Medical Statistics Course

2020-10-05

Exploratory data analysis

Data representation

Motivation

A searcher receives the dataset decathlon2 from the factoextra package. This dataset describe the performances of several athletes in two different sport events. Before starting any analysis, the searcher wants to have a feeling about the data and wants to create plot in order to chose what type of analysis would be suitable for this dataset.

```
library(factoextra)
## Loading required package: ggplot2
## Welcome! Want to learn more? See two factoextra-related books at https://goo.gl/ve3WBa
data(decathlon2)
# Display a description of the dataset
str(decathlon2)
## 'data.frame':
                    27 obs. of 13 variables:
## $ X100m
                  : num 11 10.8 11 11.3 11.1 ...
##
   $ Long.jump
                  : num 7.58 7.4 7.23 7.09 7.3 7.31 6.81 7.56 6.97 7.27 ...
   $ Shot.put
                        14.8 14.3 14.2 15.2 13.5 ...
                  : num
   $ High.jump
                         2.07 1.86 1.92 2.1 2.01 2.13 1.95 1.86 1.95 1.98 ...
##
                  : num
   $ X400m
                         49.8 49.4 48.9 50.4 48.6 ...
                  : num
##
  $ X110m.hurdle: num 14.7 14.1 15 15.3 14.2 ...
  $ Discus
                 : num 43.8 50.7 40.9 46.3 45.7 ...
  $ Pole.vault : num
                         5.02 4.92 5.32 4.72 4.42 4.42 4.92 4.82 4.72 4.62 ...
##
                         63.2 60.1 62.8 63.4 55.4 ...
   $ Javeline
                  : num
##
  $ X1500m
                  : num 292 302 280 276 268 ...
  $ Rank
                  : int 1 2 4 5 7 8 9 10 11 12 ...
                  : int 8217 8122 8067 8036 8004 7995 7802 7733 7708 7651 ...
##
   $ Competition : Factor w/ 2 levels "Decastar", "OlympicG": 1 1 1 1 1 1 1 1 1 1 1 ...
# Display the first lines of the table
head(decathlon2)
             X100m Long.jump Shot.put High.jump X400m X110m.hurdle Discus
##
## SEBRLE
             11.04
                        7.58
                                14.83
                                           2.07 49.81
                                                             14.69
                                                                    43.75
## CLAY
             10.76
                        7.40
                                14.26
                                           1.86 49.37
                                                                    50.72
                                                             14.05
## BERNARD
             11.02
                        7.23
                                14.25
                                           1.92 48.93
                                                             14.99
                                                                    40.87
## YURKOV
             11.34
                        7.09
                                15.19
                                           2.10 50.42
                                                             15.31
                                                                    46.26
## ZSIVOCZKY 11.13
                        7.30
                                13.48
                                           2.01 48.62
                                                             14.17
                                                                    45.67
## McMULLEN 10.83
                        7.31
                                13.76
                                           2.13 49.91
                                                             14.38 44.41
##
             Pole.vault Javeline X1500m Rank Points Competition
```

##	SEBRLE	5.02	63.19	291.7	1	8217	Decastar
##	CLAY	4.92	60.15	301.5	2	8122	Decastar
##	BERNARD	5.32	62.77	280.1	4	8067	Decastar
##	YURKOV	4.72	63.44	276.4	5	8036	Decastar
##	ZSIVOCZKY	4.42	55.37	268.0	7	8004	Decastar
##	McMULLEN	4.42	56.37	285.1	8	7995	Decastar

Working example

We consider the following simulated dataset: 2 continuous variable scoreX, scoreY, and a categorical variable representing the membership of group A or B.

```
set.seed(352)
sample.size<-60

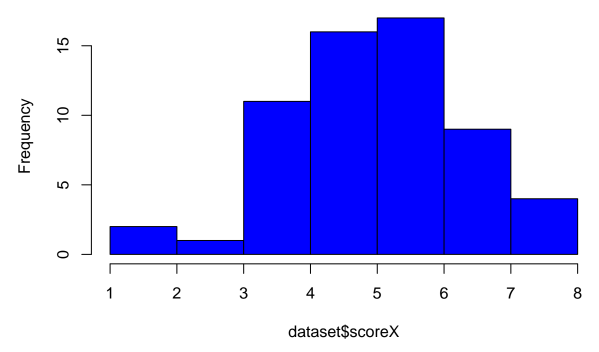
scoreX <- rnorm(sample.size, mean = 5, sd = 1.2)
scoreY <- rgamma(sample.size, shape = 2, scale = 1)
group <- rep(c('A', 'B'), each = sample.size/2)

dataset <- data.frame(scoreX = scoreX, scoreY = scoreY, group = group, stringsAsFactors = T)</pre>
```

