

# Finalcode \_rproject

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*code :-*

```
library(ggplot2)
library(tidyverse)
```

```
## -- Attaching packages ----- tidyverse 1.3.0 --
```

```
## v tibble  3.0.2    v dplyr   1.0.0
## v tidyr   1.1.0    v stringr 1.4.0
## v readr   1.3.1    v forcats 0.5.0
## v purrr   0.3.4
```

```
## -- Conflicts ----- tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()     masks stats::lag()
```

```
library(dplyr)
library(caret)
```

```
## Loading required package: lattice
```

```
##
```

```
## Attaching package: 'caret'
```

```
## The following object is masked from 'package:purrr':
```

```
##
```

```
## lift
```

```
library(mlbench)
library(tidyr)
library(Boruta)
library(data.table)
```

```
##
```

```
## Attaching package: 'data.table'
```

```
## The following objects are masked from 'package:dplyr':
```

```
##
```

```
## between, first, last
```

```

## The following object is masked from 'package:purrr':
##
##      transpose

library(modelr)
library(cowplot)

##
## *****

## Note: As of version 1.0.0, cowplot does not change the

##      default ggplot2 theme anymore. To recover the previous

##      behavior, execute:
##      theme_set(theme_cowplot())

## *****

library(randomForest)

## randomForest 4.6-14

## Type rfNews() to see new features/changes/bug fixes.

##
## Attaching package: 'randomForest'

## The following object is masked from 'package:dplyr':
##
##      combine

## The following object is masked from 'package:ggplot2':
##
##      margin

library(ROSE)

## Loaded ROSE 0.0-3

library(e1071)
library(Matrix)

##
## Attaching package: 'Matrix'

## The following objects are masked from 'package:tidyr':
##
##      expand, pack, unpack

```

```
library(xgboost)
```

```
##  
## Attaching package: 'xgboost'  
  
## The following object is masked from 'package:dplyr':  
##  
## slice
```

```
library(mlr)
```

```
## Loading required package: ParamHelpers  
  
## 'mlr' is in maintenance mode since July 2019. Future development  
## efforts will go into its successor 'mlr3' (<https://mlr3.ml-org.com>).  
  
##  
## Attaching package: 'mlr'  
  
## The following object is masked from 'package:e1071':  
##  
## impute  
  
## The following object is masked from 'package:modelr':  
##  
## resample  
  
## The following object is masked from 'package:caret':  
##  
## train
```

```
library(MLeval)
```

```
# Reading the demographics, laboratory and examination .csv files using read_csv
```

```
demo <- read_csv("C:/Semester 2/Intro to Data Mining and Processing/Project/national-health-and-nutrition-survey-2017.csv")
```

```
## Parsed with column specification:  
## cols(  
##   .default = col_double()  
## )  
  
## See spec(...) for full column specifications.
```

```
labs <- read_csv("C:/Semester 2/Intro to Data Mining and Processing/Project/national-health-and-nutrition-survey-2017.csv")
```

```
## Parsed with column specification:  
## cols(  
##   .default = col_double()  
## )  
## See spec(...) for full column specifications.
```

```
exam <- read_csv("C:/Semester 2/Intro to Data Mining and Processing/Project/national-health-and-nutrition-survey-2017.csv")
```

```
## Parsed with column specification:
## cols(
##   .default = col_double(),
##   BMIHEAD = col_logical(),
##   OHX02CTC = col_character(),
##   OHX03CTC = col_character(),
##   OHX04CTC = col_character(),
##   OHX05CTC = col_character(),
##   OHX06CTC = col_character(),
##   OHX07CTC = col_character(),
##   OHX08CTC = col_character(),
##   OHX09CTC = col_character(),
##   OHX10CTC = col_character(),
##   OHX11CTC = col_character(),
##   OHX12CTC = col_character(),
##   OHX13CTC = col_character(),
##   OHX14CTC = col_character(),
##   OHX15CTC = col_character(),
##   OHX18CTC = col_character(),
##   OHX19CTC = col_character(),
##   OHX20CTC = col_character(),
##   OHX21CTC = col_character(),
##   OHX22CTC = col_character()
##   # ... with 38 more columns
## )
## See spec(...) for full column specifications.
```

```
# Making new dataframes consisting of only selected columns
demo_df <- demo %>% select(SEQN , "GENDER" = RIAGENDR, "AGE" = RIDAGEYR)

# HbA1c is Hemoglobin A1c or Glycohemoglobin which is an indicator of diabetes
lab_df <- labs %>% select(SEQN, "HbA1c" = LBXGH)

# Creating another dataframe by inner joining the other data frames
df <- lab_df %>% inner_join(demo_df, by = "SEQN") %>%
  inner_join(exam, by = "SEQN")

summary(df)
```

```
##      SEQN      HbA1c      GENDER      AGE
## Min.   :73557 Min.   : 3.500 Min.   :1.000 Min.   : 0.00
## 1st Qu.:76092 1st Qu.: 5.200 1st Qu.:1.000 1st Qu.:10.00
## Median :78643 Median : 5.400 Median :2.000 Median :27.00
## Mean   :78645 Mean   : 5.643 Mean   :1.508 Mean   :31.63
## 3rd Qu.:81191 3rd Qu.: 5.800 3rd Qu.:2.000 3rd Qu.:52.00
## Max.   :83731 Max.   :17.500 Max.   :2.000 Max.   :80.00
##      NA's      :3170
##      PEASCST1      PEASCTM1      PEASCCT1      BPXCHR
## Min.   :1.000 Min.   : 7.0 Min.   : 1.00 Min.   : 60.0
## 1st Qu.:1.000 1st Qu.: 578.0 1st Qu.: 56.00 1st Qu.: 88.0
```

##	Median :1.000	Median : 689.0	Median : 56.00	Median :102.0	
##	Mean :1.065	Mean : 660.6	Mean : 57.83	Mean :105.5	
##	3rd Qu.:1.000	3rd Qu.: 832.0	3rd Qu.: 72.00	3rd Qu.:120.0	
##	Max. :3.000	Max. :2868.0	Max. :122.00	Max. :178.0	
##		NA's :305	NA's :9493	NA's :7852	
##	BPAARM	BPACSZ	BPXPLS	BPXPULS	
##	Min. :1.000	Min. :1.000	Min. : 40.00	Min. :1.000	
##	1st Qu.:1.000	1st Qu.:3.000	1st Qu.: 66.00	1st Qu.:1.000	
##	Median :1.000	Median :4.000	Median : 74.00	Median :1.000	
##	Mean :1.008	Mean :3.675	Mean : 74.42	Mean :1.014	
##	3rd Qu.:1.000	3rd Qu.:4.000	3rd Qu.: 82.00	3rd Qu.:1.000	
##	Max. :8.000	Max. :5.000	Max. :180.00	Max. :2.000	
##	NA's :2278	NA's :2271	NA's :2264	NA's :302	
##	BPXPTY	BPXML1	BPXSY1	BPXDI1	
##	Min. :1.000	Min. :100.0	Min. : 66.0	Min. : 0.00	
##	1st Qu.:1.000	1st Qu.:130.0	1st Qu.:106.0	1st Qu.: 58.00	
##	Median :1.000	Median :140.0	Median :116.0	Median : 66.00	
##	Mean :1.004	Mean :144.7	Mean :118.1	Mean : 65.77	
##	3rd Qu.:1.000	3rd Qu.:150.0	3rd Qu.:128.0	3rd Qu.: 76.00	
##	Max. :2.000	Max. :888.0	Max. :228.0	Max. :122.00	
##	NA's :2249	NA's :2260	NA's :2641	NA's :2641	
##	BPAEN1	BPXSY2	BPXDI2	BPAEN2	
##	Min. :1.000	Min. : 66.0	Min. : 0.00	Min. :1.000	
##	1st Qu.:2.000	1st Qu.:106.0	1st Qu.: 58.00	1st Qu.:2.000	
##	Median :2.000	Median :116.0	Median : 66.00	Median :2.000	
##	Mean :1.989	Mean :118.2	Mean : 65.24	Mean :1.954	
##	3rd Qu.:2.000	3rd Qu.:128.0	3rd Qu.: 74.00	3rd Qu.:2.000	
##	Max. :2.000	Max. :230.0	Max. :116.00	Max. :2.000	
##	NA's :2274	NA's :2404	NA's :2404	NA's :2276	
##	BPXSY3	BPXDI3	BPAEN3	BPXSY4	
##	Min. : 62	Min. : 0.00	Min. :1.000	Min. : 80.0	
##	1st Qu.:106	1st Qu.: 58.00	1st Qu.:2.000	1st Qu.:108.0	
##	Median :114	Median : 68.00	Median :2.000	Median :126.0	
##	Mean :118	Mean : 65.04	Mean :1.963	Mean :125.7	
##	3rd Qu.:128	3rd Qu.: 74.00	3rd Qu.:2.000	3rd Qu.:140.0	
##	Max. :228	Max. :118.00	Max. :2.000	Max. :212.0	
##	NA's :2405	NA's :2405	NA's :2276	NA's :9298	
##	BPXDI4	BPAEN4	BMDSTATS	BMXWT	
##	Min. : 0.00	Min. :1.000	Min. :1.00	Min. : 3.10	
##	1st Qu.: 60.00	1st Qu.:2.000	1st Qu.:1.00	1st Qu.: 37.95	
##	Median : 70.00	Median :2.000	Median :1.00	Median : 65.30	
##	Mean : 69.01	Mean :1.879	Mean :1.14	Mean : 62.60	
##	3rd Qu.: 78.00	3rd Qu.:2.000	3rd Qu.:1.00	3rd Qu.: 83.50	
##	Max. :128.00	Max. :2.000	Max. :4.00	Max. :222.60	
##	NA's :9298	NA's :9251		NA's :90	
##	BMIWT	BMXRECUM	BMIRECUM	BMXHEAD	BMIHEAD
##	Min. :1.000	Min. : 48.60	Min. :1	Min. :33.80	Mode:logical
##	1st Qu.:3.000	1st Qu.: 69.70	1st Qu.:1	1st Qu.:39.70	NA's:9813
##	Median :3.000	Median : 82.80	Median :1	Median :41.80	
##	Mean :2.966	Mean : 81.63	Mean :1	Mean :41.57	
##	3rd Qu.:3.000	3rd Qu.: 93.20	3rd Qu.:1	3rd Qu.:43.50	
##	Max. :4.000	Max. :115.10	Max. :1	Max. :46.80	
##	NA's :9429	NA's :8748	NA's :9782	NA's :9584	
##	BMXHT	BMiht	BMXBMI	BMDBMIC	

##	Min. : 79.7	Min. :1.000	Min. :12.10	Min. :1.000	
##	1st Qu.:149.5	1st Qu.:3.000	1st Qu.:19.70	1st Qu.:2.000	
##	Median :162.0	Median :3.000	Median :24.70	Median :2.000	
##	Mean :155.9	Mean :2.647	Mean :25.68	Mean :2.489	
##	3rd Qu.:171.1	3rd Qu.:3.000	3rd Qu.:30.20	3rd Qu.:3.000	
##	Max. :202.6	Max. :3.000	Max. :82.90	Max. :4.000	
##	NA's :746	NA's :9592	NA's :758	NA's :6290	
##	BMXLEG	BMILEG	BMXARML	BMIARML	BMXARMC
##	Min. :24.40	Min. :1	Min. : 9.90	Min. :1	Min. :10.40
##	1st Qu.:36.00	1st Qu.:1	1st Qu.:30.50	1st Qu.:1	1st Qu.:22.60
##	Median :38.60	Median :1	Median :35.50	Median :1	Median :29.30
##	Mean :38.58	Mean :1	Mean :33.14	Mean :1	Mean :28.49
##	3rd Qu.:41.30	3rd Qu.:1	3rd Qu.:38.10	3rd Qu.:1	3rd Qu.:34.00
##	Max. :51.90	Max. :1	Max. :47.90	Max. :1	Max. :59.40
##	NA's :2411	NA's :9463	NA's :512	NA's :9445	NA's :512
##	BMIARMC	BMXWAIST	BMIWAIST	BMXSAD1	BMXSAD2
##	Min. :1	Min. : 40.20	Min. :1	Min. :10.00	Min. :10.20
##	1st Qu.:1	1st Qu.: 71.20	1st Qu.:1	1st Qu.:17.30	1st Qu.:17.30
##	Median :1	Median : 87.80	Median :1	Median :20.70	Median :20.70
##	Mean :1	Mean : 87.27	Mean :1	Mean :21.11	Mean :21.09
##	3rd Qu.:1	3rd Qu.:102.80	3rd Qu.:1	3rd Qu.:24.40	3rd Qu.:24.38
##	Max. :1	Max. :177.90	Max. :1	Max. :40.20	Max. :40.20
##	NA's :9441	NA's :1152	NA's :9374	NA's :2595	NA's :2595
##	BMXSAD3	BMXSAD4	BMDAVSAD	BMSADCM	MGDEXSTS
##	Min. :10.90	Min. :11.0	Min. :10.10	Min. :1.000	Min. :1.000
##	1st Qu.:18.50	1st Qu.:18.5	1st Qu.:17.30	1st Qu.:1.000	1st Qu.:1.000
##	Median :22.40	Median :22.4	Median :20.70	Median :1.000	Median :1.000
##	Mean :22.23	Mean :22.2	Mean :21.11	Mean :1.176	Mean :1.132
##	3rd Qu.:25.40	3rd Qu.:25.3	3rd Qu.:24.40	3rd Qu.:1.000	3rd Qu.:1.000
##	Max. :35.60	Max. :35.6	Max. :40.10	Max. :5.000	Max. :3.000
##	NA's :9455	NA's :9455	NA's :2595	NA's :9329	NA's :1522
##	MGD050	MGD060	MGQ070	MGQ080	
##	Min. :1.000	Min. :1.000	Min. :1.000	Min. :1.000	
##	1st Qu.:2.000	1st Qu.:1.000	1st Qu.:2.000	1st Qu.:1.000	
##	Median :2.000	Median :2.000	Median :2.000	Median :1.000	
##	Mean :1.967	Mean :2.115	Mean :1.883	Mean :3.283	
##	3rd Qu.:2.000	3rd Qu.:3.000	3rd Qu.:2.000	3rd Qu.:9.000	
##	Max. :9.000	Max. :9.000	Max. :9.000	Max. :9.000	
##	NA's :4602	NA's :9621	NA's :2062	NA's :8872	
##	MGQ090	MGQ100	MGQ110	MGQ120	
##	Min. :1.000	Min. :1.000	Min. :1.000	Min. :1.000	
##	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:1.000	1st Qu.:2.000	
##	Median :2.000	Median :2.000	Median :1.000	Median :2.000	
##	Mean :1.905	Mean :1.909	Mean :2.997	Mean :1.852	
##	3rd Qu.:2.000	3rd Qu.:2.000	3rd Qu.:2.000	3rd Qu.:2.000	
##	Max. :9.000	Max. :9.000	Max. :9.000	Max. :2.000	
##	NA's :8872	NA's :2061	NA's :9058	NA's :9058	
##	MGD130	MGQ90DG	MGDSEAT	MGAPHAND	MGATHAND
##	Min. :1.000	Min. :1.000	Min. :1.000	Min. :1.0	Min. :1.0
##	1st Qu.:1.000	1st Qu.:1.000	1st Qu.:1.000	1st Qu.:1.0	1st Qu.:1.0
##	Median :1.000	Median :1.000	Median :1.000	Median :1.0	Median :2.0
##	Mean :1.142	Mean :1.014	Mean :1.041	Mean :1.5	Mean :1.5
##	3rd Qu.:1.000	3rd Qu.:1.000	3rd Qu.:1.000	3rd Qu.:2.0	3rd Qu.:2.0
##	Max. :3.000	Max. :4.000	Max. :2.000	Max. :2.0	Max. :2.0

