Linear regression

```
library(dplyr)
library(ggplot2)
library(corrplot)
```

The prediction task: to predict the amount of calories in Starbucks' drinks based on the other available nutritional data.

Load and inspect the data

The data originates from the Starbucks menu data set. It has been adapted for the class.

Load the data

```
starbucks <- read.csv("data/starbucks_calories.csv", encoding="UTF-8")
```

Inspect the data

```
str(starbucks)
```

```
## 'data.frame':
                   241 obs. of 7 variables:
                      : int 1 2 3 4 5 6 7 8 9 10 ...
   $ drink_id
                            "Brewed Coffee (Short)" "Brewed Coffee (Tall)" "Brewed Coffee (Grande)" "
## $ beverage
## $ calories
                      : int 3 4 5 5 70 100 70 100 150 110 ...
## $ fat_tot
                      : num 0.1 0.1 0.1 0.1 0.1 3.5 2.5 0.2 6 4.5 ...
  $ carbohydrates_tot: int
                            5 10 10 10 75 85 65 120 135 105 ...
##
##
   $ sugar
                      : int
                            0 0 0 0 9 9 4 14 14 6 ...
   $ proteins
                      : num 0.3 0.5 1 1 6 6 5 10 10 8 ...
```

```
# View(starbucks)
```

As the first step, we will do the prediction based on just one variable (simple regression) and then will expand to all potentially relevant variables (multiple regression).

Examining relationships among variables

Let's start examining the correlations among the variables. In particular, we are primarily interested in correlations between the outcome variable (calories) and other variables that may serve as predictors.

Select variables that might be used for predicting calories

```
sb_calories = starbucks[, 3:7]
```

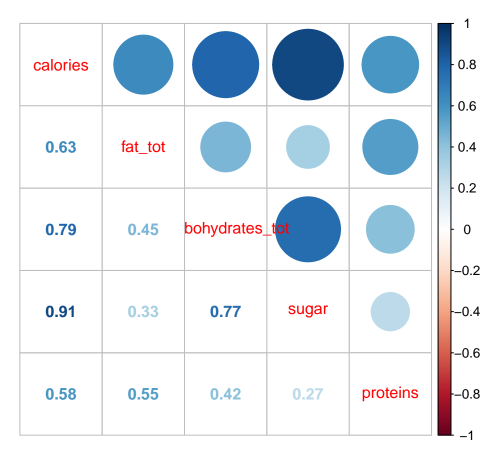
Compute correlations

```
sb_cor = cor(sb_calories)
sb_cor
```

```
##
                     calories
                                 fat_tot carbohydrates_tot
                                                               sugar proteins
## calories
                     1.0000000 0.6342758
                                                 0.7935856 0.9091753 0.5825552
                                                 0.4509799 0.3315062 0.5508502
## fat_tot
                     0.6342758 1.0000000
## carbohydrates_tot 0.7935856 0.4509799
                                                 1.0000000 0.7691970 0.4151386
## sugar
                     0.9091753 0.3315062
                                                 0.7691970 1.0000000 0.2671951
## proteins
                     0.5825552 0.5508502
                                                 0.4151386 0.2671951 1.0000000
```

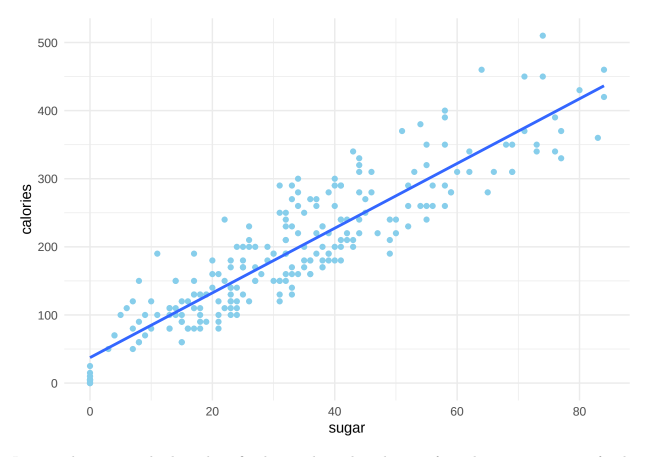
We can also plot this matrix to make it easier for inspection

```
corrplot.mixed(sb_cor)
```



Clearly all selected variables are highly correlated with calories, but especially sugar. Let's inspect that closer.

```
## 'geom_smooth()' using formula = 'y ~ x'
```



It seems that sugar and calories have fine linear relationship, thus satisfying the primary criterion for the application of linear modeling.

