# Practical session - Modèles de régression linéaire

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## IV. Application: GAFAM or BATX dataset

The data set below shows the number of monthly active users (MAU) on Facebook from 2008 to 2021 in millions (source: https://www.statista.com/statistics/264810/number-of-monthly-active-facebook-users-worldwide/). The numbers were taken from Q4 of each year except for the year 2008, whose data is only available in Q3.

```
tab <- read.table("fb_mau.txt", header=TRUE, sep=",")
tab</pre>
```

```
##
      year
            mau
## 1
     2008
            100
      2009
           360
## 3
      2010 608
## 4
      2011 845
## 5
     2012 1056
      2013 1228
## 7
      2014 1393
      2015 1591
## 8
## 9 2016 1860
## 10 2017 2129
## 11 2018 2320
## 12 2019 2498
## 13 2020 2797
## 14 2021 2912
```

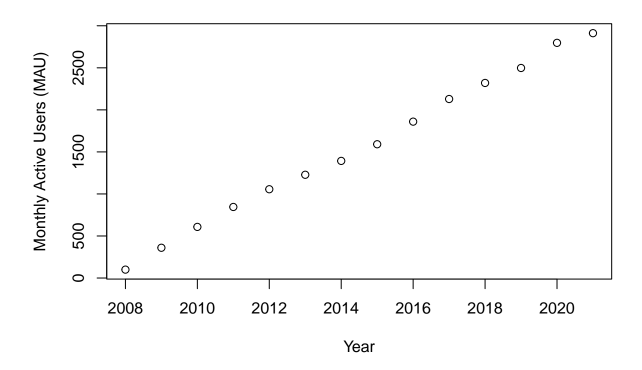
The dimension of the data set:

```
dim(tab)
```

```
## [1] 14 2
```

We then try visualizing the data in order to see if there is an apparent linear relationship between the year and the number of users.

```
plot(tab, xlab="Year", ylab="Monthly Active Users (MAU)")
```



Based on the plotted graph above, we can see that the relationship is fairly linear. Therefore, we can use a linear model to represent the relationship.

```
modreg = lm(mau ~ year, data=tab)
summary(modreg)
```

```
##
## Call:
## lm(formula = mau ~ year, data = tab)
##
##
  Residuals:
##
       Min
                1Q
                    Median
                                3Q
                                       Max
##
   -66.651 -35.664
                    -0.732
                            37.167
                                    60.701
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
## (Intercept) -4.330e+05
                           5.764e+03
                                      -75.13
                                                <2e-16 ***
                2.157e+02 2.861e+00
                                       75.40
                                               <2e-16 ***
## year
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Signif. codes:
##
## Residual standard error: 43.15 on 12 degrees of freedom
## Multiple R-squared: 0.9979, Adjusted R-squared: 0.9977
## F-statistic: 5685 on 1 and 12 DF, p-value: < 2.2e-16
```

According to the summary of the model, the estimated intercept equals  $-4.330 \times 10^5$  and the estimated

coefficient of the year variable equals  $2.157 \times 10^2$ . The model can be written in the form:

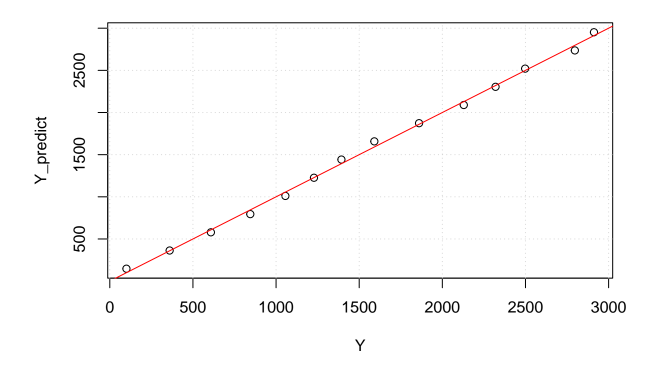
$$\hat{y} = (-4.330 \times 10^5) + (2.157 \times 10^2)x + \hat{\epsilon}$$

where x is the year variable,  $\hat{y}$  is the prediction of the MAU and  $\hat{\epsilon}$  is the residual.