##	NA's :2006	NA's :2006	NA's :4602	NA's :2006	NA's :2006
##	MGXH1T1	MGXH1T1E	MGXH2T1	MGXH2T1E	
##	Min. : 4.00	Min. :1.000	Min. : 4.00	Min. :1.000	
##	1st Qu.:20.40	1st Qu.:1.000	1st Qu.:20.70	1st Qu.:1.000	
##	Median :27.70	Median :1.000	Median :28.20	Median :1.000	
##	Mean :28.63	Mean :1.043	Mean :29.27	Mean :1.025	
##	3rd Qu.:36.70	3rd Qu.:1.000	3rd Qu.:37.40	3rd Qu.:1.000	
##	Max. :81.50	Max. :2.000	Max. :79.50	Max. :2.000	
##	NA's :2015	NA's :2015	NA's :2131	NA's :2131	
##	MGXH1T2	MGXH1T2E	MGXH2T2	MGXH2T2E	
##	Min. : 5.00	Min. :1.000	Min. : 5.00	Min. :1.000	
##	1st Qu.:21.20	1st Qu.:1.000	1st Qu.:21.60	1st Qu.:1.000	
##	Median :28.80	Median :1.000	Median :29.00	Median :1.000	
##	Mean :29.89	Mean :1.018	Mean :30.36	Mean :1.013	
##	3rd Qu.:38.10	3rd Qu.:1.000	3rd Qu.:38.80	3rd Qu.:1.000	
##	Max. :77.60	Max. :2.000	Max. :81.40	Max. :2.000	
##	NA's :2024	NA's :2024	NA's :2141	NA's :2141	
##	MGXH1T3	MGXH1T3E	MGXH2T3	MGXH2T3E	
##	Min. : 5.00	Min. :1.000	Min. : 5.10	Min. :1.000	
##	1st Qu.:21.60	1st Qu.:1.000	1st Qu.:21.80	1st Qu.:1.000	
##	Median :29.10	Median :1.000	Median :29.50	Median :1.000	
##	Mean :30.43	Mean :1.014	Mean :30.69	Mean :1.014	
##	3rd Qu.:39.00	3rd Qu.:1.000	3rd Qu.:39.40	3rd Qu.:1.000	
##	Max. :81.20	Max. :2.000	Max. :82.80	Max. :2.000	
##	NA's :2027	NA's :2027	NA's :2149	NA's :2149	
##	MGDCGSZ	OHDEXSTS	OHDESTS	OHXIMP	
##	Min. : 8.00	Min. :1.000	Min. :1.000	Min. :1.000	
##	1st Qu.: 45.60	1st Qu.:1.000	1st Qu.:1.000	1st Qu.:2.000	
##	Median : 60.30	Median :1.000	Median :1.000	Median :2.000	
##	Mean : 63.05	Mean :1.132	Mean :1.097	Mean :1.967	
##	3rd Qu.: 80.70	3rd Qu.:1.000	3rd Qu.:1.000	3rd Qu.:2.000	
##	Max. :162.80	Max. :3.000	Max. :3.000	Max. :2.000	
##	NA's :2136	NA's :391	NA's :391	NA's :5494	
##	OHX01TC	OHX02TC	OHX03TC	OHX04TC	
##	Min. :2.000	Min. :2.000	Min. :2.000	Min. :1.000	
##	1st Qu.:4.000	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:2.000	
##	Median :4.000	Median :2.000	Median :2.000	Median :2.000	
##	Mean :3.707	Mean :2.885	Mean :2.624	Mean :2.168	
##	3rd Qu.:4.000	3rd Qu.:4.000	3rd Qu.:4.000	3rd Qu.:2.000	
##	Max. :5.000	Max. :5.000	Max. :5.000	Max. :5.000	
##	NA's :845	NA's :845	NA's :845	NA's :845	
##	OHX05TC	OHX06TC	OHX07TC	OHX08TC	OHX09TC
##	Min. :1.000	Min. :1.00	Min. :1.000	Min. :1.000	Min. :1.000
##	1st Qu.:2.000	1st Qu.:2.00	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:2.000
##	Median :2.000	Median :2.00	Median :2.000	Median :2.000	Median :2.000
##	Mean :2.171	Mean :2.02	Mean :2.101	Mean :2.103	Mean :2.104
##	3rd Qu.:2.000	3rd Qu.:2.00	3rd Qu.:2.000	3rd Qu.:2.000	3rd Qu.:2.000
##	Max. :5.000	Max. :5.00	Max. :5.000	Max. :5.000	Max. :5.000
##	NA's :845	NA's :845	NA's :845	NA's :845	NA's :845
##	OHX10TC	OHX11TC	OHX12TC	OHX13TC	OHX14TC
##	Min. :1.0	Min. :1.000	Min. :1.000	Min. :1.000	Min. :2.00
##	1st Qu.:2.0	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:2.00
##	Median :2.0	Median :2.000	Median :2.000	Median :2.000	Median :2.00
##	Mean :2.1	Mean :2.017	Mean :2.172	Mean :2.175	Mean :2.62

##	3rd Qu.:2.0	3rd Qu.:2.000	3rd Qu.:2.000	3rd Qu.:2.000	3rd Qu.:4.00
##	Max. :5.0	Max. :5.000	Max. :5.000	Max. :5.000	Max. :5.00
##	NA's :845	NA's :845	NA's :845	NA's :845	NA's :845
##	OHX15TC	OHX16TC	OHX17TC	OHX18TC	OHX19TC
##	Min. :2.00	Min. :2.000	Min. :2.000	Min. :2.000	Min. :2.000
##	1st Qu.:2.00	1st Qu.:4.000	1st Qu.:4.000	1st Qu.:2.000	1st Qu.:2.000
##	Median :2.00	Median :4.000	Median :4.000	Median :2.000	Median :2.000
##	Mean :2.89	Mean :3.711	Mean :3.669	Mean :2.907	Mean :2.688
##	3rd Qu.:4.00	3rd Qu.:4.000	3rd Qu.:4.000	3rd Qu.:4.000	3rd Qu.:4.000
##	Max. :5.00	Max. :5.000	Max. :5.000	Max. :5.000	Max. :5.000
##	NA's :845	NA's :845	NA's :845	NA's :845	NA's :845
##	OHX20TC	OHX21TC	OHX22TC	OHX23TC	
##	Min. :1.000	Min. :1.000	Min. :1.000	Min. :1.000	
##	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:2.000	
##	Median :2.000	Median :2.000	Median :2.000	Median :2.000	
##	Mean :2.102	Mean :2.047	Mean :1.955	Mean :2.008	
##	3rd Qu.:2.000	3rd Qu.:2.000	3rd Qu.:2.000	3rd Qu.:2.000	
##	Max. :5.000	Max. :5.000	Max. :5.000	Max. :5.000	
##	NA's :845	NA's :845	NA's :845	NA's :845	
##	OHX24TC	OHX25TC	OHX26TC	OHX27TC	
##	Min. :1.000	Min. :1.000	Min. :1.000	Min. :1.000	
##	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:2.000	
##	Median :2.000	Median :2.000	Median :2.000	Median :2.000	
##	Mean :2.028	Mean :2.029	Mean :2.012	Mean :1.956	
##	3rd Qu.:2.000	3rd Qu.:2.000	3rd Qu.:2.000	3rd Qu.:2.000	
##	Max. :5.000	Max. :5.000	Max. :5.000	Max. :5.000	
##	NA's :845	NA's :845	NA's :845	NA's :845	
##	OHX28TC	OHX29TC	OHX30TC	OHX31TC	
##	Min. :1.000	Min. :1.000	Min. :2.000	Min. :2.000	
##	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:2.000	
##	Median :2.000	Median :2.000	Median :2.000	Median :2.000	
##	Mean :2.046	Mean :2.106	Mean :2.682	Mean :2.894	
##	3rd Qu.:2.000	3rd Qu.:2.000	3rd Qu.:4.000	3rd Qu.:4.000	
##	Max. :5.000	Max. :5.000	Max. :5.000	Max. :5.000	
##	NA's :845	NA's :845	NA's :845	NA's :845	
##	OHX32TC	OHX02CTC	OHX03CTC	OHX04CTC	
##	Min. :2.000	Length:9813	Length:9813	Length:9813	
##	1st Qu.:4.000	Class :character	Class :character	Class :character	
##	Median :4.000	Mode :character	Mode :character	Mode :character	
##	Mean :3.663				
##	3rd Qu.:4.000				
##	Max. :5.000				
##	NA's :845				
##	OHX05CTC	OHX06CTC	OHX07CTC	OHX08CTC	
##	Length:9813	Length:9813	Length:9813	Length:9813	
##	Class :character	Class :character	Class :character	Class :character	
##	Mode :character	Mode :character	Mode :character	Mode :character	
##					
##					
##					
##	OHX09CTC	OHX10CTC	OHX11CTC	OHX12CTC	
##	Length:9813	Length:9813	Length:9813	Length:9813	
##	Class :character	Class :character	Class :character	Class :character	



##	Mode :character	Mode :character	Mode :character	Mode :character
##				
##				
##				
##	OHX13CTC	OHX14CTC	OHX15CTC	OHX18CTC
##	Length:9813	Length:9813	Length:9813	Length:9813
##	Class :character	Class :character	Class :character	Class :character
##	Mode :character	Mode :character	Mode :character	Mode :character
##				
##				
##				
##	OHX19CTC	OHX20CTC	OHX21CTC	OHX22CTC
##	Length:9813	Length:9813	Length:9813	Length:9813
##	Class :character	Class :character	Class :character	Class :character
##	Mode :character	Mode :character	Mode :character	Mode :character
##				
##				
##				
##	OHX23CTC	OHX24CTC	OHX25CTC	OHX26CTC
##	Length:9813	Length:9813	Length:9813	Length:9813
##	Class :character	Class :character	Class :character	Class :character
##	Mode :character	Mode :character	Mode :character	Mode :character
##				
##				
##				
##	OHX27CTC	OHX28CTC	OHX29CTC	OHX30CTC
##	Length:9813	Length:9813	Length:9813	Length:9813
##	Class :character	Class :character	Class :character	Class :character
##	Mode :character	Mode :character	Mode :character	Mode :character
##				
##				
##				
##	OHX31CTC	OHX02CSC	OHX03CSC	OHX04CSC
##	Length:9813	Length:9813	Length:9813	Length:9813
##	Class :character	Class :character	Class :character	Class :character
##	Mode :character	Mode :character	Mode :character	Mode :character
##				
##				
##				
##	OHX05CSC	OHX06CSC	OHX07CSC	OHX08CSC
##	Length:9813	Length:9813	Length:9813	Length:9813
##	Class :character	Class :character	Class :character	Class :character
##	Mode :character	Mode :character	Mode :character	Mode :character
##				
##				
##				
##	OHX09CSC	OHX10CSC	OHX11CSC	OHX12CSC

##	Length:9813	Length:9813	Length:9813	Length:9813	
##	Class :character	Class :character	Class :character	Class :character	
##	Mode :character	Mode :character	Mode :character	Mode :character	
##					
##					
##					
##					
##	OHX13CSC	OHX14CSC	OHX15CSC	OHX18CSC	
##	Length:9813	Length:9813	Length:9813	Length:9813	
##	Class :character	Class :character	Class :character	Class :character	
##	Mode :character	Mode :character	Mode :character	Mode :character	
##					
##					
##					
##	OHX19CSC	OHX20CSC	OHX21CSC	OHX22CSC	
##	Length:9813	Length:9813	Length:9813	Length:9813	
##	Class :character	Class :character	Class :character	Class :character	
##	Mode :character	Mode :character	Mode :character	Mode :character	
##					
##					
##					
##	OHX23CSC	OHX24CSC	OHX25CSC	OHX26CSC	
##	Length:9813	Length:9813	Length:9813	Length:9813	
##	Class :character	Class :character	Class :character	Class :character	
##	Mode :character	Mode :character	Mode :character	Mode :character	
##					
##					
##					
##	OHX27CSC	OHX28CSC	OHX29CSC	OHX30CSC	
##	Length:9813	Length:9813	Length:9813	Length:9813	
##	Class :character	Class :character	Class :character	Class :character	
##	Mode :character	Mode :character	Mode :character	Mode :character	
##					
##					
##					
##	OHX31CSC	OHX02SE	OHX03SE	OHX04SE	
##	Length:9813	Min. : 0.000	Min. : 0.000	Min. :0.000	
##	Class :character	1st Qu.: 0.000	1st Qu.: 0.000	1st Qu.:0.000	
##	Mode :character	Median : 9.000	Median : 1.000	Median :0.000	
##		Mean : 5.719	Mean : 3.057	Mean :0.376	
##		3rd Qu.: 9.000	3rd Qu.: 9.000	3rd Qu.:0.000	
##		Max. :13.000	Max. :13.000	Max. :9.000	
##		NA's :6549	NA's :6549	NA's :6549	
##	OHX05SE	OHX07SE	OHX10SE	OHX12SE	OHX13SE
##	Min. :0.000	Min. :0.000	Min. :0.000	Min. :0.000	Min. :0.00
##	1st Qu.:0.000	1st Qu.:0.000	1st Qu.:0.000	1st Qu.:0.000	1st Qu.:0.00
##	Median :0.000	Median :0.000	Median :0.000	Median :0.000	Median :0.00
##	Mean :0.642	Mean :3.185	Mean :3.175	Mean :0.624	Mean :0.37
##	3rd Qu.:0.000	3rd Qu.:9.000	3rd Qu.:9.000	3rd Qu.:0.000	3rd Qu.:0.00
##	Max. :9.000	Max. :9.000	Max. :9.000	Max. :9.000	Max. :9.00

