
if(!require(sjmisc)) install.packages("dplyr", repos = repos_path)
## Loading required package: sjmisc
## Warning: package 'sjmisc' was built under R version 3.6.3
## Attaching package: 'sjmisc'
## The following object is masked from 'package:purrr':
##
##
       is_empty
## The following object is masked from 'package:tidyr':
##
##
       replace_na
```

```
## The following object is masked from 'package:tibble':
##
       add_case
##
if(!require(sjmisc)) install.packages("scales", repos = repos_path)
####################################
# Load libraries
####################################
library(lubridate)
library(tidyverse)
library(dplyr)
library(lubridate)
library(sjmisc)
library(scales)
##
## Attaching package: 'scales'
## The following object is masked from 'package:purrr':
##
##
       discard
## The following object is masked from 'package:readr':
##
##
       col_factor
####################################
# Downloading data
####################################
# Indian Live Patient Records :
 # https://www.kaggle.com/uciml/indian-liver-patient-records/
 # https://archive.ics.uci.edu/ml/machine-learning-databases/00225/Indian Liver Patient Dataset (ILPD).
url <- "https://archive.ics.uci.edu/ml/machine-learning-databases/00225/Indian Liver Patient Dataset (I
# Download csv
liverData <- read.csv(url)</pre>
# Rename columns of csv
colnames(liverData) <- c("Age", "Gender", "Total_Bilirubin", "Direct_Bilirubin", "Alkaline_Phosphotase", "Al</pre>
###################################
# Creating training and validation sets
####################################
# Validation set will be 10% of whole data
set.seed(1, sample.kind = "Rounding")
## Warning in set.seed(1, sample.kind = "Rounding"): non-uniform 'Rounding'
## sampler used
```

```
test_index <- createDataPartition(y = liverData$Dataset, times = 1, p = 0.1, list = FALSE)
training <- liverData[-test_index,]
validation <- liverData[test_index,]

# Removing the objects from environment as no longer required
rm(liverData)</pre>
```

2.2 Metrics

To evaluate the performance of classifiers, we will use following metrics:

1. Accuracy It is the ratio of number of correct predictions to the total number of input samples.

$$Accuracy = \frac{True positives + True negatives}{Total Predictions} \tag{1}$$

2. **Sensitivity** It is also referred as true positive rate or recall. It is the proportion of true positives that are correctly identified.

$$Sensitivity = \frac{Number of true positives}{Number of true positives + Number of false negatives} \tag{2}$$

3. Precision It is defined as the proportion of the true positives against all the positive results.

$$Precision = \frac{Number of true positives}{Number of true positives + Number of false positives} \tag{3}$$

4. **Specificity** It is the True negative rate. It is the proportion of true negatives that are correctly identified.

$$Specificity = \frac{Number of true negatives}{Number of true negatives + Number of false positives} \tag{4}$$

5. **F1 Score** One metric that is preferred over overall accuracy is the average of specificity and sensitivity, referred to as balanced accuracy. Because specificity and sensitivity are rates, it is more appropriate to compute the harmonic average. In fact, the F1-score is widely used to compute harmonic average of precision and recall.

$$F1Score = 2 * \frac{Precision - Recall}{Precision + Recall}$$
 (5)

3 Data Exploration

The dataset contains 11 variables namely, 'Age', 'Gender', 'Total_Bilirubin', or "Alkaline_Phosphotase". The 'Dataset' variable indicates if the liver has a disease or not. For instance, a value of 1 indicates a disease and 2 indicates no disease.

All other variables except Age", Gender", and "Dataset" represent the amount of enzymes in the blood. These variables will be used to train our machine learning models for making predictions or diagnosis.

head(training)

Age	Gender	${\bf Total_Bilirubin}$	${\bf Direct_Bilirubin}$	$Alkaline_Phosphotase$	$Alamine_Aminotransferase$	Aspartate_Aminotran
62	Male	10.9	5.5	699	64	
62	Male	7.3	4.1	490	60	
58	Male	1.0	0.4	182	14	
72	Male	3.9	2.0	195	27	
46	Male	1.8	0.7	208	19	
26	Female	0.9	0.2	154	16	

The training dataset has 523 records and there are no null values (confirmed using summary).