As for the  $R^2$ , we can see that  $R^2 = 0.9979 \approx 1$ . It is a great result since the value corresponds to the cosinus of the angle between the vector of predictions and the vector of the target values, and the closer to 0 the angle gets, the better the model becomes.

```
Y <- tab$mau
Y_predict <- predict(modreg, tab)

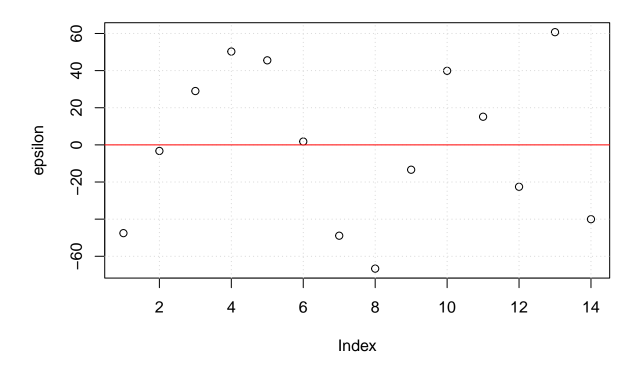
plot(Y, Y_predict)
grid()
abline(a=0, b=1, col="red")</pre>
```



In the graph  $(y, \hat{y})$  above, we can see that the plotted points are fairly close to the bisector, which indicates that the model is acceptable.

The graph below is a scatter plot of the residual for every pair of (y, hat y).

```
epsilon <- Y - Y_predict
plot(epsilon)
grid()
abline(a=0, b=0, col="red")</pre>
```



According to the residual graph or the summary of the linear model above, the absolute value of both the minimum and maximum residual approximately equal to 60, which are for the values y = 1591 and y = 2797.

### V. Medical data

```
tab <- read.table("diabetes.txt", header=TRUE, sep="\t")
```

The dimension of the data set:

```
dim(tab)
```

```
## [1] 442 11
```

The data set consists of p = 10 variables and n = 442 observations.

We now try creating a linear model with the data set.

```
modreg = lm(Y~., data=tab)
summary(modreg)
```

```
##
## Call:
```

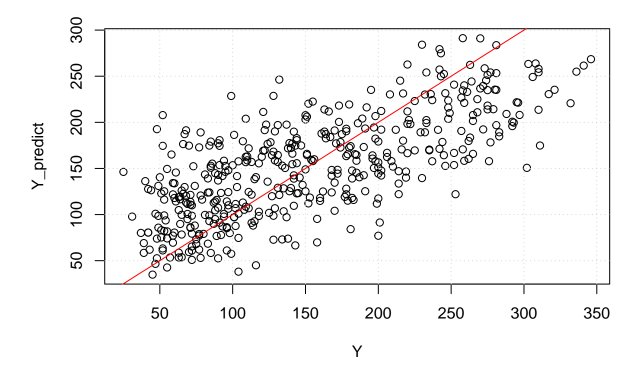
```
## lm(formula = Y ~ ., data = tab)
##
## Residuals:
##
       \mathtt{Min}
                                    3Q
                  1Q
                       Median
                                            Max
## -155.827 -38.536
                       -0.228
                                37.806
                                        151.353
##
## Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) -334.56714
                            67.45462 -4.960 1.02e-06 ***
## AGE
                 -0.03636
                             0.21704 -0.168 0.867031
## SEX
                -22.85965
                             5.83582 -3.917 0.000104 ***
## BMI
                  5.60296
                             0.71711
                                       7.813 4.30e-14 ***
## BP
                  1.11681
                             0.22524
                                      4.958 1.02e-06 ***
## S1
                             0.57333 -1.901 0.057948 .
                 -1.09000
## S2
                  0.74645
                             0.53083
                                       1.406 0.160390
## S3
                  0.37200
                             0.78246
                                       0.475 0.634723
## S4
                             5.95864
                                       1.097 0.273459
                  6.53383
## S5
                 68.48312
                            15.66972
                                       4.370 1.56e-05 ***
## S6
                  0.28012
                             0.27331
                                       1.025 0.305990
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 54.15 on 431 degrees of freedom
## Multiple R-squared: 0.5177, Adjusted R-squared: 0.5066
## F-statistic: 46.27 on 10 and 431 DF, p-value: < 2.2e-16
```

With the summary() function, we can see that the variable "SEX", "BMI", "BPI" and "S5" are the most significant variables of the model.

#### R-squared???

Using the linear model obtained above, we predict the values of y and compare them with the target values in the data set.

```
Y <- tab$Y
Y_predict <- predict(modreg, tab)
plot(Y, Y_predict)
grid()
abline(a=0, b=1, col="red")</pre>
```



Based on the  $(y, \hat{y})$  graph above, we can see that the plotted points are scattered around the bisector, with some points pretty far away from the bisector, rather that appearing on the line. This shows that the difference between predicted values and the target values are quite significant and thus the linear model is not good enough.

The scatter plot below shows the residual for every pair of (y, hat y).

```
epsilon <- Y - Y_predict
plot(epsilon)
grid()
abline(a=0, b=0, col="red")</pre>
```