##	NA's :6549	NA's :6549	NA's :6549	NA's :6549	NA's :6549
##	OHX14SE	OHX15SE	OHX18SE	OHX19SE	
##	Min. : 0.000	Min. : 0.000	Min. : 0.000	Min. : 0.000	
##	1st Qu.: 0.000	1st Qu.: 0.000	1st Qu.: 0.000	1st Qu.: 0.000	
##	Median : 1.000	Median : 9.000	Median : 9.000	Median : 1.000	
##	Mean : 3.043	Mean : 5.698	Mean : 5.425	Mean : 2.662	
##	3rd Qu.: 9.000	3rd Qu.: 9.000	3rd Qu.: 9.000	3rd Qu.: 9.000	
##	Max. :13.000	Max. :13.000	Max. :12.000	Max. :12.000	
##	NA's :6549	NA's :6549	NA's :6549	NA's :6549	
##	OHX20SE	OHX21SE	OHX28SE	OHX29SE	
##	Min. :0.000	Min. :0.0	Min. :0.000	Min. :0.000	
##	1st Qu.:0.000	1st Qu.:0.0	1st Qu.:0.000	1st Qu.:0.000	
##	Median :0.000	Median :0.0	Median :0.000	Median :0.000	
##	Mean :0.464	Mean :0.7	Mean :0.695	Mean :0.425	
##	3rd Qu.:0.000	3rd Qu.:0.0	3rd Qu.:0.000	3rd Qu.:0.000	
##	Max. :9.000	Max. :9.0	Max. :9.000	Max. :9.000	
##	NA's :6549	NA's :6549	NA's :6549	NA's :6549	
##	OHX30SE	OHX31SE	CSXEXSTS	CSXEXCMT	
##	Min. : 0.000	Min. : 0.000	Min. :1.000	Min. : 1.00	
##	1st Qu.: 0.000	1st Qu.: 0.000	1st Qu.:1.000	1st Qu.: 7.00	
##	Median : 1.000	Median : 9.000	Median :1.000	Median : 56.00	
##	Mean : 2.662	Mean : 5.472	Mean :1.207	Mean : 45.26	
##	3rd Qu.: 9.000	3rd Qu.: 9.000	3rd Qu.:1.000	3rd Qu.: 56.00	
##	Max. :12.000	Max. :12.000	Max. :3.000	Max. :122.00	
##	NA's :6549	NA's :6549	NA's :6105	NA's :9605	
##	CSQ245	CSQ241	CSQ260A	CSQ260D	CSQ260G
##	Min. :1.000	Min. :1.000	Min. :1.000	Min. :1	Min. :1
##	1st Qu.:2.000	1st Qu.:2.000	1st Qu.:1.000	1st Qu.:1	1st Qu.:1
##	Median :2.000	Median :2.000	Median :1.000	Median :1	Median :1
##	Mean :2.221	Mean :2.006	Mean :1.198	Mean :1	Mean :1
##	3rd Qu.:2.000	3rd Qu.:2.000	3rd Qu.:1.000	3rd Qu.:1	3rd Qu.:1
##	Max. :9.000	Max. :9.000	Max. :9.000	Max. :1	Max. :1
##	NA's :6256	NA's :8834	NA's :9631	NA's :9739	NA's :9530
##	CSQ260I	CSQ260N	CSQ260M	CSQ270	CSQ450
##	Min. :1	Min. :1	Min. :1	Min. :1.000	Min. : 0.00
##	1st Qu.:1	1st Qu.:1	1st Qu.:1	1st Qu.:1.000	1st Qu.: 15.00
##	Median :1	Median :1	Median :1	Median :1.000	Median : 17.00
##	Mean :1	Mean :1	Mean :1	Mean :1.534	Mean : 21.35
##	3rd Qu.:1	3rd Qu.:1	3rd Qu.:1	3rd Qu.:2.000	3rd Qu.: 27.00
##	Max. :1	Max. :1	Max. :1	Max. :9.000	Max. :100.00
##	NA's :9708	NA's :9392	NA's :7059	NA's :9530	NA's :6393
##	CSQ460	CSQ470	CSQ480	CSQ490	
##	Min. : 0.00	Min. : 0.0	Min. : 0.00	Min. : 0.00	
##	1st Qu.: 30.00	1st Qu.: 17.0	1st Qu.: 5.00	1st Qu.: 48.00	
##	Median : 35.00	Median : 27.0	Median : 9.00	Median : 55.00	
##	Mean : 41.08	Mean : 29.8	Mean : 11.46	Mean : 60.95	
##	3rd Qu.: 51.00	3rd Qu.: 37.0	3rd Qu.: 15.00	3rd Qu.: 80.00	
##	Max. :100.00	Max. :100.0	Max. :100.00	Max. :100.00	
##	NA's :6397	NA's :6401	NA's :6402	NA's :6402	
##	CSXQUIPG	CSXQUIPT	CSXNAPG	CSXNAPT	
##	Min. : 0.00	Min. :1.000	Min. : 0.00	Min. :1.000	
##	1st Qu.: 5.00	1st Qu.:2.000	1st Qu.: 15.00	1st Qu.:1.000	
##	Median : 10.00	Median :3.000	Median : 25.00	Median :1.000	
##	Mean : 14.85	Mean :3.008	Mean : 26.93	Mean :1.466	

```
## 3rd Qu.: 20.00 3rd Qu.:4.000 3rd Qu.: 35.00 3rd Qu.:1.000
## Max. :100.00 Max. :5.000 Max. :100.00 Max. :5.000
## NA's :6680 NA's :6680 NA's :6684 NA's :6596
## CSXQVISG CSXQUIST CSXSLTSG CSXSLTST
## Min. : 0.00 Min. :1.000 Min. : 2.00 Min. :1.000
## 1st Qu.: 35.00 1st Qu.:2.000 1st Qu.: 35.00 1st Qu.:1.000
## Median : 52.00 Median :2.000 Median : 50.00 Median :1.000
## Mean : 53.21 Mean :2.251 Mean : 52.06 Mean :1.064
## 3rd Qu.: 65.00 3rd Qu.:2.000 3rd Qu.: 63.00 3rd Qu.:1.000
## Max. :100.00 Max. :5.000 Max. :100.00 Max. :5.000
## NA's :6699 NA's :6699 NA's :6596 NA's :6596
## CSXNASG CSXNAST CSXTSEQ CSXCHOOD
## Min. : 0.00 Min. :1.000 Length:9813 Min. :1.000
## 1st Qu.: 18.25 1st Qu.:1.000 Class :character 1st Qu.:2.000
## Median : 30.00 Median :1.000 Mode :character Median :2.000
## Mean : 33.08 Mean :1.242 Mean :2.152
## 3rd Qu.: 42.00 3rd Qu.:1.000 3rd Qu.:2.000
## Max. :100.00 Max. :5.000 Max. :4.000
## NA's :6595 NA's :6595 NA's :6286
## CSXSOD CSXSMKOD CSXLEAOD CSXSOAOD
## Min. :1.000 Min. :1.000 Min. :1.000 Min. :1.000
## 1st Qu.:1.000 1st Qu.:3.000 1st Qu.:3.000 1st Qu.:1.000
## Median :1.000 Median :3.000 Median :3.000 Median :1.000
## Mean :1.428 Mean :2.903 Mean :2.816 Mean :1.145
## 3rd Qu.:1.000 3rd Qu.:3.000 3rd Qu.:3.000 3rd Qu.:1.000
## Max. :4.000 Max. :4.000 Max. :4.000 Max. :4.000
## NA's :6288 NA's :6290 NA's :6293 NA's :6293
## CSXGRAOD CSXONOD CSXNGSOD CSXSLTRT
## Min. :1.000 Min. :1.000 Min. :1.000 Min. : 1.00
## 1st Qu.:2.000 1st Qu.:3.000 1st Qu.:4.000 1st Qu.: 35.00
## Median :2.000 Median :3.000 Median :4.000 Median : 53.00
## Mean :2.156 Mean :2.992 Mean :3.751 Mean : 55.75
## 3rd Qu.:3.000 3rd Qu.:3.000 3rd Qu.:4.000 3rd Qu.: 74.00
## Max. :4.000 Max. :4.000 Max. :4.000 Max. :100.00
## NA's :6294 NA's :6294 NA's :6294 NA's :8218
## CSXSLTRG CSXNART CSXNARG CSAEFFRT
## Min. :1.000 Min. : 0.00 Min. :1.000 Min. :1.000
## 1st Qu.:1.000 1st Qu.: 16.00 1st Qu.:1.000 1st Qu.:1.000
## Median :1.000 Median : 26.00 Median :1.000 Median :1.000
## Mean :1.066 Mean : 31.53 Mean :1.199 Mean :1.709
## 3rd Qu.:1.000 3rd Qu.: 42.00 3rd Qu.:1.000 3rd Qu.:2.000
## Max. :5.000 Max. :100.00 Max. :5.000 Max. :5.000
## NA's :8218 NA's :8200 NA's :8200 NA's :6276
```

```
# Keeping columns which have less than or equal to 50% missing values (NAs)
```

```
DF <- df[apply(df,2,function(x) mean(is.na(x))<=.50)]
```

```
# We can see that there are no more columns which have more than 50% missing values
```

```
write_csv(DF, "C:/Semester 2/Intro to Data Mining and Processing/Project/national-health-and-nutrition-")
```

```
# Discarding remaining irrelevant features from dataframe and selecting only
```

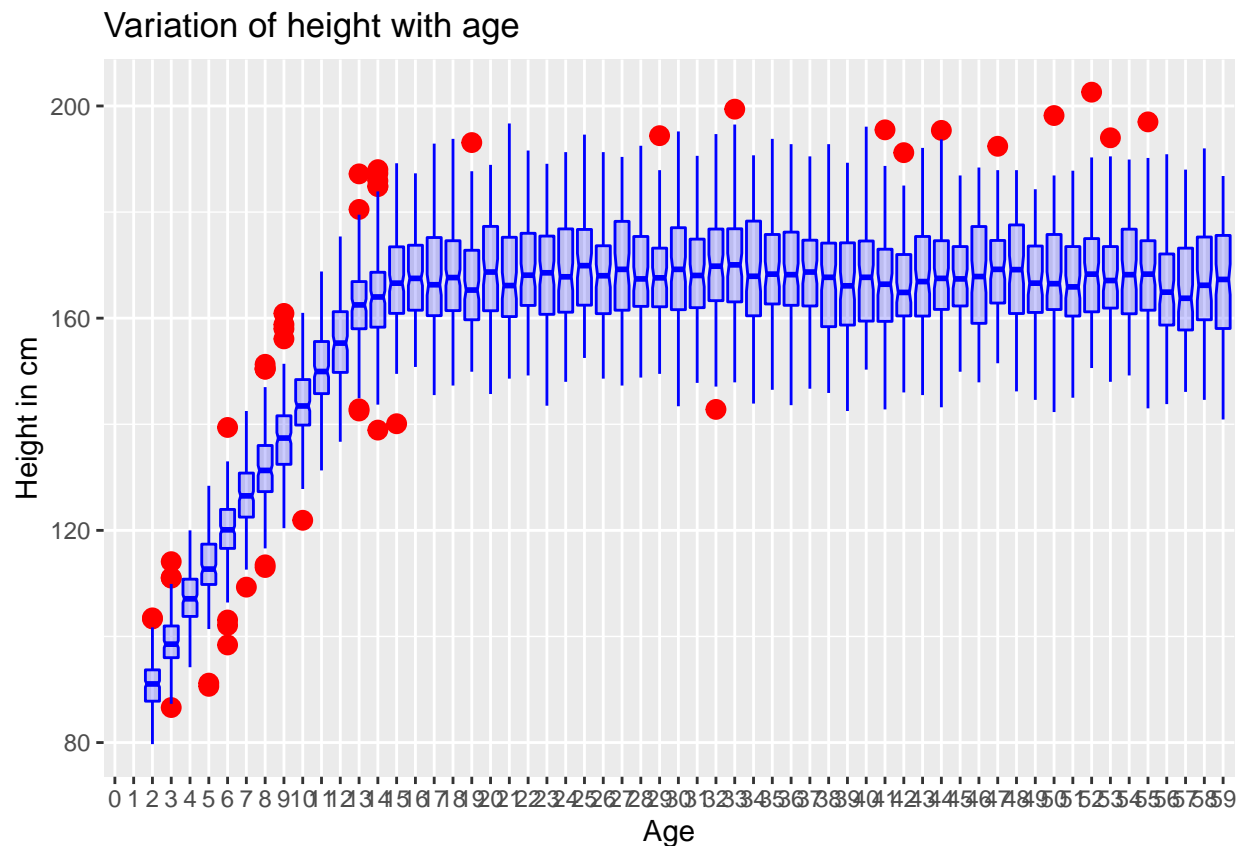
```

#useful features
# Gender not mentioned in documentation

DF1 <- DF %>% select(SEQN, HbA1c, AGE, "BP_SYS" = BPXSY1,
                    "BP_DIA" = BPXD11, "WEIGHT" = BMXWT, "HEIGHT" = BMXHT,
                    "BMI" = BMXBMI, "UPPER_LEG_LENGTH" = BMXLEG,
                    "UPPER_ARM_LENGTH" = BMXARML, "ARM_CIRCUM" = BMXARMC,
                    "WAIST_CIRCUMF" = BMXWAIST, "SAG_ABDOMINAL_DIA" = BMDAVSAD)

# 1) How does height vary with age? Are there any anomalies?
# Plot an age vs Height Graph
DF1 %>%
  filter(AGE<60) %>%
  ggplot(aes(x = as.factor(AGE), y = HEIGHT)) +
  geom_boxplot(color = "blue", fill = "blue",
              alpha = 0.2, notch = TRUE, notchwidth = 0.8, outlier.colour = "red",
              outlier.fill = "red", outlier.size = 3, outlier.alpha = 1) +
  labs(title = "Variation of height with age",
       x = "Age",
       y = "Height in cm")

```



```

# Till about 18 years height increases with age, but after that it seems to
# be constant which is as expected

```

```

# Removing observations less than 5 years old
DF1 <- subset(DF1, AGE>=5)

# Imputing missing values of remaining columns with median
f=function(x){
  x<-as.numeric(as.character(x)) #first convert each column into numeric if it
                                #is from factor
  x[is.na(x)] =median(x, na.rm=TRUE) #convert the item with NA to median value
                                #from the column
  x #display the column
}

Final_DF = data.frame(apply(DF1,2,f))
f_df <- data.frame(apply(DF1,2,f))

summary(Final_DF)

```

```

##      SEQN      HbA1c      AGE      BP_SYS
## Min.   :73557  Min.   : 3.50  Min.   : 5.0  Min.   : 66.0
## 1st Qu.:76118  1st Qu.: 5.20  1st Qu.:15.0  1st Qu.:108.0
## Median :78682  Median : 5.40  Median :34.0  Median :116.0
## Mean   :78669  Mean   : 5.59  Mean   :36.3  Mean   :117.8
## 3rd Qu.:81210  3rd Qu.: 5.60  3rd Qu.:56.0  3rd Qu.:124.0
## Max.   :83731  Max.   :17.50  Max.   :80.0  Max.   :228.0
##      BP_DIA      WEIGHT      HEIGHT      BMI
## Min.   : 0.00  Min.   : 13.10  Min.   : 90.6  Min.   :12.10
## 1st Qu.: 60.00  1st Qu.: 53.30  1st Qu.:153.9  1st Qu.:20.90
## Median : 66.00  Median : 69.80  Median :163.2  Median :25.50
## Mean   : 65.81  Mean   : 70.38  Mean   :160.4  Mean   :26.39
## 3rd Qu.: 74.00  3rd Qu.: 86.60  3rd Qu.:171.8  3rd Qu.:30.60
## Max.   :122.00  Max.   :222.60  Max.   :202.6  Max.   :82.90
## UPPER_LEG_LENGTH UPPER_ARM_LENGTH  ARM_CIRCUM  WAIST_CIRCUMF
## Min.   :24.40  Min.   :17.0  Min.   :13.90  Min.   : 41.50
## 1st Qu.:36.40  1st Qu.:33.5  1st Qu.:26.10  1st Qu.: 76.00
## Median :38.60  Median :36.0  Median :30.50  Median : 89.80
## Mean   :38.58  Mean   :35.4  Mean   :30.31  Mean   : 90.01
## 3rd Qu.:41.00  3rd Qu.:38.4  3rd Qu.:34.50  3rd Qu.:103.20
## Max.   :51.90  Max.   :47.9  Max.   :59.40  Max.   :177.90
## SAG ABDOMINAL_DIA
## Min.   :10.10
## 1st Qu.:17.90
## Median :20.70
## Mean   :21.05
## 3rd Qu.:23.60
## Max.   :40.10

```

```

# Changing the HbA1c values to yes or no diabetes

#No Diabetes 6.4 and lower Yes Diabetes 6.5 or higher

Final_DF$HbA1c <- cut(Final_DF$HbA1c,
                      breaks = c(-Inf, 6.5, Inf),
                      labels = c("NO", "YES"),

```

```

right = FALSE) # Interval is closed on the left and open
                #on right

names(Final_DF)[names(Final_DF) == "HbA1c"] <- "Diabetes" # Renaming column
str(Final_DF)

```

```

## 'data.frame': 8489 obs. of 13 variables:
## $ SEQN : num 73557 73558 73559 73560 73561 ...
## $ Diabetes : Factor w/ 2 levels "NO","YES": 2 2 2 1 1 1 1 1 1 1 ...
## $ AGE : num 69 54 72 9 73 56 61 56 65 26 ...
## $ BP_SYS : num 122 156 140 108 136 160 118 128 140 106 ...
## $ BP_DIA : num 72 62 90 38 86 84 80 74 78 60 ...
## $ WEIGHT : num 78.3 89.5 88.9 32.2 52 105 93.4 61.8 65.3 47.1 ...
## $ HEIGHT : num 171 177 175 137 162 ...
## $ BMI : num 26.7 28.6 28.9 17.1 19.7 41.7 35.7 26.5 22 20.3 ...
## $ UPPER_LEG_LENGTH : num 39.2 40 40 33.5 36.3 34.2 37.1 32.4 40 34.4 ...
## $ UPPER_ARM_LENGTH : num 40.2 41.5 41 29.5 37.5 36.2 39.3 33.5 40.3 32.6 ...
## $ ARM_CIRCUM : num 35.3 34.7 33.5 21 25.2 41.8 38 29 27.5 25.8 ...
## $ WAIST_CIRCUMF : num 100 107.6 109.2 61 89.8 ...
## $ SAG ABDOMINAL_DIA: num 20.6 24.4 25.6 14.9 20.7 29.1 26.7 19.9 20 14.5 ...