Before doing any statistical analysis it is important to look at the distributions of the variables available. Indeed, looking at the shape of the distributions can orient the searcher towards specific analysis methods. Let's look at the distribution of the variable scoreX by plotting an histogram of the variable. This can be done using the function hist in R:

```
#plot the histogram of a variable
hist(dataset$scoreX, col = 'blue', bg = "gray")
```

Histogram of dataset\$scoreX

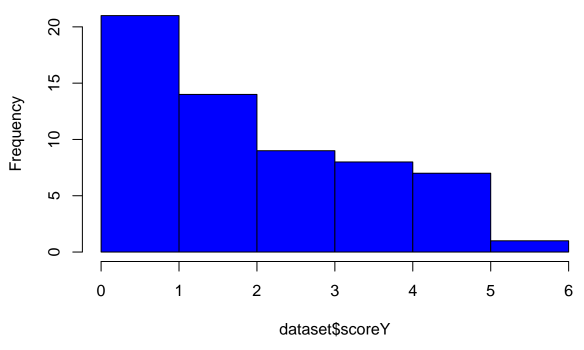


The histogram plot is the count of the number of observations falling into a specific bin. A bin, being an

interval used to divide the space of the variable. By using this plot we can get a good approximation of the shape of the distribution of the values of a variable. In our case we simulated the variables using a normal distribution. It also allows us to detect possible extreme values or skewness of the distributions as it can be seen in the variable scoreY.

```
#plot the histogram of a variable
hist(dataset$scoreY, col='blue')
```

Histogram of dataset\$scoreY



We can also display histograms per group to compare the shape of the distributions between two groups. To do so we need to split the data in two:

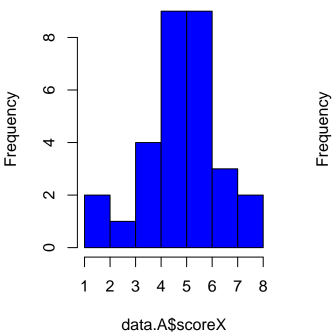
```
#Creation dataset group A
data.A <- dataset[dataset$group == 'A',]

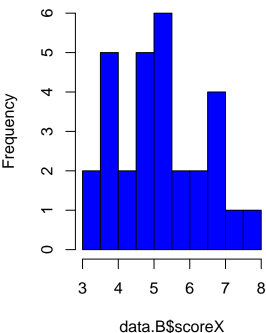
#Creation dataset group B
data.B <- dataset[dataset$group == 'B',]

#plot the histograms of scoreX for both groups
par(mfrow = c(1,2))
hist(data.A$scoreX, col = 'blue', main = 'histogram scoreX group A')
hist(data.B$scoreX, col = 'blue', main = 'histogram scoreX group B')</pre>
```

histogram scoreX group A

histogram scoreX group B





Here, the histograms of the variable <code>scoreX</code> look quite different. However, the values from both groups are drawn using the same distribution. The differences observed are only due to randomness. We can, for example, add to these histograms the density plot of the distribution used to simulate the <code>scoreX</code> variable:

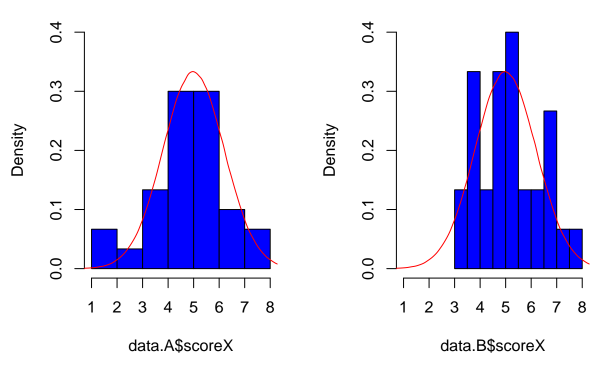
```
#simulation of values from a gaussian distribution with mean 5 and standard deviation 1.2
d <- rnorm(1000000, mean = 5, sd = 1.2)

#plot the histograms of scoreX for both groups with added density
par(mfrow = c(1,2))

hist(data.A$scoreX, col = 'blue', prob = TRUE, main = 'histogram scoreX group A', ylim = c(0, 0.4), xlines(density(d), col = 'red')
hist(data.B$scoreX, col = 'blue', prob = TRUE, main = 'histogram scoreX group B', ylim = c(0, 0.4), xlines(density(d), col = 'red')</pre>
```

