# first lesson

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#### Data

For this work we used the dataset Stroke Prediction Dataset retrieved from Kaggle.

We used the dataset to predict whether a patient is likely to get stroke based on the input parameters like gender, age, various diseases, and smoking status. Each row in the data provides relavant information about the patient.

Below the information about the attributes:

- 1. id: unique identifier
- 2. gender: "Male", "Female" or "Other"
- 3. age: age of the patient
- 4. hypertension: 0 if the patient doesn't have hypertension, 1 if the patient has hypertension
- 5. heart disease: 0 if the patient doesn't have any heart diseases, 1 if the patient has a heart disease
- 6. ever married: "No" or "Yes"
- 7. work\_type: "children", "Govt\_jov", "Never\_worked", "Private" or "Self-employed"
- 8. Residence\_type: "Rural" or "Urban"
- 9. avg\_glucose\_level: average glucose level in blood
- 10. bmi: body mass index
- 11. smoking\_status: "formerly smoked", "never smoked", "smokes" or "Unknown"\*
- 12. stroke: 1 if the patient had a stroke or 0 if not

#### Load the dataset

Set the current directory as working directory and load the dataset.

```
setwd(dirname(rstudioapi::getSourceEditorContext()$path))
source("../utils/plots.R")
source("../utils/data_analysis.R")
dataset = read.csv("stroke.csv")
str(dataset)
```

```
'data.frame':
                   5110 obs. of 12 variables:
                             9046 51676 31112 60182 1665 56669 53882 10434 27419 60491 ...
##
  $ id
                       : int
##
                              "Male" "Female" "Male" "Female" ...
   $ gender
                       : num 67 61 80 49 79 81 74 69 59 78 ...
##
  $ age
                             0 0 0 0 1 0 1 0 0 0 ...
   $ hypertension
                       : int
   $ heart_disease
                             1 0 1 0 0 0 1 0 0 0 ...
##
                       : int
                              "Yes" "Yes" "Yes" "Yes" ...
##
   $ ever_married
                       : chr
                              "Private" "Self-employed" "Private" "Private" ...
##
  $ work_type
                       : chr
   $ Residence_type
                              "Urban" "Rural" "Rural" "Urban" ...
                       : chr
```

<sup>\*</sup>Note: "Unknown" in smoking\_status means that the information is unavailable for this patient

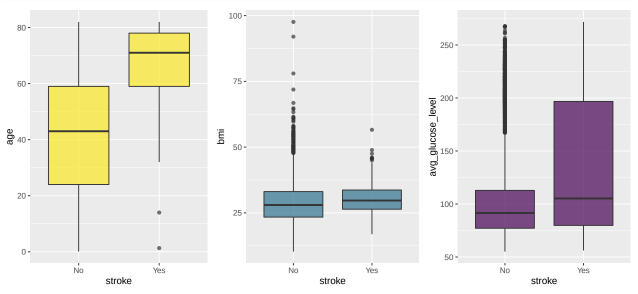
```
## $ avg_glucose_level: num 229 202 106 171 174 ...
## $ bmi : chr "36.6" "N/A" "32.5" "34.4" ...
## $ smoking_status : chr "formerly smoked" "never smoked" "never smoked" "smokes" ...
## $ stroke : int 1 1 1 1 1 1 1 1 1 1 ...
```