```

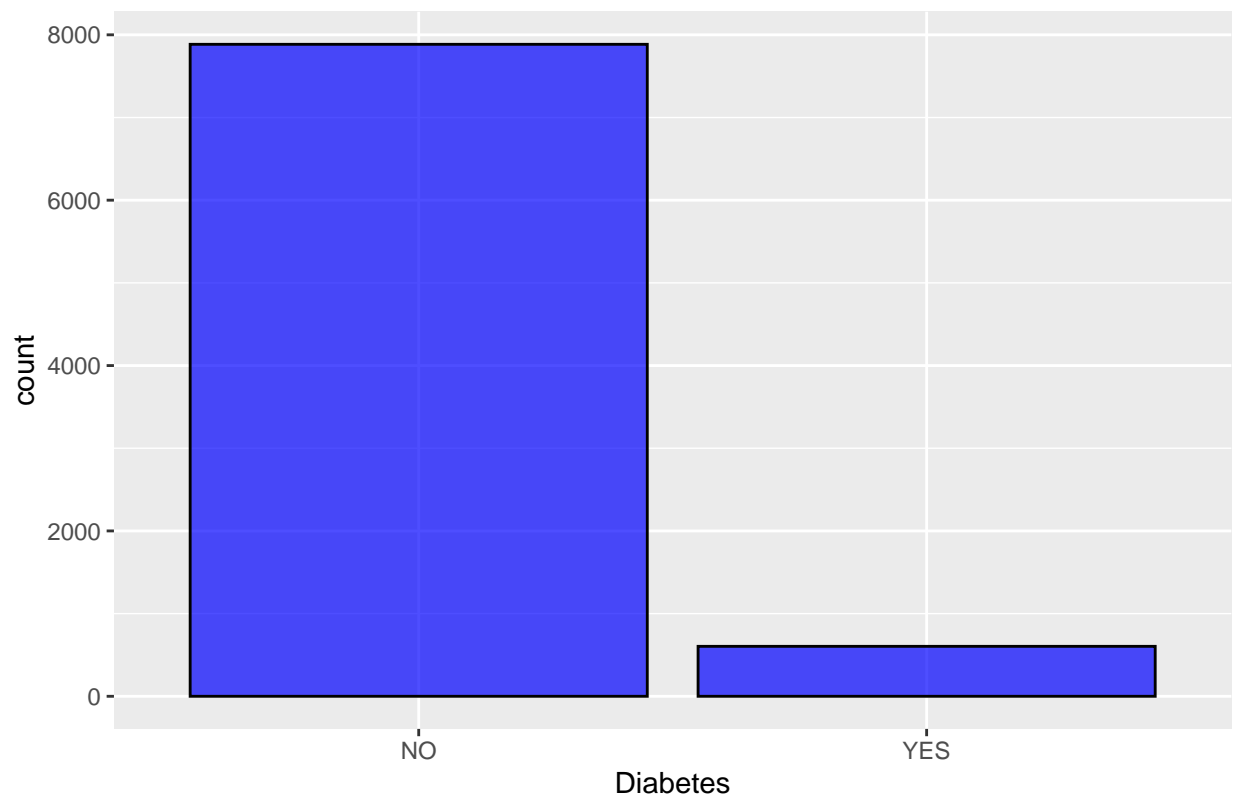
*# 2) Number of people who have vs don't have diabetes*

```

Final_DF %>%
  ggplot(aes(x = Diabetes)) + geom_bar(data = Final_DF, fill = "blue",
                                       color = "black", alpha = 0.7) +
  labs(title = "Number of people that have diabetes vs number of people that don't have diabetes")

```

Number of people that have diabetes vs number of people that don't have



```
###
#EDA
###

#SEQN, HbA1c, AGE, "BP_SYS" = BPXSY1,
#"BP_DIA" = BPXDI1, "WEIGHT" = BMXWT, "HEIGHT" = BMXHT,
#"BMI" = BMXBMI, "UPPER_LEG_LENGTH" = BMXLEG,
#"UPPER_ARM_LENGTH" = BMXARML, "ARM_CIRCUM" = BMXARMC,
#"WAIST_CIRCUMF" = BMXWAIST, "SAG ABDOMINAL_DIA" = BMDAVSAD

# Make a dataset with people with only diabetes

diabetes_data <- Final_DF %>%
  filter(Diabetes == "YES")

x1 <- diabetes_data %>%
  filter(AGE>5) %>%
  filter(AGE<=15) %>%
  count(Diabetes == "YES")

x2 <- diabetes_data %>%
  filter(AGE>15) %>%
  filter(AGE<=25) %>%
  count(Diabetes == "YES")
```



```

x3 <- diabetes_data %>%
  filter(AGE>25) %>%
  filter(AGE<=35) %>%
  count(Diabetes == "YES")

x4 <- diabetes_data %>%
  filter(AGE>35) %>%
  filter(AGE<=45) %>%
  count(Diabetes == "YES")

x5 <- diabetes_data %>%
  filter(AGE>45) %>%
  filter(AGE<=55) %>%
  count(Diabetes == "YES")

x6 <- diabetes_data %>%
  filter(AGE>55) %>%
  filter(AGE<=65) %>%
  count(Diabetes == "YES")

x7 <- diabetes_data %>%
  filter(AGE>65) %>%
  count(Diabetes == "YES")

age_groups <- x1 %>% full_join(x2)%>% full_join(x3)%>%
  full_join(x4)%>% full_join(x5)%>% full_join(x6)%>% full_join(x7)

## Joining, by = c("Diabetes == \"YES\"", "n")

## Joining, by = c("Diabetes == \"YES\"", "n")
## Joining, by = c("Diabetes == \"YES\"", "n")
## Joining, by = c("Diabetes == \"YES\"", "n")
## Joining, by = c("Diabetes == \"YES\"", "n")

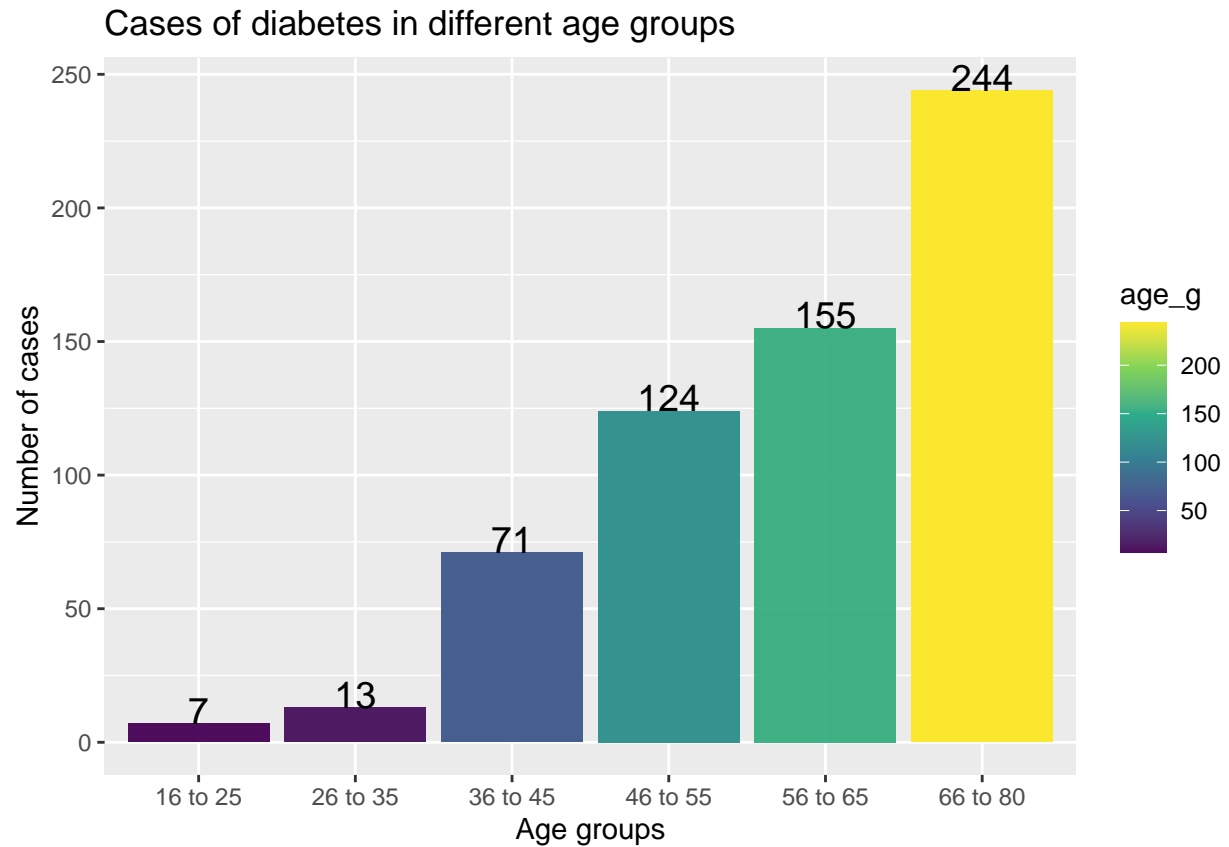
col_names <- c("16 to 25", "26 to 35", "36 to 45", "46 to 55", "56 to 65",
               "66 to 80" )
age_g <- c( 7, 13, 71, 124, 155, 244)

x8 <- data.frame(col_names, age_g)
x8 <- x8 %>%
  group_by(age_g)

# 3) What age group seems to have diabetes more frequently

ggplot(x8, aes(x=col_names,y = age_g, fill = age_g)) + geom_col() +
  scale_fill_viridis_c(alpha = 0.95) +
  geom_text(aes(label = age_g), vjust = 0.01, size = 5) +
  labs(title = "Cases of diabetes in different age groups",
       x = "Age groups",
       y = "Number of cases")

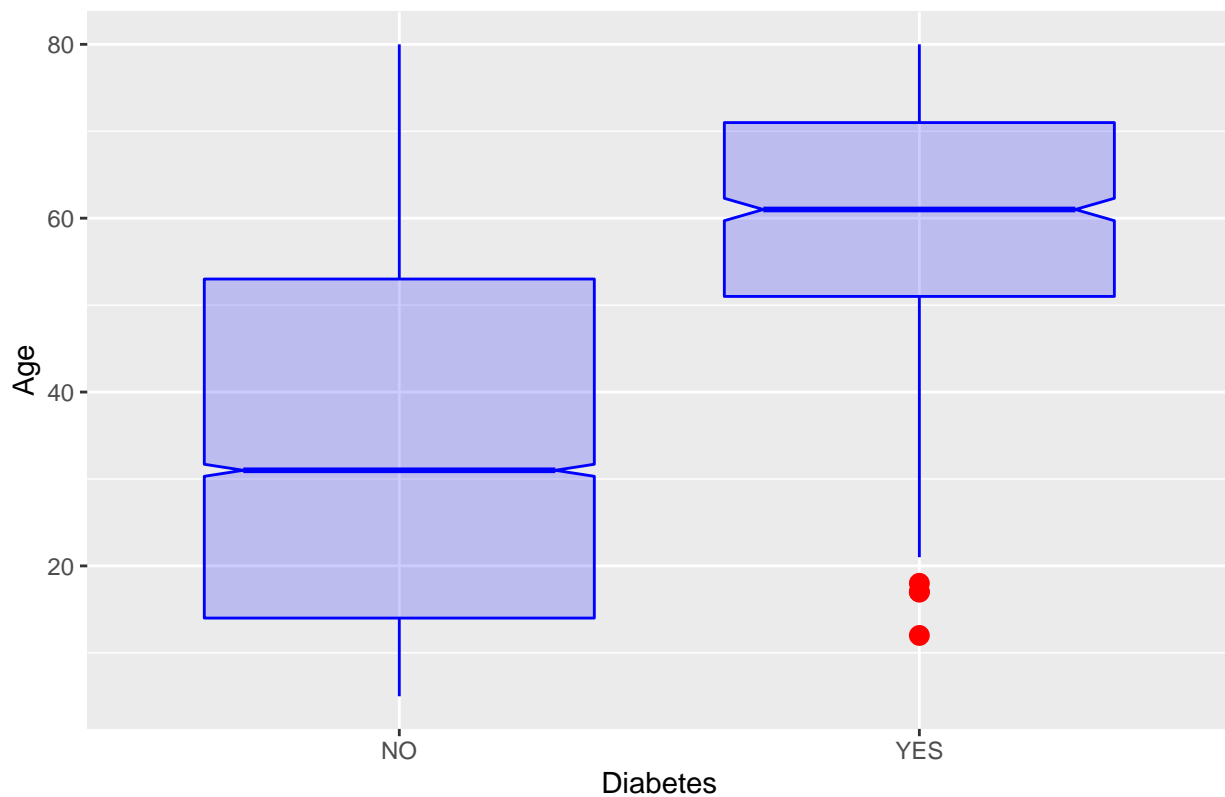
```



*# 4) Distribution of people having diabetes with AGE*

```
ggplot(Final_DF, aes(x = Diabetes, y = AGE)) +  
  geom_boxplot(color = "blue", fill = "blue", alpha = 0.2,  
               notch = TRUE, notchwidth = 0.8, outlier.colour = "red",  
               outlier.size = 3, outlier.alpha = 1) +  
  labs(title = "Distribution of people having diabetes with age",  
       x = "Diabetes",  
       y = "Age")
```

Distribution of people having diabetes with age



*# 5) Relation between BMI and diabetes*

```
bmi_under <- diabetes_data %>%
  filter(BMI<18.5) %>%
  count(Diabetes == "YES")

bmi_normal <- diabetes_data %>%
  filter(BMI>=18.5) %>%
  filter(BMI<25) %>%
  count(Diabetes == "YES")

bmi_over <- diabetes_data %>%
  filter(BMI>=25) %>%
  filter(BMI<30) %>%
  count(Diabetes == "YES")

bmi_obese <- diabetes_data %>%
  filter(BMI>=30) %>%
  count(Diabetes == "YES")

bmi_groups <- bmi_under %>% full_join(bmi_normal)%>% full_join(bmi_over)%>%
  full_join(bmi_obese)
```

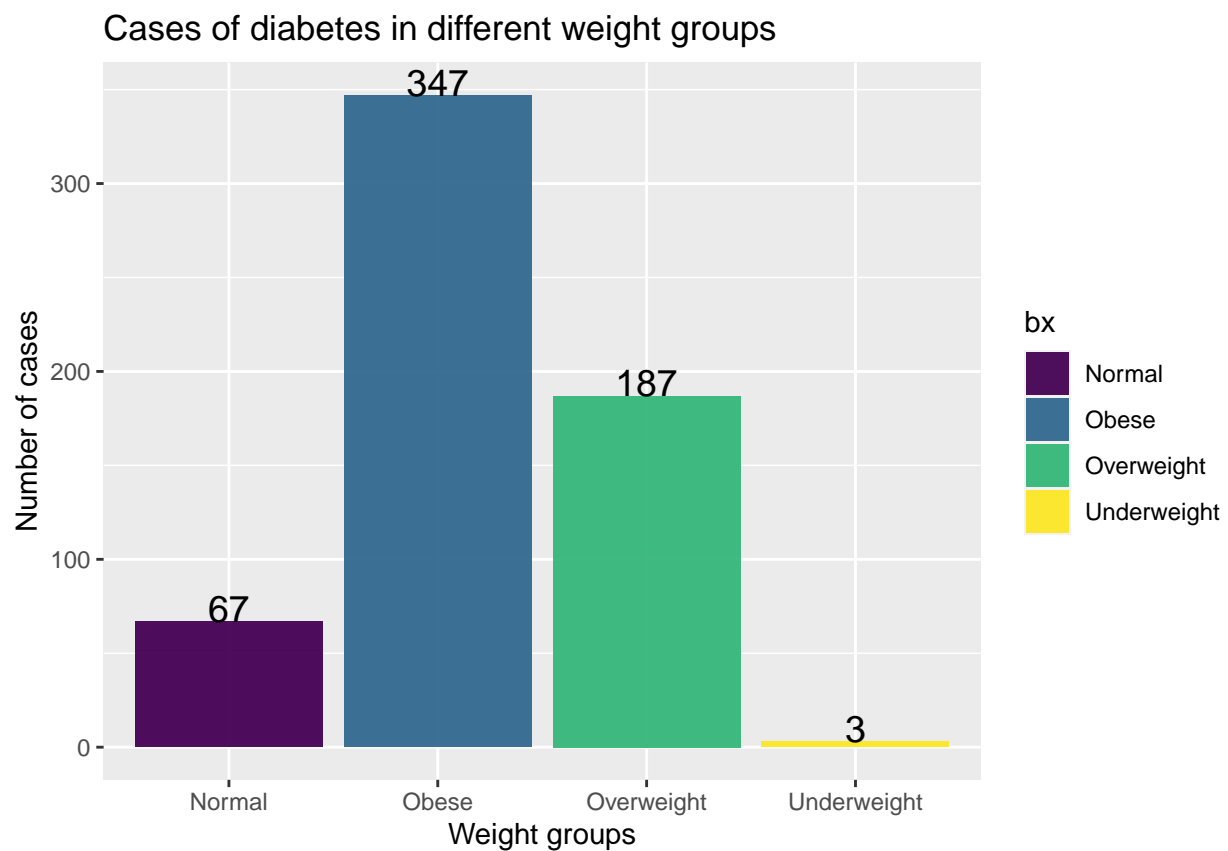
```
## Joining, by = c("Diabetes == \"YES\"", "n")
```

```
## Joining, by = c("Diabetes == \"YES\"", "n")
## Joining, by = c("Diabetes == \"YES\"", "n")

bx <- c("Underweight", "Normal", "Overweight", "Obese")
by <- c( 3, 67, 187, 347)

bd <- data.frame(bx, by)

ggplot(bd, aes(x=bx, y = by, fill = bx)) + geom_col() +
  scale_fill_viridis_d(alpha = 0.95) +
  geom_text(aes(label = by), vjust = 0.01, size = 5) +
  labs(title = "Cases of diabetes in different weight groups",
       x = "Weight groups",
       y = "Number of cases")
```