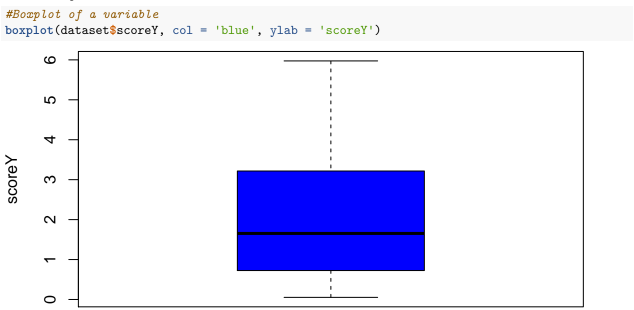
histogram scoreX group A

histogram scoreX group B



As we can see, it is not possible to determine from these plots that the distribution of score x in both groups are different. This is why statistical testing is needed to compare both group and draw conclusions.

Another important and useful way to represent the data are the boxplots. Boxplots are graphical representations of summary measures of a distribution represented as the name indicates in the shape of a box. We will draw a boxplot of the variable scorey:

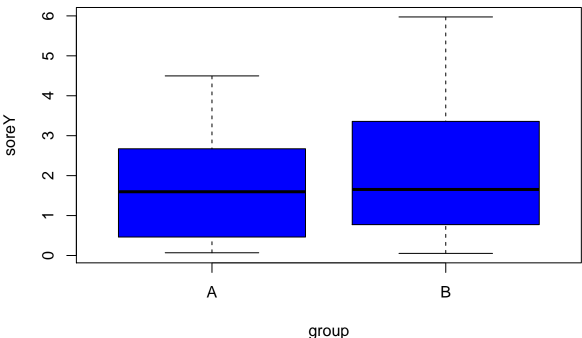


The box represents the space between the first and the third quartile values of the variable. Quartiles are the three values of a variables splitting the variables in 4 equal parts. The thick line represent the median of the variables, i.e. the second quartile which has the particularity to split that variable into two equal parts. Finally, the top and bottom lines represent a space equal to 1.5 the boxsize from nearest edge of the box.

Any values above or below these lines are represented as points and are considered as extreme values and possible outliers.

On the contrary of the histograms where we need to split the data to plot histograms for different groups, representing boxplots of a same variable for different groups is really easy and just require the use of a formula R object as shown below:





As we can see in these box plots, the median value for both groups is really close but we can observe a larger box for group B indicating more variations in this group. However, statistical methods are needed to prove if this difference is real or not.

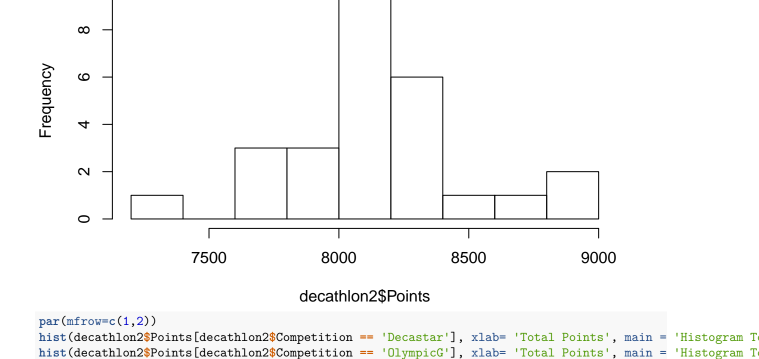
Quick tasks

Let us go back to the decathlon2 dataset. Use both histograms and boxplots to represent the distribution of the variable Points overall and by competitions.