### Data Preparation

```
We converted the categorical variables in factors.
dataset$id = NULL
dataset$gender = factor(dataset$gender)
dataset$hypertension = factor(dataset$hypertension, levels = c(0,1), labels = c("No", "Yes"))
dataset$heart_disease = factor(dataset$heart_disease, levels = c(0,1), labels = c("No", "Yes"))
dataset$ever_married = factor(dataset$ever_married)
dataset$work_type = factor(dataset$work_type)
dataset$residence_type = factor(dataset$Residence_type)
dataset$Residence_type = NULL
dataset$smoking_status = factor(dataset$smoking_status)
dataset$stroke = factor(dataset$stroke, levels = c(0,1), labels = c("No", "Yes"))
dataset$bmi = as.numeric(dataset$bmi)
str(dataset)
                   5110 obs. of 11 variables:
## 'data.frame':
## $ gender
                       : Factor w/ 3 levels "Female", "Male", ...: 2 1 2 1 1 2 2 1 1 1 ...
                       : num 67 61 80 49 79 81 74 69 59 78 ...
## $ age
## $ hypertension
                      : Factor w/ 2 levels "No", "Yes": 1 1 1 1 2 1 2 1 1 1 ...
                      : Factor w/ 2 levels "No", "Yes": 2 1 2 1 1 1 2 1 1 1 ...
## $ heart_disease
                      : Factor w/ 2 levels "No", "Yes": 2 2 2 2 2 2 2 1 2 2 ...
## $ ever married
                       : Factor w/ 5 levels "children", "Govt_job",...: 4 5 4 4 5 4 4 4 4 ...
## $ work_type
## $ avg_glucose_level: num 229 202 106 171 174 ...
## $ bmi
                     : num 36.6 NA 32.5 34.4 24 29 27.4 22.8 NA 24.2 ...
## $ smoking status : Factor w/ 4 levels "formerly smoked",..: 1 2 2 3 2 1 2 2 4 4 ...
                       : Factor w/ 2 levels "No", "Yes": 2 2 2 2 2 2 2 2 2 ...
## $ stroke
## $ residence_type : Factor w/ 2 levels "Rural", "Urban": 2 1 1 2 1 2 1 2 1 2 ...
```

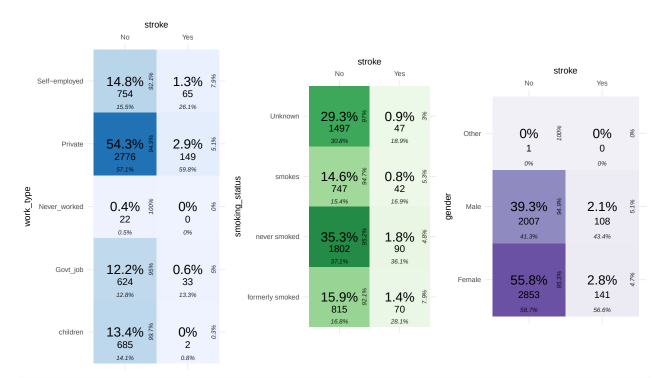
#### **Data Visualization**

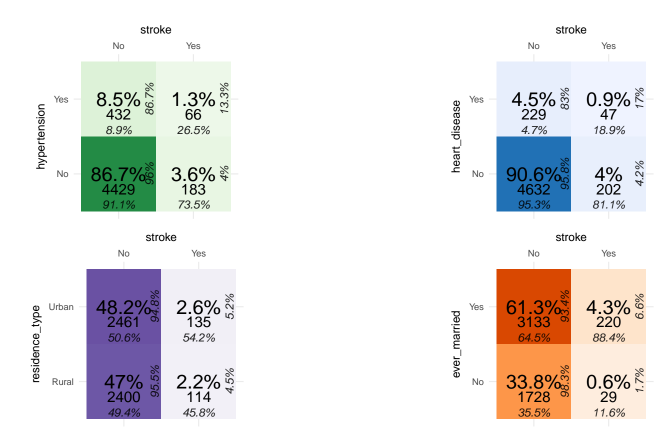
Boxplots to visualize the stroke status in association with age, bmi and glucose level.



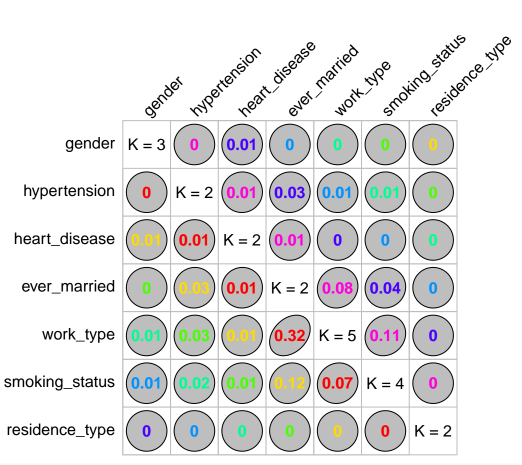
- 1. Age there is a relation with age: older people are more likely to have a stroke
- 2. Bmi there is no evident relation between stroke and bmi
- 3. Averagre glucose level the higher the level of glucose, the higher the relation with stroke

Then we built some matrices to show the relationship between stroke and some **categorical variables** that have different levels.