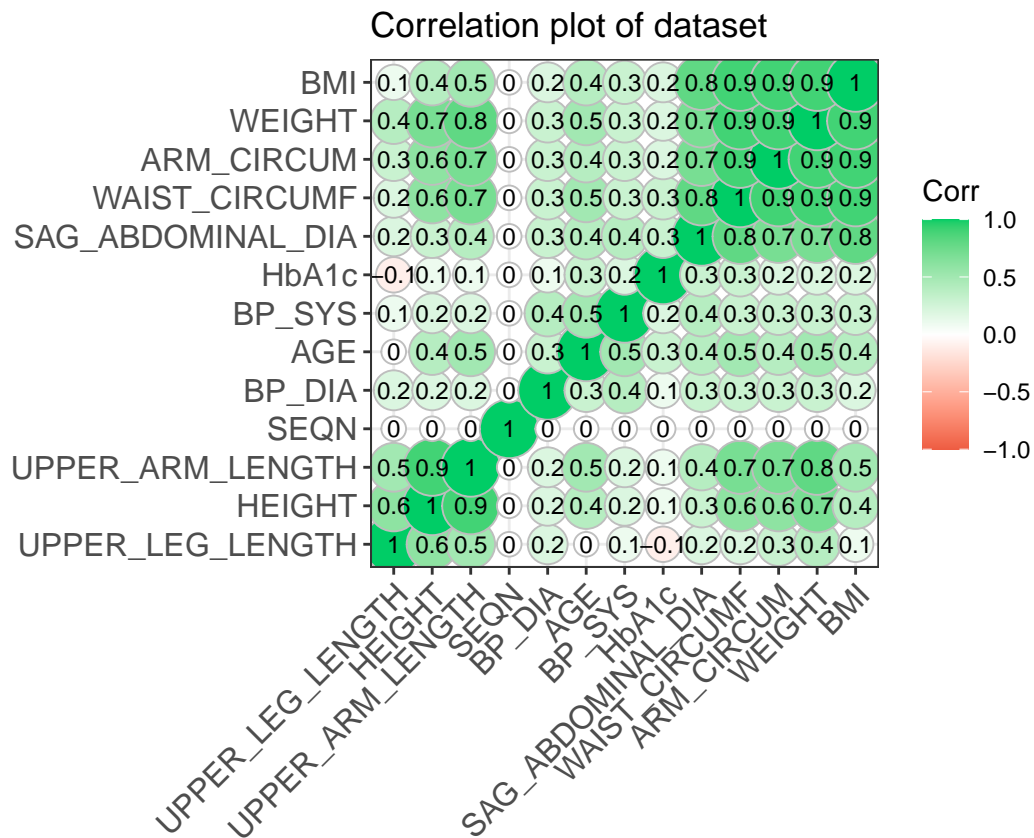


```
#Correlation Plot
library(ggcorrplot)

corr <- round(cor(f_df), 1)

ggcorrplot(corr, hc.order = TRUE,
           lab = TRUE,
           lab_size = 3,
           method = "circle",
           colors = c("tomato2", "white", "springgreen3"),
```

```
title = "Correlation plot of dataset",
ggtheme = theme_bw())
```



```
## Feature importance using Boruta Package
```

```
boruta_output <- Boruta(Diabetes ~ . , data=na.omit(Final_DF), maxRuns = 200,
doTrace=2)
```

```
## 1. run of importance source...
```

```
## 2. run of importance source...
```

```
## 3. run of importance source...
```

```
## 4. run of importance source...
```

```
## 5. run of importance source...
```

```
## 6. run of importance source...
```

```
## 7. run of importance source...
```

```
## 8. run of importance source...
```

```
## 9. run of importance source...
## 10. run of importance source...
## 11. run of importance source...
## After 11 iterations, +48 secs:
## confirmed 11 attributes: AGE, ARM_CIRCUM, BMI, BP_DIA, BP_SYS and 6 more;
## still have 1 attribute left.
## 12. run of importance source...
## 13. run of importance source...
## 14. run of importance source...
## 15. run of importance source...
## 16. run of importance source...
## 17. run of importance source...
## 18. run of importance source...
## 19. run of importance source...
## 20. run of importance source...
## 21. run of importance source...
## After 21 iterations, +1.5 mins:
## rejected 1 attribute: SEQN;
## no more attributes left.
```

```
print(boruta_output)
```

```
## Boruta performed 21 iterations in 1.48958 mins.
## 11 attributes confirmed important: AGE, ARM_CIRCUM, BMI, BP_DIA,
## BP_SYS and 6 more;
## 1 attributes confirmed unimportant: SEQN;
```

```
#names(boruta_output)
```

```
boruta_signif <- getSelectedAttributes(boruta_output, withTentative = TRUE)
print(boruta_signif)
```

```
## [1] "AGE"           "BP_SYS"        "BP_DIA"
## [4] "WEIGHT"        "HEIGHT"        "BMI"
## [7] "UPPER_LEG_LENGTH" "UPPER_ARM_LENGTH" "ARM_CIRCUM"
## [10] "WAIST_CIRCUMF" "SAG ABDOMINAL_DIA"
```

```
roughFixMod <- TentativeRoughFix(boruta_output)
boruta_signif <- getSelectedAttributes(roughFixMod)
print(boruta_signif)
```

```
## [1] "AGE"          "BP_SYS"       "BP_DIA"
## [4] "WEIGHT"       "HEIGHT"       "BMI"
## [7] "UPPER_LEG_LENGTH" "UPPER_ARM_LENGTH" "ARM_CIRCUM"
## [10] "WAIST_CIRCUMF" "SAG ABDOMINAL_DIA"
```

```
# Variable Importance Scores
imps <- attStats(roughFixMod)
imps2 = imps[imps$decision != 'Rejected', c('meanImp', 'decision')]
imps2[order(-imps2$meanImp), ] # descending sort
```

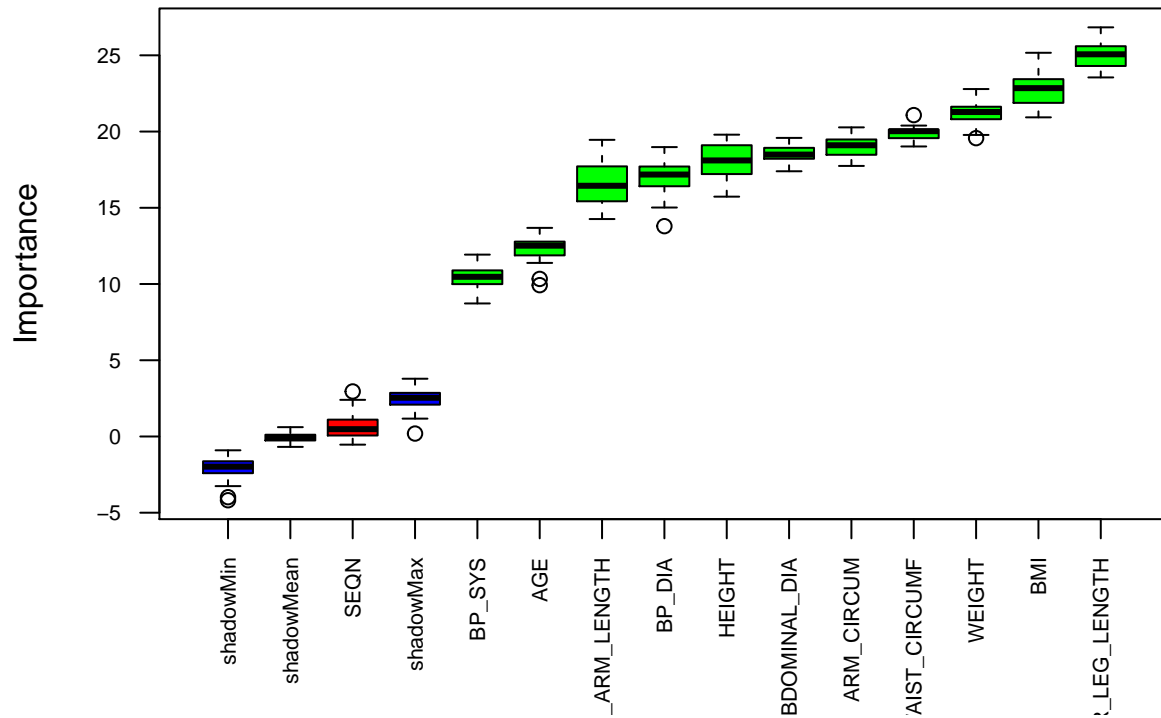
```
##          meanImp decision
## UPPER_LEG_LENGTH 25.05469 Confirmed
## BMI              22.80465 Confirmed
## WEIGHT           21.22875 Confirmed
## WAIST_CIRCUMF    19.89443 Confirmed
## ARM_CIRCUM       19.01839 Confirmed
## SAG ABDOMINAL_DIA 18.52763 Confirmed
## HEIGHT           18.12503 Confirmed
## BP_DIA           16.99411 Confirmed
## UPPER_ARM_LENGTH 16.55303 Confirmed
## AGE              12.32013 Confirmed
## BP_SYS           10.36542 Confirmed
```

```
setDT(imps2, keep.rownames = TRUE)[, ]
```

```
##          rn meanImp decision
## 1:          AGE 12.32013 Confirmed
## 2:         BP_SYS 10.36542 Confirmed
## 3:         BP_DIA 16.99411 Confirmed
## 4:         WEIGHT 21.22875 Confirmed
## 5:         HEIGHT 18.12503 Confirmed
## 6:          BMI 22.80465 Confirmed
## 7: UPPER_LEG_LENGTH 25.05469 Confirmed
## 8: UPPER_ARM_LENGTH 16.55303 Confirmed
## 9:      ARM_CIRCUM 19.01839 Confirmed
## 10:   WAIST_CIRCUMF 19.89443 Confirmed
## 11: SAG ABDOMINAL_DIA 18.52763 Confirmed
```

```
# Plot variable importance
plot(boruta_output, cex.axis=.7, las=2, xlab="", main="Variable Importance")
```