Train-test split

Before going any further, we first need to split the data into training and test sets. We will build our models using training data only, while test data will be used for model evaluation.

There are 2 key requirements when splitting the data into training and test sets: 1) random selection of observations to go into training and test sets 2) ensuring the same distribution of the outcome variable in both parts (train and test) We can achieve both things by using the createDataPartition function from the caret package.

library(caret)

```
set.seed(2024)
training_indices = createDataPartition(sb_calories$calories, p = 0.75, list = FALSE)
sb_train = sb_calories[training_indices, ]
sb_test = sb_calories[-training_indices, ]
```

Check the distribution of the calories variable on the train and test sets

```
summary(sb_train$calories)
```

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.0 120.0 180.0 192.6 260.0 510.0
```

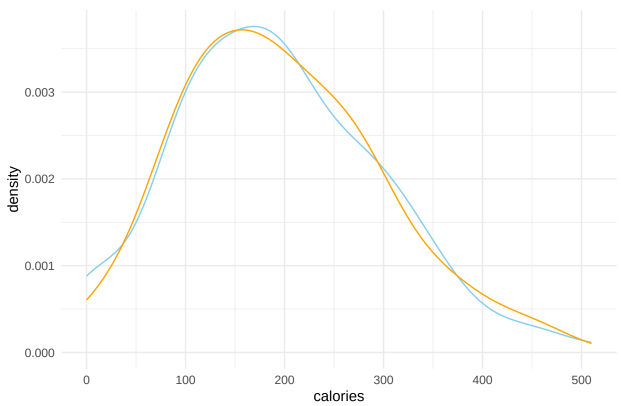
summary(sb_test\$calories)

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.0 120.0 185.0 195.7 260.0 460.0
```

We can do that visually as well

```
ggplot() +
  geom_density(data = sb_train, mapping = aes(x=calories), color = "skyblue") +
  geom_density(data = sb_test, mapping = aes(x=calories), color = "orange") +
  labs(title = "Distribution of the calories variable on the train and test sets") +
  theme_minimal()
```

Distribution of the calories variable on the train and test sets



Simple linear regression

Create the model

```
lm1 = lm(calories ~ sugar, data = sb_train)
```

Print the model

summary(lm1)

```
##
## Call:
## lm(formula = calories ~ sugar, data = sb_train)
##
## Residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -78.821 -34.102 -9.812 24.042 115.751
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 34.8118
                           6.1384
                                  5.671 5.52e-08 ***
                           0.1622 29.951 < 2e-16 ***
## sugar
                4.8573
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 42.61 on 181 degrees of freedom
## Multiple R-squared: 0.8321, Adjusted R-squared: 0.8312
## F-statistic: 897.1 on 1 and 181 DF, p-value: < 2.2e-16
```

Get the estimated paramter values

```
coef(lm1)
```

```
## (Intercept) sugar
## 34.811751 4.857267
```

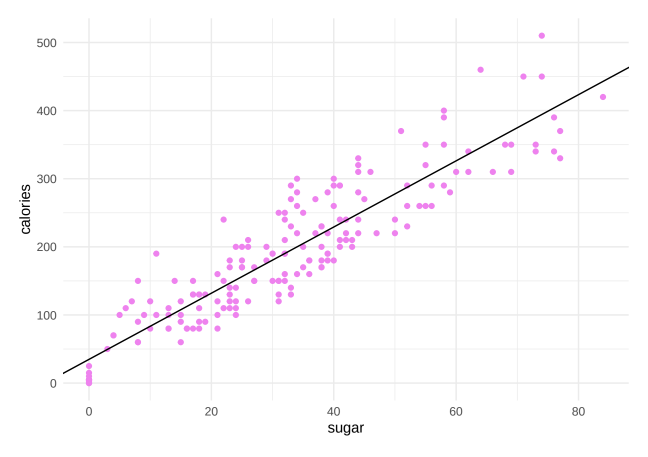
and the confidence interval for the parameters...

```
confint(lm1, level = 0.95)
```

```
## 2.5 % 97.5 %
## (Intercept) 22.69981 46.923688
## sugar 4.53727 5.177263
```