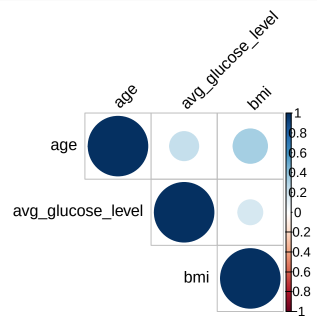




After we built some plots to show the qualitative and quantitative correlation.



```
# TODO find a way to plot table and image side-by-side
quantitative_vars = c('age', 'avg_glucose_level', 'bmi')
corr <- rcorr(as.matrix(dataset %>% select(all_of(quantitative_vars))))
corrplot(corr$r, type = "upper", tl.col = "black", tl.srt = 45)
```



#flattenCorrMatrix(corr\$r, corr\$P)

### **Data Manipulation**

In order to facilitate our work, we removed the level **other** from the variable gender and the level **never\_worked** from the variable work\_type, since they are not relevant for the analysis.

```
# rm gender "other"
dataset$gender <- as.character(dataset$gender)
stroke <- dataset[dataset$gender == "Male" | dataset$gender == "Female",]
stroke$gender <- as.factor(stroke$gender)

# rm "Never_worked"
dataset$work_type <- as.character(dataset$work_type)
stroke <- dataset[!dataset$work_type == "Never_worked",]
stroke$work_type <- as.factor(stroke$work_type)</pre>
```

We removed the NA values from the **bmi** variable. In order to replace them, we predicted their values using a fitted tree

```
a fitted tree.
stroke <- as tibble(stroke)</pre>
sum(is.na(stroke$bmi))
## [1] 201
missing_index <- which(is.na(stroke$bmi))</pre>
X <- stroke[missing index,]</pre>
train_v <- stroke[-c(missing_index),]</pre>
tree = caret::train(bmi ~ .,
                     data=train_v,
                     method="rpart",
                     trControl = trainControl(method = "cv"))
bmi_pred <- predict(tree, newdata = X)</pre>
stroke[missing_index,"bmi"] <- bmi_pred</pre>
sum(is.na(stroke$bmi))
## [1] 0
sum(is.na(stroke))
## [1] 0
# check for other NAs
for (i in 1:11) {
  print(which(is.na(stroke[,i])))
}
## integer(0)
```

```
# clean Glob_Env
rm(train_v,X,tree, bmi_pred,i,missing_index)
```

# Divide in training and test set

```
set.seed(42)
stroke <- as.data.frame(stroke)
for (i in 1:11) {
   levels(stroke[,i]) <- make.names(c(levels(stroke[,i])))
}

split_train_test <- createDataPartition(y = stroke$stroke, p=0.5, list = F)
train <- stroke[split_train_test,]
test <- stroke[-split_train_test,]</pre>
```

### SMOTE algorithm

Since our dataset has a problem of under sampling, we used the SMOTE algorithm to create synthetic new data.

```
# new smote algo
```