## Variable Importance



```
## Remove SEQN from df

Final_DF_1 <- subset(Final_DF , select = -c(SEQN))

#####
#Fitting Models
#####

#####
##Splitting Data
#####

dat_part <- resample_partition(Final_DF_1,
                               c(train = 0.8,
                                 test = 0.2))

train <- as.tibble(dat_part$train)
test <- as.tibble(dat_part$test)

#####
#Oversampling
#####

train.rose <- ovun.sample(Diabetes ~ . , data = train, method = "over",
```



```
seed = 1)$data

table(train.rose$Diabetes)
```

```
##
## NO YES
## 6320 6268
```

```
table(train$Diabetes)
```

```
##
## NO YES
## 6320 471
```

```
summary(train)
```

```
## Diabetes      AGE      BP_SYS      BP_DIA      WEIGHT
## NO :6320  Min.   : 5.00  Min.   : 66.0  Min.   : 0.00  Min.   : 13.10
## YES: 471  1st Qu.:15.00  1st Qu.:108.0  1st Qu.: 60.00  1st Qu.: 53.60
##           Median :34.00  Median :116.0  Median : 66.00  Median : 69.80
##           Mean   :36.34  Mean   :117.7  Mean   : 65.69  Mean   : 70.58
##           3rd Qu.:56.00  3rd Qu.:124.0  3rd Qu.: 74.00  3rd Qu.: 86.75
##           Max.   :80.00  Max.   :228.0  Max.   :122.00  Max.   :222.60
##      HEIGHT      BMI      UPPER_LEG_LENGTH UPPER_ARM_LENGTH
## Min.   : 90.6  Min.   :12.10  Min.   :24.40  Min.   :17.00
## 1st Qu.:154.1  1st Qu.:21.00  1st Qu.:36.40  1st Qu.:33.50
## Median :163.2  Median :25.50  Median :38.60  Median :36.00
## Mean   :160.5  Mean   :26.43  Mean   :38.59  Mean   :35.43
## 3rd Qu.:171.9  3rd Qu.:30.60  3rd Qu.:41.00  3rd Qu.:38.40
## Max.   :202.6  Max.   :82.90  Max.   :51.90  Max.   :47.90
##      ARM_CIRCUM  WAIST_CIRCUMF  SAG ABDOMINAL_DIA
## Min.   :13.90  Min.   : 41.50  Min.   :10.10
## 1st Qu.:26.20  1st Qu.: 76.20  1st Qu.:17.90
## Median :30.50  Median : 89.80  Median :20.70
## Mean   :30.34  Mean   : 90.14  Mean   :21.07
## 3rd Qu.:34.50  3rd Qu.:103.30  3rd Qu.:23.70
## Max.   :59.40  Max.   :172.50  Max.   :40.10
```

```
summary(train.rose)
```

```
## Diabetes      AGE      BP_SYS      BP_DIA      WEIGHT
## NO :6320  Min.   : 5.00  Min.   : 66.0  Min.   : 0.00  Min.   : 13.10
## YES:6268  1st Qu.:29.00  1st Qu.:112.0  1st Qu.: 60.00  1st Qu.: 65.00
##           Median :51.00  Median :118.0  Median : 66.00  Median : 78.40
##           Mean   :47.33  Mean   :123.3  Mean   : 67.01  Mean   : 80.41
##           3rd Qu.:66.00  3rd Qu.:134.0  3rd Qu.: 76.00  3rd Qu.: 95.70
##           Max.   :80.00  Max.   :228.0  Max.   :122.00  Max.   :222.60
##      HEIGHT      BMI      UPPER_LEG_LENGTH UPPER_ARM_LENGTH
## Min.   : 90.6  Min.   :12.1  Min.   :24.4  Min.   :17.00
## 1st Qu.:156.6  1st Qu.:23.9  1st Qu.:35.7  1st Qu.:34.80
## Median :164.7  Median :28.4  Median :38.6  Median :36.70
```

```
## Mean :163.4 Mean :29.4 Mean :38.2 Mean :36.47
## 3rd Qu.:172.9 3rd Qu.:33.7 3rd Qu.:40.7 3rd Qu.:39.00
## Max. :202.6 Max. :82.9 Max. :51.9 Max. :47.90
## ARM_CIRCUM WAIST_CIRCUMF SAG ABDOMINAL DIA
## Min. :13.90 Min. : 41.50 Min. :10.10
## 1st Qu.:29.20 1st Qu.: 87.70 1st Qu.:20.40
## Median :32.60 Median : 99.05 Median :22.50
## Mean :32.62 Mean : 99.28 Mean :23.28
## 3rd Qu.:36.40 3rd Qu.:112.60 3rd Qu.:26.60
## Max. :59.40 Max. :172.50 Max. :40.10
```

```
#####
#Undersampling
#####
```

```
train.rose_under <- ovun.sample(Diabetes~. , data = train, method = "under",
                                N = 964, seed = 1)$data
```

```
table(train.rose_under$Diabetes)
```

```
##
## NO YES
## 493 471
```

```
table(train$Diabetes)
```

```
##
## NO YES
## 6320 471
```

```
summary(train)
```

```
## Diabetes AGE BP_SYS BP_DIA WEIGHT
## NO :6320 Min. : 5.00 Min. : 66.0 Min. : 0.00 Min. : 13.10
## YES: 471 1st Qu.:15.00 1st Qu.:108.0 1st Qu.: 60.00 1st Qu.: 53.60
## Median :34.00 Median :116.0 Median : 66.00 Median : 69.80
## Mean :36.34 Mean :117.7 Mean : 65.69 Mean : 70.58
## 3rd Qu.:56.00 3rd Qu.:124.0 3rd Qu.: 74.00 3rd Qu.: 86.75
## Max. :80.00 Max. :228.0 Max. :122.00 Max. :222.60
## HEIGHT BMI UPPER_LEG_LENGTH UPPER_ARM_LENGTH
## Min. : 90.6 Min. :12.10 Min. :24.40 Min. :17.00
## 1st Qu.:154.1 1st Qu.:21.00 1st Qu.:36.40 1st Qu.:33.50
## Median :163.2 Median :25.50 Median :38.60 Median :36.00
## Mean :160.5 Mean :26.43 Mean :38.59 Mean :35.43
## 3rd Qu.:171.9 3rd Qu.:30.60 3rd Qu.:41.00 3rd Qu.:38.40
## Max. :202.6 Max. :82.90 Max. :51.90 Max. :47.90
## ARM_CIRCUM WAIST_CIRCUMF SAG ABDOMINAL DIA
## Min. :13.90 Min. : 41.50 Min. :10.10
## 1st Qu.:26.20 1st Qu.: 76.20 1st Qu.:17.90
## Median :30.50 Median : 89.80 Median :20.70
## Mean :30.34 Mean : 90.14 Mean :21.07
## 3rd Qu.:34.50 3rd Qu.:103.30 3rd Qu.:23.70
## Max. :59.40 Max. :172.50 Max. :40.10
```

```
summary(train.rose_under)
```

```
## Diabetes      AGE      BP_SYS      BP_DIA      WEIGHT
## NO :493   Min.    : 5.00   Min.    : 72.0   Min.    : 0.00   Min.    : 14.2
## YES:471   1st Qu.:30.00   1st Qu.:114.0   1st Qu.: 60.00   1st Qu.: 65.1
##           Median :52.00   Median :118.0   Median : 66.00   Median : 77.5
##           Mean    :47.74   Mean    :123.6   Mean    : 66.96   Mean    : 79.9
##           3rd Qu.:66.00   3rd Qu.:134.0   3rd Qu.: 74.00   3rd Qu.: 95.2
##           Max.    :80.00   Max.    :204.0   Max.    :112.00   Max.    :195.4
##      HEIGHT      BMI      UPPER_LEG_LENGTH UPPER_ARM_LENGTH
## Min.    :103.1   Min.    :12.60   Min.    :26.00   Min.    :20.10
## 1st Qu.:156.6   1st Qu.:24.15   1st Qu.:36.00   1st Qu.:34.90
## Median :164.4   Median :28.30   Median :38.60   Median :36.60
## Mean    :163.2   Mean    :29.28   Mean    :38.26   Mean    :36.45
## 3rd Qu.:172.4   3rd Qu.:33.42   3rd Qu.:40.50   3rd Qu.:39.00
## Max.    :202.6   Max.    :71.50   Max.    :51.90   Max.    :46.80
##  ARM_CIRCUM  WAIST_CIRCUMF  SAG ABDOMINAL_DIA
## Min.    :14.20   Min.    : 41.50   Min.    :11.90
## 1st Qu.:29.20   1st Qu.: 87.95   1st Qu.:20.50
## Median :32.40   Median : 98.50   Median :22.40
## Mean    :32.52   Mean    : 99.07   Mean    :23.28
## 3rd Qu.:36.20   3rd Qu.:112.03   3rd Qu.:26.32
## Max.    :55.70   Max.    :161.00   Max.    :40.10
```

```
#####
#Logistic Regression
#####
```

```
#####
# Normal Sample
#####
```

```
log_train <- train
log_test  <- test
```

```
fit_log_normal <- glm(Diabetes ~ . , family = binomial(link = "logit"),
                      data = log_train)
```

```
summary(fit_log_normal)
```

```
##
## Call:
## glm(formula = Diabetes ~ . , family = binomial(link = "logit"),
##      data = log_train)
##
## Deviance Residuals:
##      Min       1Q   Median       3Q      Max
## -1.6246  -0.3633  -0.1743  -0.0768   3.2927
##
## Coefficients:
##              Estimate Std. Error z value Pr(>|z|)
## (Intercept)   -11.532152    2.592321  -4.449 8.64e-06 ***
```

```
## AGE                0.042433    0.003771   11.253   < 2e-16 ***
## BP_SYS             0.011741    0.003109    3.776 0.000159 ***
## BP_DIA            -0.008955    0.004095   -2.187 0.028770 *
## WEIGHT            -0.007088    0.014427   -0.491 0.623196
## HEIGHT             0.037294    0.016703    2.233 0.025567 *
## BMI                0.011801    0.039505    0.299 0.765159
## UPPER_LEG_LENGTH  -0.100827    0.022078   -4.567 4.95e-06 ***
## UPPER_ARM_LENGTH  -0.028332    0.033103   -0.856 0.392070
## ARM_CIRCUM         0.033358    0.022002    1.516 0.129485
## WAIST_CIRCUMF      0.004263    0.009240    0.461 0.644549
## SAG_ABDOMINAL_DIA  0.146868    0.028026    5.240 1.60e-07 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##      Null deviance: 3422.3  on 6790  degrees of freedom
## Residual deviance: 2538.0  on 6779  degrees of freedom
## AIC: 2562
##
## Number of Fisher Scoring iterations: 7
```

```
log_test <-
  log_test %>%
  add_predictions(fit_log_normal, type = "response") %>%
  mutate(pred_dia = ifelse(pred > 0.12,
                           "YES", "NO"),
         correct = Diabetes == pred_dia )

table(log_test$Diabetes, log_test$pred_dia)[,1:2] # Confusion Matrix
```

```
##
##      NO  YES
## NO 1343 222
## YES  60  73
```

```
# Sensitivity = 0.6885246
# Specificity = 0.8261421
# Accuracy = 0.8162544

#####
# Oversampling
#####

log_train_over <- train.rose
log_test_over <- test

fit_log_over <- glm(Diabetes ~ . , family = binomial(link = "logit"),
                   data = log_train_over)
summary(fit_log_over)
```

```
##
## Call:
## glm(formula = Diabetes ~ ., family = binomial(link = "logit"),
##      data = log_train_over)
##
## Deviance Residuals:
##      Min       1Q   Median       3Q      Max
## -2.78023  -0.61224  -0.08069   0.74336   2.58414
##
## Coefficients:
##              Estimate Std. Error z value Pr(>|z|)
## (Intercept)  -10.869641   1.318447  -8.244 < 2e-16 ***
## AGE           0.049885   0.001627  30.653 < 2e-16 ***
## BP_SYS       0.011280   0.001573   7.170 7.49e-13 ***
## BP_DIA      -0.004989   0.001977  -2.523  0.0116 *
## WEIGHT      -0.012812   0.007715  -1.661  0.0968 .
## HEIGHT       0.047557   0.008503   5.593 2.24e-08 ***
## BMI          0.052907   0.021286   2.486  0.0129 *
## UPPER_LEG_LENGTH -0.118967  0.009697 -12.268 < 2e-16 ***
## UPPER_ARM_LENGTH -0.025298  0.015423  -1.640  0.1009
## ARM_CIRCUM    0.015696  0.009922   1.582  0.1137
## WAIST_CIRCUMF  0.008598  0.003783   2.273  0.0230 *
## SAG_ABDOMINAL_DIA 0.129030  0.012592  10.247 < 2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##      Null deviance: 17450  on 12587  degrees of freedom
## Residual deviance: 11046  on 12576  degrees of freedom
## AIC: 11070
##
## Number of Fisher Scoring iterations: 6
```

```
log_test_over <-
  log_test_over %>%
  add_predictions(fit_log_over, type = "response") %>%
  mutate(pred_dia = ifelse(pred > 0.5,
                           "YES", "NO"),
         correct = Diabetes == pred_dia )

table(log_test_over$Diabetes, log_test_over$pred_dia)[,1:2] # Confusion Matrix
```

```
##
##      NO  YES
## NO  1171 394
## YES   26 107
```

```
#sensitivity = 0.852459
```

```
#specificity = 0.732868
```

```
#accuracy = 0.0.7414065

#####
# Undersampling
#####

log_train_under <- train.rose_under
log_test_under <- test

fit_log_under <- glm(Diabetes ~ . , family = binomial(link = "logit"),
                     data = log_train_under)
summary(fit_log_under)
```

```
##
## Call:
## glm(formula = Diabetes ~ . , family = binomial(link = "logit"),
##      data = log_train_under)
##
## Deviance Residuals:
##      Min       1Q   Median       3Q      Max
## -2.8199  -0.6447  -0.0817   0.7464   2.6018
##
## Coefficients:
##              Estimate Std. Error z value Pr(>|z|)
## (Intercept)  -11.843795   3.852714  -3.074  0.00211 **
## AGE           0.044722   0.005878   7.608 2.78e-14 ***
## BP_SYS       0.010542   0.005626   1.874  0.06096 .
## BP_DIA      -0.003661   0.007015  -0.522  0.60172
## WEIGHT      -0.015189   0.021990  -0.691  0.48973
## HEIGHT       0.061413   0.025104   2.446  0.01443 *
## BMI          0.075009   0.058514   1.282  0.19988
## UPPER_LEG_LENGTH -0.151973  0.036268  -4.190 2.79e-05 ***
## UPPER_ARM_LENGTH -0.025603  0.055616  -0.460  0.64527
## ARM_CIRCUM    0.016594  0.034555   0.480  0.63108
## WAIST_CIRCUMF  0.004152  0.013954   0.298  0.76606
## SAG ABDOMINAL_DIA 0.134150  0.046995   2.855  0.00431 **
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##      Null deviance: 1335.89  on 963  degrees of freedom
## Residual deviance:  856.14  on 952  degrees of freedom
## AIC: 880.14
##
## Number of Fisher Scoring iterations: 6
```

```
log_test_under <-
  log_test_under %>%
  add_predictions(fit_log_under, type = "response") %>%
  mutate(pred_dia = ifelse(pred > 0.5,
                           "YES", "NO"),
         correct = Diabetes == pred_dia )
```

```
table(log_test_under$Diabetes, log_test_under$pred_dia)[,1:2] # Confusion Matrix
```

```
##
##          NO  YES
##   NO  1186  379
##   YES   30  103
```

```
#sensitivity = 0.8852459
```

```
#specificity = 0.7277919
```

```
#accuracy = 0.7391048
```

```
#####
##Random Forests
#####
```

```
## Normal Sample
```

```
summary(train)
```

```
## Diabetes      AGE      BP_SYS      BP_DIA      WEIGHT
## NO :6320  Min.   : 5.00  Min.   : 66.0  Min.   : 0.00  Min.   : 13.10
## YES: 471  1st Qu.:15.00  1st Qu.:108.0  1st Qu.: 60.00  1st Qu.: 53.60
##          Median :34.00  Median :116.0  Median : 66.00  Median : 69.80
##          Mean   :36.34  Mean   :117.7  Mean   : 65.69  Mean   : 70.58
##          3rd Qu.:56.00  3rd Qu.:124.0  3rd Qu.: 74.00  3rd Qu.: 86.75
##          Max.   :80.00  Max.   :228.0  Max.   :122.00  Max.   :222.60
##      HEIGHT      BMI      UPPER_LEG_LENGTH  UPPER_ARM_LENGTH
## Min.   : 90.6  Min.   :12.10  Min.   :24.40  Min.   :17.00
## 1st Qu.:154.1  1st Qu.:21.00  1st Qu.:36.40  1st Qu.:33.50
## Median :163.2  Median :25.50  Median :38.60  Median :36.00
## Mean   :160.5  Mean   :26.43  Mean   :38.59  Mean   :35.43
## 3rd Qu.:171.9  3rd Qu.:30.60  3rd Qu.:41.00  3rd Qu.:38.40
## Max.   :202.6  Max.   :82.90  Max.   :51.90  Max.   :47.90
##  ARM_CIRCUM  WAIST_CIRCUMF  SAG_ABDOMINAL_DIA
## Min.   :13.90  Min.   : 41.50  Min.   :10.10
## 1st Qu.:26.20  1st Qu.: 76.20  1st Qu.:17.90
## Median :30.50  Median : 89.80  Median :20.70
## Mean   :30.34  Mean   : 90.14  Mean   :21.07
## 3rd Qu.:34.50  3rd Qu.:103.30  3rd Qu.:23.70
## Max.   :59.40  Max.   :172.50  Max.   :40.10
```

```
model_normal_rf <- randomForest(Diabetes ~ ., data = train, ntree = 1000,
                                proximity = TRUE) # Prox = true returns
                                                    #proximity matrix
```

```
model_normal_rf
```

```
##
## Call:
## randomForest(formula = Diabetes ~ ., data = train, ntree = 1000,      proximity = TRUE)
##           Type of random forest: classification
##           Number of trees: 1000
## No. of variables tried at each split: 3
##
##           OOB estimate of  error rate: 7.16%
## Confusion matrix:
##           NO YES class.error
## NO  6289  31 0.004905063
## YES  455  16 0.966029724
```

```
prediction <- as.data.frame((predict(model_normal_rf, newdata = test)))

confusionMatrix((predict(model_normal_rf, newdata = test)), test$Diabetes,
                positive = "YES") # Bad Sensitivity but good specificity
```

```
## Confusion Matrix and Statistics
##
##           Reference
## Prediction  NO  YES
##           NO 1562 128
##           YES   3   5
##
##           Accuracy : 0.9229
##           95% CI : (0.9091, 0.9351)
## No Information Rate : 0.9217
## P-Value [Acc > NIR] : 0.4511
##
##           Kappa : 0.0626
##
## Mcnemar's Test P-Value : <2e-16
##
##           Sensitivity : 0.037594
##           Specificity : 0.998083
##           Pos Pred Value : 0.625000
##           Neg Pred Value : 0.924260
##           Prevalence : 0.078327
##           Detection Rate : 0.002945
## Detection Prevalence : 0.004711
##           Balanced Accuracy : 0.517839
##
##           'Positive' Class : YES
##
```

```
#sensitivity = 0.049180
#specificity = 0.995558
#accuracy = 0.9276
# save the model to disk
```



```
saveRDS(model_normal_rf, "C:/Semester 2/Intro to Data Mining and Processing/Project/national-health-and
```

```
## Oversampling
```

```
model <- randomForest(Diabetes ~ ., data = train.rose, ntree = 1000,  
                      proximity = TRUE) # Prox = true returns proximity matrix
```

```
model
```

```
##
```

```
## Call:
```

```
## randomForest(formula = Diabetes ~ ., data = train.rose, ntree = 1000, proximity = TRUE)
```

```
## Type of random forest: classification
```

```
## Number of trees: 1000
```

```
## No. of variables tried at each split: 3
```

```
##
```

```
## OOB estimate of error rate: 0.91%
```

```
## Confusion matrix:
```

```
## NO YES class.error
```

```
## NO 6206 114 0.01803797
```

```
## YES 0 6268 0.00000000
```

```
confusionMatrix((predict(model, newdata = test)), test$Diabetes,  
                positive = "YES") # Bad Sensitivity but good specificity
```

```
## Confusion Matrix and Statistics
```

```
##
```

```
## Reference
```

```
## Prediction NO YES
```

```
## NO 1541 116
```

```
## YES 24 17
```

```
##
```

```
## Accuracy : 0.9176
```

```
## 95% CI : (0.9034, 0.9302)
```

```
## No Information Rate : 0.9217
```

```
## P-Value [Acc > NIR] : 0.7531
```

```
##
```

```
## Kappa : 0.1646
```

```
##
```

```
## McNemar's Test P-Value : 1.461e-14
```

```
##
```

```
## Sensitivity : 0.12782
```

```
## Specificity : 0.98466
```

```
## Pos Pred Value : 0.41463
```

```
## Neg Pred Value : 0.92999
```

```
## Prevalence : 0.07833
```

```
## Detection Rate : 0.01001
```

```
## Detection Prevalence : 0.02415
```

```
## Balanced Accuracy : 0.55624
```

```
##
```

```
## 'Positive' Class : YES
```

```
##
```

```

#sensitivity = 0.07377

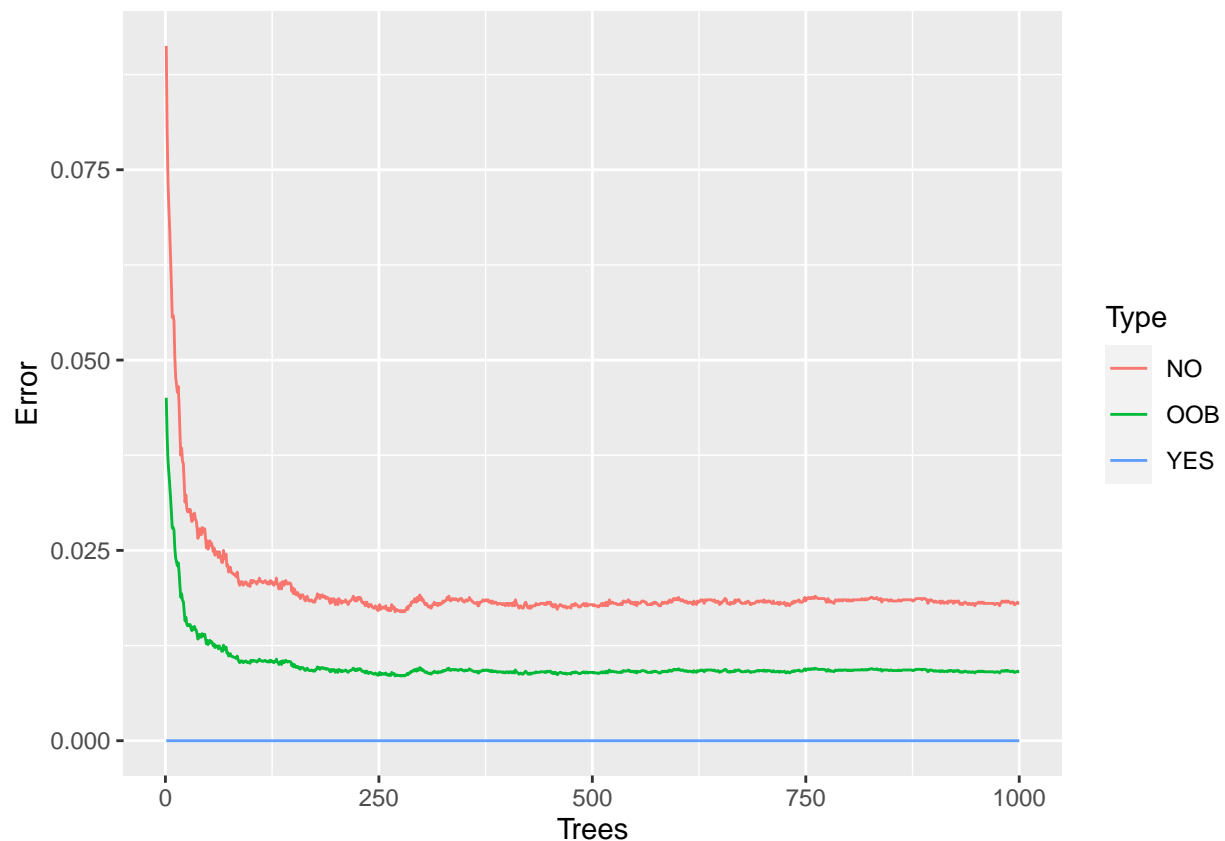
#specificity = 0.98731

#accuracy = 0.9217

oob.error.data <- data.frame(
  Trees = rep(1:nrow(model$err.rate), times=3),
  Type = rep(c("OOB", "NO", "YES"), each = nrow(model$err.rate)),
  Error=c(model$err.rate[, "OOB"],
          model$err.rate[, "NO"],
          model$err.rate[, "YES"]))

ggplot(data = oob.error.data, aes(x=Trees, y=Error)) +
  geom_line(aes(color = Type))