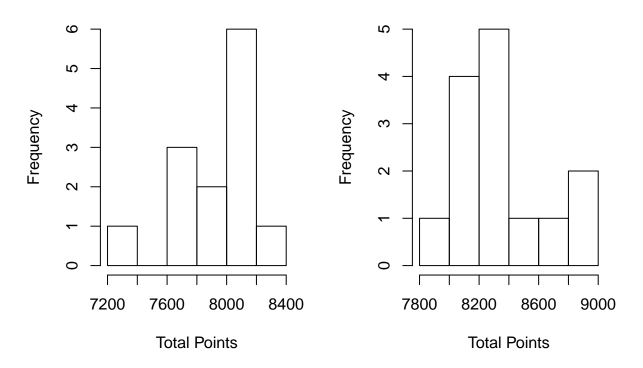
hist(decathlon2\$Points)

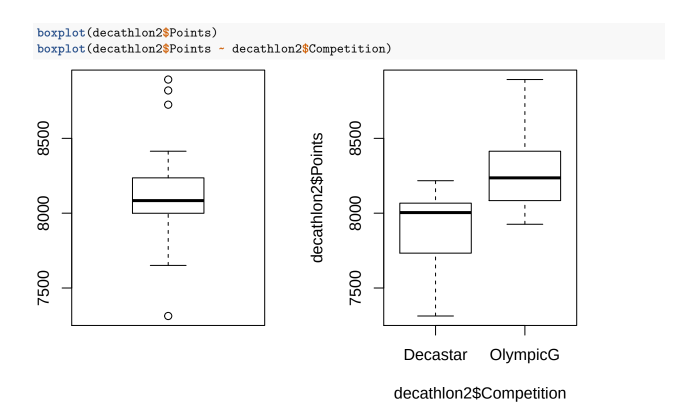
Histogram of decathlon2\$Points

10



Histogram Total points at Decastlistogram Total points at Olympic G





Mean and variance

Motivation

After looking at the distributions of the several variables contained in decathlon2, the searcher wants to obtain several summary measures for these variables.

```
library(factoextra)
data(decathlon2)
```

Working example

We consider the same dataset that we simulated in the previous part:

```
set.seed(352)
sample.size<-60

scoreX <- rnorm(sample.size, mean = 5, sd = 1.2)
scoreY <- rgamma(sample.size, shape = 2, scale = 1)
group <- rep(c('A', 'B'), each = sample.size/2)

dataset <- data.frame(scoreX = scoreX, scoreY = scoreY, group = group, stringsAsFactors = T)</pre>
```

By looking at the histograms and boxplots created we could see possible differences between the variables in the different groups. We now want to derive summary measures in order to corroborate what the eyes could see. The most common summary measures of continuous variables are the mean and the standard deviation.

The sample mean represents the central value of the combined values of all samples for a specific variable. The mean is computed as the sum of all the values of a variable divided by the total number of values measured:

$$\overline{x} = \frac{\sum_{i}^{N} (x_i)}{N}$$

We want to compute mean values of scoreX and scoreY for the groups A and B.

```
#Creation dataset group A containing only scoreX and scoreY
data.A <- dataset[dataset$group == 'A', c('scoreX', 'scoreY')]

#Creation dataset group B containing only scoreX and scoreY
data.B <- dataset[dataset$group == 'B', c('scoreX', 'scoreY')]

#Computation of the mean for each variable of the dataset for group A
mean.variables.groupA <-apply(data.A, 2, mean, na.rm=T)
mean.variables.groupA

## scoreX scoreY
## 4.880274 1.750616

#Computation of the mean for each variable of the dataset for group B
mean.variables.groupB <-apply(data.B, 2, mean, na.rm=T)
mean.variables.groupB
```

scoreX scoreY ## 5.148834 2.197438

We can see slightly different values for both variables in both groups. However, the mean gives us only information about the central values for both variables. It is important to introduce a measure of variation in order to determine how values are distributed 'around' the mean. To do so, we use the standard deviation.

The standard deviation represents the amount of variation of a set of value. It measures the amount of dispersion from the mean of the set of values and can be computed as:

$$\sigma = \sqrt{\left(\frac{\sum_{i}^{N} (x_i - \overline{x})^2}{N}\right)}$$

We will now compute the standard deviation for both variables in each group:

```
#Computation of the standard deviation for both variables of the dataset in group A
sd.variables.groupA <- apply(data.A, 2, sd, na.rm=T)
sd.variables.groupA

## scoreX scoreY
## 1.370947 1.331300

#Computation of the standard deviation for both variables of the dataset in group B
sd.variables.groupB <- apply(data.B, 2, sd, na.rm=T)
sd.variables.groupB</pre>
```

```
## scoreX scoreY
## 1.14792 1.61810
```

A large standard deviation illustrates the fact that one or several values can be very different from the mean while a small standard deviation shows that most values are very close to the mean.