### Logistic Regression

## smoking\_statussmokes
## smoking\_statusUnknown

```
Logit <- glm(stroke~., data=train, family = binomial(link = "logit"))</pre>
summary(Logit)
##
## Call:
## glm(formula = stroke ~ ., family = binomial(link = "logit"),
##
       data = train)
##
## Deviance Residuals:
                  1Q
                        Median
                                      3Q
                                                Max
## -1.24055 -0.32175 -0.16709 -0.08114
                                            3.03598
## Coefficients:
                               Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                             -2.039e+01 5.667e+02 -0.036
                                                             0.9713
## genderMale
                              2.056e-01 1.993e-01 1.031
                                                             0.3023
## genderOther
                             -1.452e+01 1.075e+04 -0.001
                                                             0.9989
## age
                              7.149e-02 8.197e-03
                                                     8.721
                                                             <2e-16 ***
                              5.664e-01 2.223e-01
                                                     2.548
## hypertensionYes
                                                             0.0108 *
## heart_diseaseYes
                              3.453e-01 2.671e-01 1.293
                                                             0.1960
                             -1.706e-01 3.134e-01 -0.544
## ever_marriedYes
                                                             0.5863
## work_typeGovt_job
                              1.236e+01 5.667e+02
                                                     0.022
                                                             0.9826
## work_typePrivate
                              1.286e+01 5.667e+02
                                                     0.023
                                                             0.9819
                              1.251e+01 5.667e+02
## work_typeSelf.employed
                                                     0.022
                                                             0.9824
## avg_glucose_level
                              2.993e-03 1.706e-03
                                                     1.755
                                                             0.0793
## bmi
                              8.237e-03 1.596e-02
                                                     0.516
                                                             0.6058
```

## smoking\_statusnever.smoked -1.453e-01 2.462e-01 -0.590

-2.820e-01 3.120e-01 -0.904

3.513e-01 2.904e-01

0.5550

0.2265

0.3660

1.210

As we can see, the only significant variables are **age** and **hypertension**, that have both an high positive impact on the increase of the logit probability.

Also about the **smoking status** we can see, as expected, that it has a positive impact and a negatie one when the individual has never smoked.

```
# TODO clean code and stuff
# data Preprocessing, Econding with OneHotEncoding ------
#dummy <- dummyVars(" ~ gender + work_type + smoking_status + ever_married + Residence_type", data=stro
#newdata <- data.frame(predict(dummy, newdata = stroke))</pre>
#a <- stroke[,2:4]
#b <- stroke[,8:9]
#dt <- cbind(a, b, newdata, stroke['stroke'])
\#dt \leftarrow as \ tibble(dt)
#y <- stroke['stroke']</pre>
# Fill missing bmi data w/ tree prediction
# TODO keep this way? should we just remove them?
#
# missing_index <-which(is.na(dataset$bmi))</pre>
# train_set <- dataset[-c(missing_index),]</pre>
#
# tree = caret::train(bmi ~ .,
#
                      data=train_set,
#
                      method="rpart",
#
                      trControl = trainControl(method = "cv"))
# predicted_bmi <- predict(tree, newdata = dataset[missing_index,])</pre>
# # What? Why?
# # x <- mean(bmi_pred)
# # bmi pred[202] <- x
# #########################
# dataset[missing_index,"bmi"] <- predicted_bmi</pre>
# dataset = na.omit(dataset)
# # Check quantitative correlation is under control w/ new data
# TODO side by side :)
```

```
# new_corr <- rcorr(as.matrix(dataset %>% select(all_of(quantitative_vars))))
# flattenCorrMatrix(new corr$r, new corr$P)
# grid.arrange(corrplot(corr$r,
                                  type = "upper", tl.col = "black", tl.srt = 45),
               corrplot(new_corr$r, type = "upper", tl.col = "black", tl.srt = 45),
#
              ncol=2, nrow=1)
# Solve the under sampling problem with SMOTE algho to create synth new data
# dataset <- as.data.frame(lapply(dataset, as.factor)) # What? Why?</pre>
# trainSplit <- DMwR::SMOTE(stroke ~ ., dt, perc.over = 2000, perc.under=10)</pre>
# dt_synth<- rbind(trainSplit,dt)</pre>
# dt_synth$work_type.Never_worked <- NULL # Again, but why :D</pre>
# dt_synth$avg_glucose_level <- as.numeric(dt_synth$avg_glucose_level)
# dt_synth$bmi <- as.numeric(dt_synth$bmi)</pre>
# dt_synth$age <- as.numeric(dt_synth$age)</pre>
# TODO Show some stats of new dataset
# TODO recheck correlations, maybe w/ function to plot both side-by-side
# TODO add models/predictions
```