```



```

oob.values <- vector(length = 12)
for(i in 1:12){
  temp.model <- randomForest(Diabetes~., data = train.rose, mtry=i,
                             ntree = 1000)
  oob.values[i] <- temp.model$err.rate[nrow(temp.model$err.rate),1]
}

oob.values # mtry = 1 gave the best oob

```

```
## [1] 0.005878615 0.008102955 0.008817922 0.010168414 0.010962822 0.012313314
## [7] 0.012472196 0.013504925 0.015729266 0.016523673 0.017556403 0.017635844
```

```
# save the model to disk
```

```
saveRDS(model, "C:/Semester 2/Intro to Data Mining and Processing/Project/national-health-and-nutrition")
```

```
#code to load model
```

```
#super_model <- readRDS("")
```

```
#print(super_model)
```

```
## Undersampling
```

```
model_under <- randomForest(Diabetes ~ ., data = train.rose_under, ntree = 1000,
                             proximity = TRUE) # Prox = true returns proximity
                                                #matrix
```

```
model_under
```

```
##
```

```
## Call:
```

```
## randomForest(formula = Diabetes ~ ., data = train.rose_under, ntree = 1000, proximity = TRUE)
```

```
## Type of random forest: classification
```

```
## Number of trees: 1000
```

```
## No. of variables tried at each split: 3
```

```
##
```

```
## OOB estimate of error rate: 20.75%
```

```
## Confusion matrix:
```

```
## NO YES class.error
```

```
## NO 364 129 0.2616633
```

```
## YES 71 400 0.1507431
```

```
confusionMatrix(predict(model_under, test), test$Diabetes, positive = "YES")
```

```
## Confusion Matrix and Statistics
```

```
##
```

```
## Reference
```

```
## Prediction NO YES
```

```
## NO 1160 31
```

```
## YES 405 102
```

```
##
```

```
## Accuracy : 0.7432
```

```
## 95% CI : (0.7217, 0.7639)
```

```
## No Information Rate : 0.9217
```

```
## P-Value [Acc > NIR] : 1
```

```
##
```

```
## Kappa : 0.2222
```

```
##
```

```
## McNemar's Test P-Value : <2e-16
```

```
##
```

```
## Sensitivity : 0.76692
```

```
## Specificity : 0.74121
```

```
##          Pos Pred Value : 0.20118
##          Neg Pred Value : 0.97397
##          Prevalence : 0.07833
##          Detection Rate : 0.06007
##          Detection Prevalence : 0.29859
##          Balanced Accuracy : 0.75407
##
##          'Positive' Class : YES
##
```

```
# Good sensitivity
```

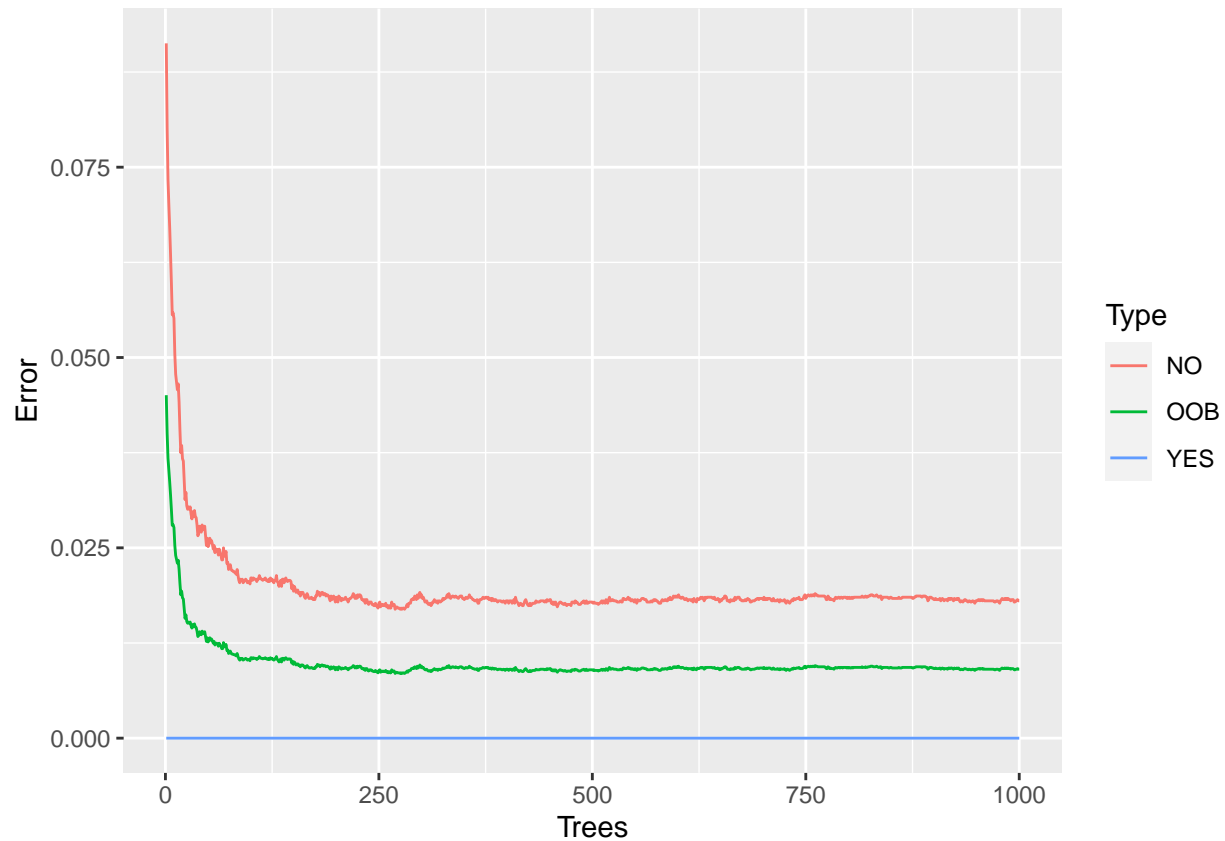
```
#sensitivity = Sensitivity : 0.90164
```

```
#specificity = 0.71447
```

```
#accuracy = 0.7279
```

```
oob.error.data <- data.frame(
  Trees = rep(1:nrow(model$err.rate), times=3),
  Type = rep(c("OOB", "NO", "YES"), each = nrow(model$err.rate)),
  Error=c(model$err.rate[, "OOB"],
          model$err.rate[, "NO"],
          model$err.rate[, "YES"]))

ggplot(data = oob.error.data, aes(x=Trees, y=Error)) +
  geom_line(aes(color = Type))
```



```
oob.values <- vector(length = 12)
for(i in 1:12){
  temp.model <- randomForest(Diabetes~., data = train.rose, mtry=i,
                             ntree = 1000)
  oob.values[i] <- temp.model$err.rate[nrow(temp.model$err.rate),1]
} ##ntrees = 630 seems to be the best

oob.values # mtry = 1 gave the best oob
```

```
## [1] 0.006037496 0.007785192 0.009056244 0.010168414 0.010565618 0.012233874
## [7] 0.013584366 0.013902129 0.014617096 0.015252622 0.017079759 0.017079759
```

```
saveRDS(model_under, "C:/Semester 2/Intro to Data Mining and Processing/Project/national-health-and-nutrition-survey-iv")
```

```
#tuning the model
```

```
model_under_tuned <- randomForest(Diabetes ~ ., data = train.rose_under,
                                   ntree = 1000, proximity = TRUE)
# Prox = true returns proximity matrix
model_under_tuned
```

```
##
## Call:
## randomForest(formula = Diabetes ~ ., data = train.rose_under,          ntree = 1000, proximity = TRUE)
```

```
##           Type of random forest: classification
##           Number of trees: 1000
## No. of variables tried at each split: 3
##
##           OOB estimate of  error rate: 20.85%
## Confusion matrix:
##           NO YES class.error
## NO   365 128   0.2596349
## YES   73 398   0.1549894
```

```
confusionMatrix(predict(model_under_tuned, test), test$Diabetes,
                  positive = "YES")
```

```
## Confusion Matrix and Statistics
##
##           Reference
## Prediction  NO  YES
##           NO 1160  27
##           YES 405 106
##
##           Accuracy : 0.7456
##           95% CI : (0.7242, 0.7662)
##           No Information Rate : 0.9217
##           P-Value [Acc > NIR] : 1
##
##           Kappa : 0.234
##
## Mcnemar's Test P-Value : <2e-16
##
##           Sensitivity : 0.79699
##           Specificity : 0.74121
##           Pos Pred Value : 0.20744
##           Neg Pred Value : 0.97725
##           Prevalence : 0.07833
##           Detection Rate : 0.06243
##           Detection Prevalence : 0.30094
##           Balanced Accuracy : 0.76910
##
##           'Positive' Class : YES
##
```

```
#Sensitivity : 0.90984
#Specificity : 0.71256
#Accuracy : 0.7267
```

```
#####
##xgboost
#####

#####
##NORMAL DATA
#####
```

```

df_train_0 <- train.rose
df_test_0 <- test

setDT(df_train_0)
setDT(df_test_0)

# Using one hot encoding

labels_0 <- df_train_0$Diabetes
ts_label_0 <- df_test_0$Diabetes

new_tr_0 <- model.matrix(~.+0, data = df_train_0[, -c("Diabetes"), with=F])
new_ts_0 <- model.matrix(~.+0, data = df_test_0[, -c("Diabetes"), with=F])

#convert factor to numeric
labels_0 <- as.numeric(labels_0)-1
ts_label_0 <- as.numeric(ts_label_0)-1

dtrain_0 <- xgb.DMatrix(data = new_tr_0, label = labels_0)
dtest_0 <- xgb.DMatrix(data = new_ts_0, label=ts_label_0)

#default parameters
params <- list(booster = "gbtree", objective = "binary:logistic", eta=0.3,
               gamma=0, max_depth=6, min_child_weight=1, subsample=1,
               colsample_bytree=1)

xgbcv_0 <- xgb.cv( params = params, data = dtrain_0, nrounds = 100, nfold = 5,
                  showsd = T, stratified = T, print.every.n = 10,
                  early.stop.round = 20, maximize = F)

```

```

## [1] train-error:0.152546+0.002483 test-error:0.160630+0.006881
## Multiple eval metrics are present. Will use test_error for early stopping.
## Will train until test_error hasn't improved in 20 rounds.
##
## [11] train-error:0.100652+0.002405 test-error:0.113283+0.005454
## [21] train-error:0.073006+0.003341 test-error:0.093503+0.005552
## [31] train-error:0.048677+0.002178 test-error:0.073881+0.006044
## [41] train-error:0.028837+0.001852 test-error:0.059104+0.004167
## [51] train-error:0.018529+0.001077 test-error:0.052431+0.003392
## [61] train-error:0.011618+0.000785 test-error:0.045758+0.003083
## [71] train-error:0.006117+0.000324 test-error:0.039641+0.002927
## [81] train-error:0.003555+0.000291 test-error:0.035669+0.002546
## [91] train-error:0.001966+0.000368 test-error:0.032412+0.002671
## [100] train-error:0.001231+0.000342 test-error:0.031618+0.002422