```
sprintf("Rows of training dataset = %d", nrow(training))
## [1] "Rows of training dataset = 523"
print("========")
```

[1] "======="

```
summary(training)
```

```
Gender
                                Total_Bilirubin Direct_Bilirubin
##
        Age
          : 4.00
##
   Min.
                   Female:125
                                Min. : 0.40
                                               Min. : 0.100
##
   1st Qu.:33.00
                   Male :398
                                1st Qu.: 0.80
                                                1st Qu.: 0.200
##
   Median :45.00
                                Median: 1.00
                                                Median : 0.300
##
   Mean
          :45.33
                                Mean : 3.22
                                                Mean : 1.446
                                3rd Qu.: 2.60
##
   3rd Qu.:58.00
                                                3rd Qu.: 1.300
##
   Max.
          :90.00
                                Max.
                                       :75.00
                                                Max.
                                                      :19.700
##
##
   Alkaline_Phosphotase Alamine_Aminotransferase Aspartate_Aminotransferase
                        Min. : 10.00
         : 63.0
##
   Min.
                                                 Min.
                                                       : 10.0
   1st Qu.: 176.0
                                  24.00
                                                 1st Qu.:
##
                        1st Qu.:
                                                           25.0
   Median : 208.0
                                  35.00
                                                 Median: 41.0
##
                        Median :
##
   Mean : 289.9
                        Mean :
                                  76.34
                                                 Mean : 105.0
   3rd Qu.: 298.0
                        3rd Qu.: 60.00
                                                 3rd Qu.: 86.5
##
          :1896.0
                               :1680.00
                                                       :4929.0
##
   Max.
                        Max.
                                                 Max.
##
##
   Total_Protiens
                     Albumin
                                  Albumin_and_Globulin_Ratio
                                                                Dataset
##
                         :0.900
   Min.
         :2.70
                                  Min.
                                         :0.3000
                                                                   :1.000
                  Min.
                                                             Min.
##
   1st Qu.:5.80
                  1st Qu.:2.600
                                  1st Qu.:0.7000
                                                             1st Qu.:1.000
##
   Median:6.60
                  Median :3.100
                                  Median :0.9300
                                                             Median :1.000
##
   Mean
         :6.49
                  Mean
                        :3.147
                                  Mean
                                        :0.9458
                                                             Mean :1.281
   3rd Qu.:7.20
                  3rd Qu.:3.800
                                  3rd Qu.:1.1000
                                                             3rd Qu.:2.000
          :9.50
##
   Max.
                  Max.
                         :5.500
                                  Max.
                                         :2.8000
                                                             Max.
                                                                   :2.000
##
                                  NA's
                                         :4
```

3.0.1 Data Wrangling

The variable "Dataset" is our prediction and we will use it to analyse the performance of machine lerning models. To improve readability, we create a new column namely "Liver Disease", which will have two values:

1. Malignant (M) indicating that the patient has a liver disease.

2. Benign (B) indicating that the patient has no no liver disease.

After creating a new column, wr delete the 'Dataset' variable. We apply these operations to both training and validation datasets.

```
# Adding a new column, which will contain the disease information
training <- transform(training, Disease= ifelse(Dataset==1, "M","B"))
validation <- transform(validation, Disease= ifelse(Dataset==1, "M","B"))

# Deleting the column 'Dataset' as no longer required
training<-within(training, rm(Dataset))
validation<-within(validation, rm(Dataset))

# Displaying the first siz rows
head(training)</pre>
```

Age	Gender	${\bf Total_Bilirubin}$	${\bf Direct_Bilirubin}$	$Alkaline_Phosphotase$	$Alamine_Aminotransferase$	Aspartate_Aminotran
62	Male	10.9	5.5	699	64	
62	Male	7.3	4.1	490	60	
58	Male	1.0	0.4	182	14	
72	Male	3.9	2.0	195	27	
46	Male	1.8	0.7	208	19	
26	Female	0.9	0.2	154	16	

References

- [1] Ethan Du-Crowa, Lucy Warrenb, Susan M Astleya and Johan Hullemanc,"Is there a safety-net effect with Computer-Aided Detection (CAD)?", Medical Imaging 2019.
- [2] Eugene, R., Sorrell, Michael F.; Maddrey, Willis C., "Schiff's Diseases of the Liver", 10th Edition, Lippincott Williams & Wilkins by Schiff.
- [3] Bendi, Venkata . R, M. S. Prasad Babu, and N. B. Venkateswarlu, "Critical Comparative Study of Liver Patients from USA and INDIA: An Exploratory Analysis", International Journal of Computer Science Issues, May 2012.