This two measure are very useful to describe a distribution and are very important in statistical testing as we will see later in the course. However, mean and standard deviation are not perfect to describe all distributions.

Indeed, one weakness of the mean is the lack of robustness to extreme values. The mean can be strongly influenced by the presence of a proportion of extremely large or small values. It is, therefore, not the best

summary measure for skewed distributions. In such case, quantiles of a set of values are preferred.

Quantiles are cut points that are dividing a set of values in subsets of same size. Quartiles represented in the boxplot are 3 values that split a set of values into 4 parts of equal sizes. A quantile particularly interesting to define a central point of a set of values is the median.

The median is the value that splits a set of values into two equal parts. By definition this descriptive measure is more robust to extreme values as it's value is not influenced directly by the extreme values. Indeed having a maximum value of 100 or 1000 will not influence the value of the median as opposed to the mean.

We can compute easily the median in R:

5.060254 1.653759

```
#computation of the median for both variables in group A
median.variables.groupA <- apply(data.A, 2, median)
median.variables.groupA

## scoreX scoreY
## 4.926336 1.596064

#computation of the median for both variables in group B
median.variables.groupB <- apply(data.B, 2, median)
median.variables.groupB</pre>
## scoreX scoreY
```

We can see that for both variables, the median for both groups are closer to each other. This is particularly the case for <code>scoreY</code>, which was identified as skewed when plotting the histograms. By randomness, due to the presence of larger values in group B the mean was really different from group A. Both set of values in group A and group B are coming from the same distribution and no differences should be seen. This is the case when we use the median.

It can also be of interest to look at the other quantiles of the set of values. Let's take the variable ScoreY as an example:

```
#computation of quantile of the variable scoreY for group A
quantile(data.A$scoreY)
##
           0%
                     25%
                                 50%
                                            75%
                                                      100%
## 0.06849886 0.48053515 1.59606376 2.63762448 4.49709044
#computation of quantile of the variable scoreY for group B
quantile(data.B$scoreY)
##
                     25%
                                 50%
                                            75%
                                                      100%
## 0.05282527 0.77458628 1.65375910 3.34040589 5.97353896
```

By default the function quantile provides the minimum, the maximum and the quartiles of a set of values. In this example we can see that the distance between quantile in both groups is small until the median. The third quartile and the maximum values explain the difference obtain in the mean for the variable scoreY between group A and group B.

As for the mean, we can also measure the amount of variation of the variable based on the median. This measure is called the median absolute variation (MAD). It is obtained by calculating the median value of the distance to the median, as described by the formula below:

```
## scoreX scoreY
## 1.134796 1.626960

#computation of the median absolute deviation for group A
mad.variables.groupB <- apply(data.B, 2, mad)
mad.variables.groupB

## scoreX scoreY</pre>
```

This measure of dispersion is also robust to extreme values. Indeed, when looking at the formulas to compute the standard deviation and the median absolute deviation, we can see that for the standard deviation the distance to the mean is squared while for the MAD this is not the case.

Quick tasks

1.282703 1.723980

Let us go back to the decathlon2 dataset and compute the mean, and the standard deviation.

```
data.Decastar <- decathlon2[decathlon2$Competition == 'Decastar', c(2:10,12)]
data.OlympicGames <- decathlon2[decathlon2$Competition == 'OlympicG', c(2:10,12)]
mean.variables.Decastar <- apply(data.Decastar, 2, mean, na.rm=T)
mean.variables.OlympicGames <- apply(data.OlympicGames, 2, mean, na.rm=T)
sd.variables.Decastar <- apply(data.Decastar, 2, sd, na.rm=T)
sd.variables.OlympicGames <- apply(data.OlympicGames, 2, sd, na.rm=T)</pre>
```

Now compute the median, and the median absolute deviation for each event and the total number of points for each competition.

```
data.Decastar <- decathlon2[decathlon2$Competition == 'Decastar', c(1:10,12)]
data.OlympicGames <- decathlon2[decathlon2$Competition == 'OlympicG', c(1:10,12)]
mean.variables.Decastar <- apply(data.Decastar, 2, median, na.rm=T)
mean.variables.OlympicGames <- apply(data.OlympicGames, 2, median, na.rm=T)
sd.variables.Decastar <- apply(data.Decastar, 2, mad, na.rm=T)
sd.variables.OlympicGames <- apply(data.OlympicGames, 2, mad, na.rm=T)</pre>
```