We would interpret this as follows: in 95% of samples taken from the population of Starbucks drinks, the estimated value of the parameter associated with the sugar variable will be in the (4.53, 5.18) range.

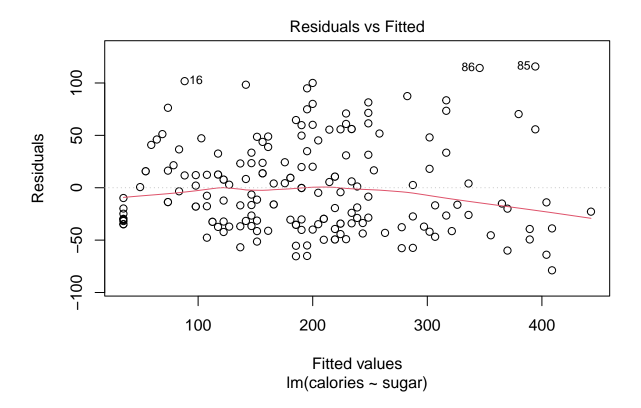
Plot the regression line

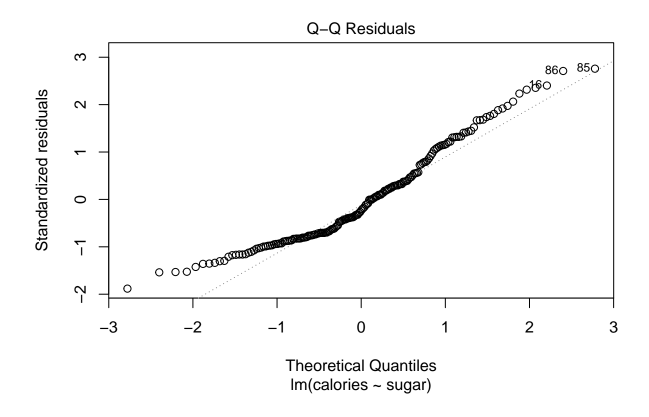


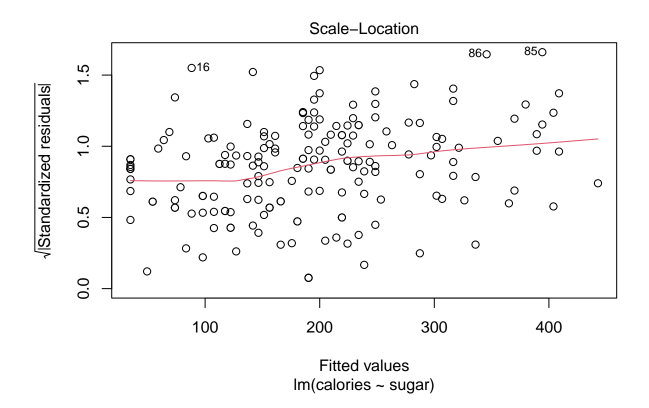
Check if the assumptions for linear models are met, using the plot function. The function will create 4 plots:

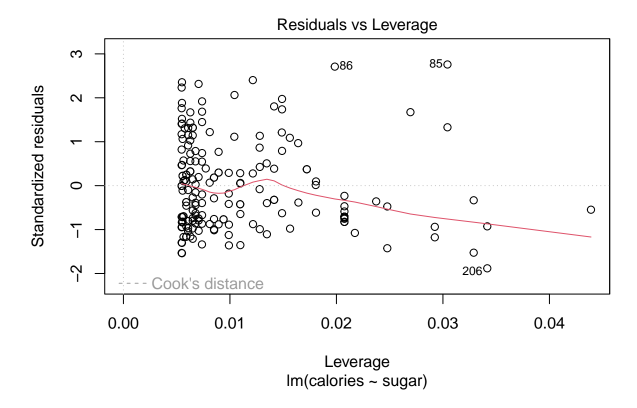
- The 1st plot, Residual vs Fitted value, is used for checking if the linearity assumption is satisfied.
- The 2nd plot, Normal Q-Q plot, tells us if residuals are normally distributed.
- The 3rd plot, **Scale-Location**, is used for checking the assumption known as homoscedasticity that the residuals have equal variance.
- The 4th plot, **Residuals vs Leverage**, is used for spotting the presence of highly influential observations; those are the observations that, if included in or excluded from the model, can significantly affect the values of the model parameters. Such observations are identified among those with unusually high/low output values (outliers) and/or unusually high/low predictor values (high leverage).

plot(lm1)



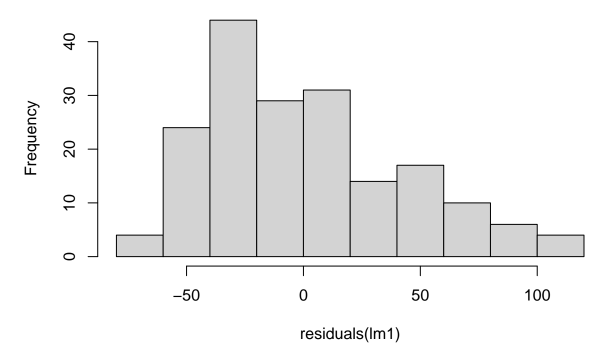






hist(residuals(lm1))

Histogram of residuals(Im1)



So, out of the four assumptions, one is not satisfied - normality of residuals. This may affect the prediction intervals (see below), that is, lead to incorrect confidence and prediction intervals, since those intervals are calculated based on the assumption that the residuals are normally distributed.

Make predictions and evaluate the model on the test data

```
lm1_predicted = predict(lm1, newdata = sb_test)
sb_test |> select(sugar, calories) |> mutate(calories_pred = lm1_predicted) |> head()
##
      sugar calories calories_pred
## 5
          9
                   70
                           78.52715
## 8
         14
                  100
                          102.81348
## 9
         14
                  150
                          102.81348
## 12
         17
                  190
                          117.38528
## 17
         17
                  110
                          117.38528
## 22
         20
                  180
                          131.95708
```

We can also get confidence and prediction intervals for our predictions

```
lm1_pred_ci = predict(lm1, newdata = sb_test, interval = "confidence", level = 0.95)
sb_test |> select(sugar, calories) |> cbind(lm1_pred_ci) |> head()

## sugar calories fit lwr upr
## 5 9 70 78.52715 68.77442 88.27988
```

```
## 8 14 100 102.81348 94.23291 111.39405

## 9 14 150 102.81348 94.23291 111.39405

## 12 17 190 117.38528 109.43607 125.33450

## 17 17 110 117.38528 109.43607 125.33450

## 22 20 180 131.95708 124.56810 139.34606
```

Confidence intervals reflects the uncertainty around the mean predictions. For example, for a drink with 9g of sugar the expected (mean) value of calories will range between 68.77 and 88.28

```
lm1_pred_pi = predict(lm1, newdata = sb_test, interval = "prediction", level = 0.95)
sb_test |> select(sugar, calories) |> cbind(lm1_pred_pi) |> head()
```

```
##
      sugar calories
                            fit
                                      lwr
                                                upr
## 5
          9
                  70 78.52715 -6.118175 163.1725
## 8
         14
                 100 102.81348 18.295192 187.3318
## 9
         14
                 150 102.81348 18.295192 187.3318
## 12
         17
                 190 117.38528 32.928753 201.8418
## 17
         17
                 110 117.38528 32.928753 201.8418
## 22
         20
                 180 131.95708 47.551440 216.3627
```

Prediction interval reflects uncertainty related to a particular instance. For example, for a particular drink with 9g of sugar the value of calories will range between -6.12 and 123.18. However, note that due to the lack of normality of residuals (the assumption that was violated), the prediction intervals for this particular model can be considered trustworthy