```

```

# lowest in 100th iteration

```

```

#first default - model training

```

```

xgb1_0 <- xgb.train (params = params, data = dtrain_0, nrounds = 79, watchlist =
                    list(val=dtest_0,train=dtrain_0), print.every.n = 10,
                    early.stop.round = 10, maximize = F ,
                    eval_metric = "error")

```

```
## [17:19:17] WARNING: amalgamation/../src/learner.cc:480:
## Parameters: { early_stop_round, print_every_n } might not be used.
##
## This may not be accurate due to some parameters are only used in language bindings but
## passed down to XGBoost core. Or some parameters are not used but slip through this
## verification. Please open an issue if you find above cases.
##
##
## [1] val-error:0.272674 train-error:0.151970
## Multiple eval metrics are present. Will use train_error for early stopping.
## Will train until train_error hasn't improved in 10 rounds.
##
## [11] val-error:0.220848 train-error:0.100493
## [21] val-error:0.192580 train-error:0.080473
## [31] val-error:0.163722 train-error:0.049571
## [41] val-error:0.146054 train-error:0.033445
## [51] val-error:0.133098 train-error:0.020416
## [61] val-error:0.118963 train-error:0.012313
## [71] val-error:0.108363 train-error:0.006435
## [79] val-error:0.107774 train-error:0.004131
```

#### *#model prediction*

```
xgbpred_0 <- predict(xgb1_0,dtest_0)
xgbpred_0 <- ifelse(xgbpred_0 > 0.5,'YES','NO')
xgbpred1_0 <- as.factor(xgbpred_0)
confusionMatrix(xgbpred1_0, df_test_0$Diabetes, positive = "YES")
```

#### ## Confusion Matrix and Statistics

```
##
##           Reference
## Prediction  NO  YES
##           NO 1469  86
##           YES  96  47
##
##           Accuracy : 0.8928
##           95% CI : (0.8771, 0.9071)
##           No Information Rate : 0.9217
##           P-Value [Acc > NIR] : 1.0000
##
##           Kappa : 0.2823
##
## Mcnemar's Test P-Value : 0.5047
##
##           Sensitivity : 0.35338
##           Specificity : 0.93866
##           Pos Pred Value : 0.32867
##           Neg Pred Value : 0.94469
##           Prevalence : 0.07833
##           Detection Rate : 0.02768
##           Detection Prevalence : 0.08422
##           Balanced Accuracy : 0.64602
##
##           'Positive' Class : YES
##
```



```

# Bad sensitivity

#Sensitivity : 0.27049
#Specificity : 0.93147
#Accuracy : 0.884

#####
##OVERSAMPLED DATA
#####
df_train <- train.rose
df_test <- test

setDT(df_train)
setDT(df_test)

# Using one hot encoding

labels <- df_train$Diabetes
ts_label <- df_test$Diabetes

new_tr <- model.matrix(~.+0, data = df_train[, -c("Diabetes"), with=F])
new_ts <- model.matrix(~.+0, data = df_test[, -c("Diabetes"), with=F])

#convert factor to numeric
labels <- as.numeric(labels)-1
ts_label <- as.numeric(ts_label)-1

dtrain <- xgb.DMatrix(data = new_tr, label = labels)
dtest <- xgb.DMatrix(data = new_ts, label=ts_label)

#default parameters
params <- list(booster = "gbtree", objective = "binary:logistic", eta=0.3,
              gamma=0, max_depth=6, min_child_weight=1, subsample=1,
              colsample_bytree=1)

xgbcv <- xgb.cv( params = params, data = dtrain, nrounds = 100, nfold = 5,
                showsd = T, stratified = T, print.every.n = 10,
                early.stop.round = 20, maximize = F)

## [1] train-error:0.155009+0.003644 test-error:0.163012+0.004003
## Multiple eval metrics are present. Will use test_error for early stopping.
## Will train until test_error hasn't improved in 20 rounds.
##
## [11] train-error:0.098526+0.003562 test-error:0.113758+0.007134
## [21] train-error:0.074198+0.001372 test-error:0.093104+0.002586
## [31] train-error:0.046751+0.002370 test-error:0.071893+0.005021
## [41] train-error:0.028539+0.001770 test-error:0.058865+0.004396
## [51] train-error:0.018212+0.000991 test-error:0.051954+0.005838
## [61] train-error:0.010605+0.001297 test-error:0.044645+0.005286
## [71] train-error:0.005680+0.000484 test-error:0.038846+0.004493
## [81] train-error:0.002900+0.000500 test-error:0.035907+0.004191
## [91] train-error:0.001708+0.000346 test-error:0.034000+0.003694

```

```
## [100]      train-error:0.001251+0.000204      test-error:0.033206+0.003949
```

```
# lowest in 100th iteration
```

```
#first default - model training
```

```
xgb1 <- xgb.train (params = params, data = dtrain, nrounds = 79, watchlist =  
                  list(val=dtest,train=dtrain), print.every.n = 10,  
                  early.stop.round = 10, maximize = F , eval_metric = "error")
```

```
## [17:19:21] WARNING: amalgamation/./src/learner.cc:480:
```

```
## Parameters: { early_stop_round, print_every_n } might not be used.
```

```
##
```

```
## This may not be accurate due to some parameters are only used in language bindings but  
## passed down to XGBoost core. Or some parameters are not used but slip through this  
## verification. Please open an issue if you find above cases.
```

```
##
```

```
##
```

```
## [1] val-error:0.272674 train-error:0.151970
```

```
## Multiple eval metrics are present. Will use train_error for early stopping.
```

```
## Will train until train_error hasn't improved in 10 rounds.
```

```
##
```

```
## [11] val-error:0.220848 train-error:0.100493
```

```
## [21] val-error:0.192580 train-error:0.080473
```

```
## [31] val-error:0.163722 train-error:0.049571
```

```
## [41] val-error:0.146054 train-error:0.033445
```

```
## [51] val-error:0.133098 train-error:0.020416
```

```
## [61] val-error:0.118963 train-error:0.012313
```

```
## [71] val-error:0.108363 train-error:0.006435
```

```
## [79] val-error:0.107774 train-error:0.004131
```

```
#model prediction
```

```
xgbpred <- predict (xgb1,dtest)
```

```
xgbpred <- ifelse (xgbpred > 0.5,'YES','NO')
```

```
xgbpred1 <- as.factor(xgbpred)
```

```
confusionMatrix(xgbpred1 , df_test$Diabetes, positive = "YES")
```

```
## Confusion Matrix and Statistics
```

```
##
```

```
##           Reference
```

```
## Prediction  NO  YES
```

```
##           NO 1469  86
```

```
##           YES  96  47
```

```
##
```

```
##           Accuracy : 0.8928
```

```
##           95% CI : (0.8771, 0.9071)
```

```
## No Information Rate : 0.9217
```

```
## P-Value [Acc > NIR] : 1.0000
```

```
##
```

```
##           Kappa : 0.2823
```

```
##
```

```
## McNemar's Test P-Value : 0.5047
```

```
##
```

```
##           Sensitivity : 0.35338
```

```
##           Specificity : 0.93866
##           Pos Pred Value : 0.32867
##           Neg Pred Value : 0.94469
##           Prevalence : 0.07833
##           Detection Rate : 0.02768
##           Detection Prevalence : 0.08422
##           Balanced Accuracy : 0.64602
##
##           'Positive' Class : YES
##
```

```
# very low sensitivity but high specificity
```

```
#Sensitivity : 0.27049
#Specificity : 0.93147
#Accuracy : 0.884
```

```
#####
##UNDERSAMPLED DATA
#####
```

```
df_train1 <- train.rose_under
df_test1 <- test
```

```
setDT(df_train1)
setDT(df_test1)
```

```
# Using one hot encoding
```

```
labels1 <- df_train1$Diabetes
ts_label1 <- df_test1$Diabetes
```

```
new_tr1 <- model.matrix(~.+0, data = df_train1[, -c("Diabetes"), with=F])
```

```
new_ts1 <- model.matrix(~.+0, data = df_test1[, -c("Diabetes"), with=F])
```

```
#convert factor to numeric
```

```
labels1 <- as.numeric(labels1)-1
ts_label1 <- as.numeric(ts_label1)-1
```

```
dtrain1 <- xgb.DMatrix(data = new_tr1, label = labels1)
```

```
dtest1 <- xgb.DMatrix(data = new_ts1, label=ts_label1)
```

```
#default parameters
```

```
xgbcv1 <- xgb.cv( params = params, data = dtrain1, nrounds = 100,
                  nfold = 5, showsd = T, stratified = T, print.every.n = 10,
                  early.stop.round = 20, maximize = F)
```

```
## [1] train-error:0.148340+0.010841 test-error:0.243771+0.036014
## Multiple eval metrics are present. Will use test_error for early stopping.
## Will train until test_error hasn't improved in 20 rounds.
##
## [11] train-error:0.068983+0.012089 test-error:0.220947+0.027666
## [21] train-error:0.029043+0.006480 test-error:0.205397+0.020386
```

```
## [31] train-error:0.006224+0.001720    test-error:0.217832+0.030496
## [41] train-error:0.002334+0.001271    test-error:0.218890+0.024266
## Stopping. Best iteration:
## [21] train-error:0.029043+0.006480    test-error:0.205397+0.020386
```

```
# lowest in 6th iteration
```

```
#first default - model training
```

```
xgb1_1 <- xgb.train (params = params, data = dtrain1, nrounds = 79, watchlist =
                    list(val=dtest1,train=dtrain1), print.every.n = 10,
                    early.stop.round = 10, maximize = F ,
                    eval_metric = "error")
```

```
## [17:19:23] WARNING: amalgamation/./src/learner.cc:480:
## Parameters: { early_stop_round, print_every_n } might not be used.
##
## This may not be accurate due to some parameters are only used in language bindings but
## passed down to XGBoost core. Or some parameters are not used but slip through this
## verification. Please open an issue if you find above cases.
##
##
## [1] val-error:0.269729 train-error:0.153527
## Multiple eval metrics are present. Will use train_error for early stopping.
## Will train until train_error hasn't improved in 10 rounds.
##
## [11] val-error:0.263251 train-error:0.097510
## [21] val-error:0.259717 train-error:0.040456
## [31] val-error:0.256773 train-error:0.022822
## [41] val-error:0.249706 train-error:0.007261
## [51] val-error:0.254417 train-error:0.003112
## [61] val-error:0.252061 train-error:0.002075
## Stopping. Best iteration:
## [56] val-error:0.247939 train-error:0.002075
```

```
#model prediction
```

```
xgbpred11 <- predict (xgb1_1,dtest1)
xgbpred11 <- ifelse (xgbpred11 > 0.5,'YES','NO')
xgbpred12 <- as.factor(xgbpred11)
confusionMatrix(xgbpred12 , df_test$Diabetes, positive = "YES")
```

```
## Confusion Matrix and Statistics
```

```
##
```

```
##           Reference
## Prediction  NO  YES
##           NO 1175  31
##           YES 390 102
```

```
##
```

```
##           Accuracy : 0.7521
##           95% CI : (0.7308, 0.7724)
## No Information Rate : 0.9217
## P-Value [Acc > NIR] : 1
```

```
##
```

```
##           Kappa : 0.2316
```

```
##
## McNemar's Test P-Value : <2e-16
##
##      Sensitivity : 0.76692
##      Specificity : 0.75080
##      Pos Pred Value : 0.20732
##      Neg Pred Value : 0.97430
##      Prevalence : 0.07833
##      Detection Rate : 0.06007
##      Detection Prevalence : 0.28975
##      Balanced Accuracy : 0.75886
##
##      'Positive' Class : YES
##
```

```
#Sensitivity : 0.86885
#Specificity : 0.72145
#Accuracy : 0.732
```

```
#####
##SVM
#####

#####
#normal sample
#####
svm_model_normal <- svm(Diabetes ~ ., data=train)
summary(svm_model_normal)
```

```
##
## Call:
## svm(formula = Diabetes ~ ., data = train)
##
##
## Parameters:
##   SVM-Type:  C-classification
##   SVM-Kernel: radial
##       cost:  1
##
## Number of Support Vectors:  1244
##
## ( 471 773 )
##
##
## Number of Classes:  2
##
## Levels:
##  NO YES
```

```
confusionMatrix(predict (svm_model_normal, test) , df_test$Diabetes,
                  positive = "YES")
```

```
## Confusion Matrix and Statistics
```

```
##
##           Reference
## Prediction  NO  YES
##           NO 1565 133
##           YES  0   0
##
##           Accuracy : 0.9217
##           95% CI : (0.9079, 0.934)
##           No Information Rate : 0.9217
##           P-Value [Acc > NIR] : 0.5231
##
##           Kappa : 0
##
## Mcnemar's Test P-Value : <2e-16
##
##           Sensitivity : 0.00000
##           Specificity : 1.00000
##           Pos Pred Value :      NaN
##           Neg Pred Value : 0.92167
##           Prevalence : 0.07833
##           Detection Rate : 0.00000
##           Detection Prevalence : 0.00000
##           Balanced Accuracy : 0.50000
##
##           'Positive' Class : YES
##
```

```
# Sensitivity : 0.00000
# Specificity : 1.00000
# Accuracy : 0.9282

#####
#oversampled
#####

svm_model_over <- svm(Diabetes ~ ., data=train.rose)
summary(svm_model_over)
```

```
##
## Call:
## svm(formula = Diabetes ~ ., data = train.rose)
##
##
## Parameters:
##   SVM-Type:  C-classification
##   SVM-Kernel: radial
##           cost: 1
##
## Number of Support Vectors: 5483
##
## ( 2654 2829 )
##
##
## Number of Classes: 2
```

```
##
## Levels:
## NO YES
```

```
confusionMatrix(predict (svm_model_over, test) , df_test$Diabetes,
                  positive = "YES")
```

```
## Confusion Matrix and Statistics
##
##           Reference
## Prediction  NO  YES
##           NO 1156  30
##           YES  409 103
##
##           Accuracy : 0.7415
##           95% CI : (0.7199, 0.7621)
##           No Information Rate : 0.9217
##           P-Value [Acc > NIR] : 1
##
##           Kappa : 0.2227
##
## Mcnemar's Test P-Value : <2e-16
##
##           Sensitivity : 0.77444
##           Specificity : 0.73866
##           Pos Pred Value : 0.20117
##           Neg Pred Value : 0.97470
##           Prevalence : 0.07833
##           Detection Rate : 0.06066
##           Detection Prevalence : 0.30153
##           Balanced Accuracy : 0.75655
##
##           'Positive' Class : YES
##
```

```
#Sensitivity : 0.88525
#Specificity : 0.71510
#Accuracy : 0.7273
```

```
#####
#undersampled
#####
```

```
svm_model_under <- svm(Diabetes ~ ., data=train.rose_under)
summary(svm_model_under)
```

```
##
## Call:
## svm(formula = Diabetes ~ ., data = train.rose_under)
##
##
## Parameters:
```

```
## SVM-Type: C-classification
## SVM-Kernel: radial
## cost: 1
##
## Number of Support Vectors: 513
##
## ( 245 268 )
##
##
## Number of Classes: 2
##
## Levels:
## NO YES
```

```
confusionMatrix(predict (svm_model_under, test) , df_test$Diabetes,
                  positive = "YES")
```

```
## Confusion Matrix and Statistics
##
##           Reference
## Prediction  NO  YES
##           NO 1153  25
##           YES 412 108
##
##           Accuracy : 0.7426
##           95% CI : (0.7211, 0.7633)
##           No Information Rate : 0.9217
##           P-Value [Acc > NIR] : 1
##
##           Kappa : 0.2354
##
## Mcnemar's Test P-Value : <2e-16
##
##           Sensitivity : 0.81203
##           Specificity : 0.73674
##           Pos Pred Value : 0.20769
##           Neg Pred Value : 0.97878
##           Prevalence : 0.07833
##           Detection Rate : 0.06360
##           Detection Prevalence : 0.30624
##           Balanced Accuracy : 0.77439
##
##           'Positive' Class : YES
##
```

```
# Sensitivity : 0.92623
# Specificity : 0.67893
# Accuracy : 0.6967
```

```
#####
#Comparing Results
#####
```