To evaluate the model, we will compute - on the test set - two evaluation measures: R2 and RMSE

```
R2 = (TSS - RSS)/TSS RMSE = sqrt(RSS/n)
```

```
compute_eval_measures <- function(y_train, y_test, y_pred) {
  tss = sum((y_test - mean(y_train))^2)
  rss = sum((y_test - y_pred)^2)
  r2 = (tss - rss)/tss

n = length(y_test)
  rmse = sqrt(rss/n)

return(c(R2=r2, RMSE=rmse))
}</pre>
```

```
round(lm1_eval, 4)
```

```
## R2 RMSE
## 0.8057 43.9478
```

To get a perspective of how large the error is, we'll compare it with the mean value of the response variable on the test set.

```
lm1_rmse = as.numeric(lm1_eval['RMSE'])
lm1_rmse/mean(sb_test$calories)
```

```
## [1] 0.2245791
```

It's a fairly large error - about 22% of the average value.

Multiple linear regression

We've seen that in addition to sugar, the other three variables are also highly correlated with calories (as the outcome variable). Let's see if by adding them to the model we can improve the prediction performance.

```
lm2 = lm(calories ~ fat_tot + sugar + proteins + carbohydrates_tot, data = sb_train)
summary(lm2)
```

```
##
## Call:
## lm(formula = calories ~ fat_tot + sugar + proteins + carbohydrates_tot,
##
       data = sb_train)
##
## Residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
## -24.610 -4.832 -1.140
                             3.079 36.794
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
                                 1.40170
                                          1.540
## (Intercept)
                     2.15915
                                                    0.125
## fat tot
                                 0.28259 31.429
                     8.88152
                                                   <2e-16 ***
## sugar
                     3.88674
                                 0.06095 63.765
                                                   <2e-16 ***
## proteins
                     5.06721
                                 0.17064
                                         29.696
                                                   <2e-16 ***
## carbohydrates_tot 0.01387
                                 0.01515
                                           0.916
                                                    0.361
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## Residual standard error: 8.729 on 178 degrees of freedom
## Multiple R-squared: 0.9931, Adjusted R-squared: 0.9929
## F-statistic: 6378 on 4 and 178 DF, p-value: < 2.2e-16
```

Note that in the presence of sugar, carbohydrates_tot are not relevant for the prediction of calories. This is intuitively clear, but might also be explained by the high correlation of sugar and carbohydrates_tot variables (see the correlations plot above). This points to a potential problem of multicollinearity that is often present in case of multiple linear regression and should be avoided as it tends to make the resulting model unstable and/or plainly wrong.

Let's check if we have the case of multicollinearity in our data

```
library(performance)
```

One way to check for multicollinearity is to compute the Variance Inflation Factor (VIF):

```
lm2_collinearity = check_collinearity(1m2)
lm2_collinearity
```

```
## # Check for Multicollinearity
##
## Low Correlation
##
##
                              VIF 95% CI Increased SE Tolerance Tolerance 95% CI
                  Term VIF
                                                             0.60
                                                                       [0.48, 0.71]
##
              fat_tot 1.66 [1.41, 2.08]
                                                   1.29
##
                 sugar 3.37 [2.69, 4.31]
                                                   1.83
                                                             0.30
                                                                       [0.23, 0.37]
##
             proteins 1.61 [1.37, 2.01]
                                                   1.27
                                                             0.62
                                                                       [0.50, 0.73]
    carbohydrates_tot 3.76 [2.99, 4.83]
                                                             0.27
                                                                       [0.21, 0.33]
##
                                                   1.94
```

Variance Inflation Factor (VIF) is an often used way to check for multicollinearity. If the VIF value of a variable is high, it means the information in that variable is already explained by other predictors present in the given model, which means that the variable is redundant. If interested in how VIF is computed, this article offers a nice and simple explanation.

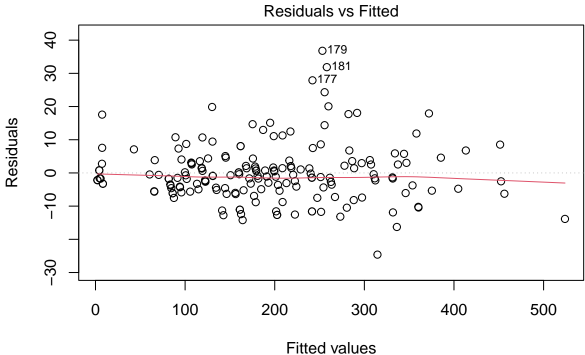
As a rule of thumb, variables having $\sqrt{VIF} > 2$ are problematic. Here, it is not the case. When it is the case, we remove the variable with the high VIF value from the data set and build a new model without it. If there are several such variables, we remove them in a step-wise manner, starting from the one with the highest VIF value.

Even though the presence of the carbohydrates_tot variable is not causing the problem of multicollinearity and from that perspective can be kept in the model, it proved to be irrelevant for the prediction of the outcome variable and thus it should be better removed from the model. As a general rule, if two (or more) models are of similar performance, it is better to keep the simpler model, since such a model is less prone to overfitting.

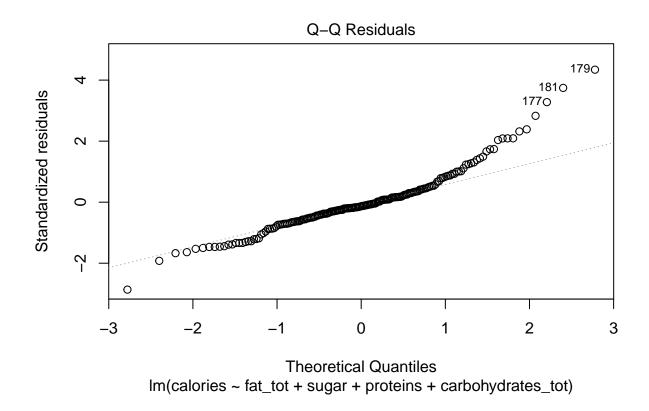
```
lm3 = lm(calories ~ fat_tot + sugar + proteins, data = sb_train)
summary(lm3)
```

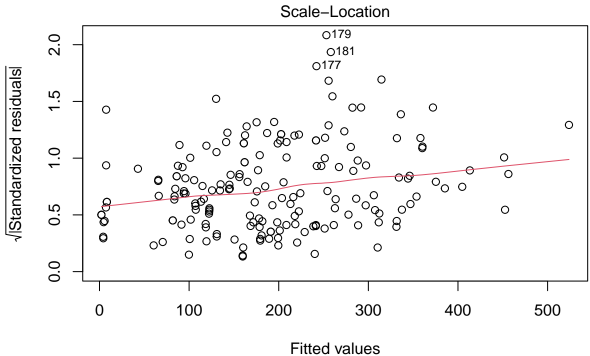
```
##
## Call:
## lm(formula = calories ~ fat_tot + sugar + proteins, data = sb_train)
##
## Residuals:
##
       Min
                1Q
                    Median
                                 3Q
                                        Max
   -25.030
            -4.795
                    -1.190
                              3.233
                                     36.190
##
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                2.21598
                           1.39969
                                      1.583
                                               0.115
## fat tot
                8.92549
                                    32.065
                                              <2e-16 ***
                           0.27835
## sugar
                3.93163
                           0.03621 108.568
                                              <2e-16 ***
                5.09198
                           0.16840
                                     30.237
                                              <2e-16 ***
## proteins
## Signif. codes:
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 8.725 on 179 degrees of freedom
## Multiple R-squared: 0.993, Adjusted R-squared: 0.9929
## F-statistic: 8511 on 3 and 179 DF, p-value: < 2.2e-16
```

Check the four assumtions of linear models:

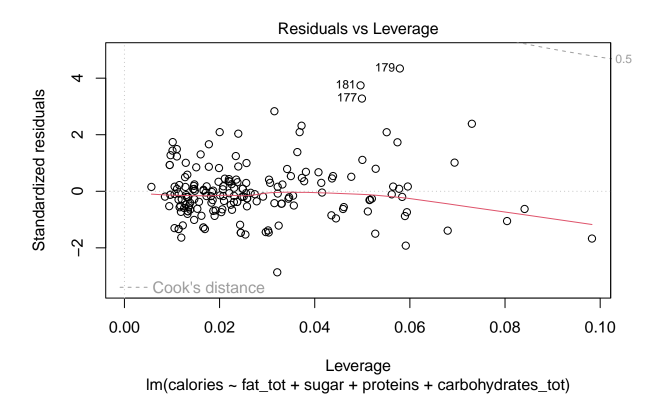


Im(calories ~ fat_tot + sugar + proteins + carbohydrates_tot)



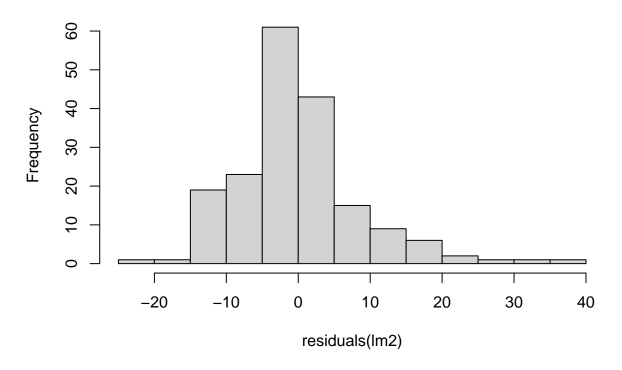


Im(calories ~ fat_tot + sugar + proteins + carbohydrates_tot)



hist(residuals(lm2))

Histogram of residuals(Im2)



This time, all assumptions hold

Let's check **confidence** and **prediction** intervals for predictions based on this new model; this time, the intervals should be trustworthy since the all the assumptions are satisfied

```
lm3_pred_ci = predict(lm3, newdata = sb_test, interval = "confidence", level = 0.95)
sb_test |> select(sugar, calories) |> cbind(lm3_pred_ci) |> head()
```

```
##
      sugar calories
                            fit
                                       lwr
## 5
                                 66.77282
          9
                  70
                      69.04508
                                           71.31735
## 8
         14
                  100 109.96371 107.15340 112.77401
## 9
         14
                 150 161.73157 159.14509 164.31806
## 12
         17
                 190 192.63592 189.77384 195.49800
                  110 118.08579 116.31730 119.85428
## 17
         17
         20
                 180 171.30387 169.33108 173.27666
## 22
```

```
lm3_pred_pi = predict(lm3, newdata = sb_test, interval = "predict", level = 0.95)
sb_test |> select(sugar, calories) |> cbind(lm3_pred_pi) |> head()
```

```
##
      sugar calories
                            fit
                                       lwr
                                                 upr
## 5
          9
                       69.04508
                                 51.67778
                                            86.41239
## 8
         14
                  100 109.96371
                                 92.51785 127.40956
## 9
         14
                  150 161.73157 144.32037 179.14278
## 12
         17
                  190 192.63592 175.18165 210.09019
## 17
         17
                  110 118.08579 100.77719 135.39439
                  180 171.30387 153.97320 188.63454
## 22
         20
```

We will now make predictions on the test set and compute evaluation measures

```
lm3_predicted = predict(lm3, newdata = sb_test)
```

```
## R2 RMSE
## 0.9856 11.9769
```

```
lm3_rmse = as.numeric(lm3_eval['RMSE'])
lm3_rmse/mean(sb_test$calories)
```

```
## [1] 0.06120335
```

Compare the performance of the latest model with the initial model

```
rbind(lm1_eval, lm3_eval) |> round(4)

## R2 RMSE
## lm1_eval 0.8057 43.9478
## lm3_eval 0.9856 11.9769
```

Obviously the latest model is much better than the initial one.

Note: linear regression is not restricted to numerical variables; that is, a linear model can be built with categorical variables, as well, or a mix of numerical and categorical variables. However, model interpretation in that case is more difficult than when only numerical variables are used and thus is out of the scope of this course.