Correlation

Motivation

We consider again the dataset decathlon2, after looking at the distributions of each variables measured in the dataset, we are now interested in possible relationships between the variables.

```
library(factoextra)
data(decathlon2)
```

Are those variables related or independent and can we quantify it? The measure of relatedness between variables is called correlation. Several types of correlation exist and do not measure exactly the same thing. This is what we are going to explore in this section.

Working example

In the previous section, 2 unrelated variables were simulated. In this example we will create a dataset by simulating 10 more or less correlated variables.

```
library(mvtnorm)
set.seed(352)
sample.size <- 60

correlations <- runif(45, -1, 1)
correlation.matrix <- matrix(0, 10, 10)
correlation.matrix[upper.tri(correlation.matrix, diag=FALSE)] <- correlations
correlation.matrix[lower.tri(correlation.matrix, diag=FALSE)] <- t(correlation.matrix)[lower.tri(corrediag(correlation.matrix) <- 1
standard.deviations <- rep(1,10)
covariance.matrix<- diag(standard.deviations) %*% correlation.matrix %*% diag(standard.deviations)

dataset<-as.data.frame(rmvnorm( sample.size, mean=rep(0,10), covariance.matrix))
colnames(dataset) <- paste('Variable', c(1:10), sep='')</pre>
```

By definition the correlation is a measure that ranges from -1 to 1 and is symmetrical and scale-invariant, i.e. multiplying a set of value of a variable by a positive number (except 0) will not impact the correlation of this variable with another variables. -1 and 1 represent perfect correlation while 0 represents absence of correlation. Having a negative correlation between two variables means that large values of one variable is associated with small values of another variable. These properties are similar for each type of correlation. Three main correlations are used.

The most common one is the Pearson correlation also called linear correlation and can be computed via the formula:

$$r = \frac{\sum_{i} (x_i - \overline{x})(y_i - \overline{y})}{\sum_{i} (x_i - \overline{x})^2 \sum_{i} (y_i - \overline{y})^2}$$

To compute the different correlations in r we can use the function cor:

```
#computation of the pearson correlation between Variable1 and Variable2
cor(dataset$Variable1,dataset$Variable2, method = 'pearson', use = 'complete.obs')
```

[1] 0.1912794

Here we can see that the correlation is negative, so high values of Variable1 correspond to small values of Variable2. The absolute value of the correlation 0.1912794 tells us that large variation to the mean in Variable1 does not necessarily mean the same amount of variation to the mean for Variable2.

In our dataset we have 10 variables and we don't want to test each pair of variables at a time. Hopefully, we can do it all at once:

```
#computation of the Pearson correlation matrix between all variables of the dataset
correlation.pearson <- cor(dataset, method = 'pearson', use = 'complete.obs')
correlation.pearson</pre>
```

```
##
              Variable1
                         Variable2
                                     Variable3
                                               Variable4
                                                          Variable5
## Variable1
             1.00000000 0.19127937 0.21752551 -0.33143225 -0.03092937
## Variable2
             0.19127937 1.00000000 -0.51763726 -0.01338385 -0.49874314
             0.21752551 \ -0.51763726 \ 1.000000000 \ -0.30236418 \ -0.07174202
## Variable3
## Variable4 -0.33143225 -0.01338385 -0.30236418 1.00000000 -0.03811203
## Variable5 -0.03092937 -0.49874314 -0.07174202 -0.03811203 1.00000000
## Variable6 -0.53182411 -0.35594681 0.04107634 0.14017732 0.58564298
## Variable7
             ## Variable8 -0.24406243 -0.28428073 0.11700153 -0.14327223 -0.33169360
## Variable9
             0.54566738 0.09543885 0.45465891 -0.58902336 -0.17757922
## Variable10 -0.32526916 -0.47961035 -0.13380258 0.25004362 0.45375074
##
              Variable6 Variable7 Variable8 Variable9 Variable10
```

```
## Variable1
             -0.53182411 0.50528333 -0.2440624 0.54566738 -0.3252692
## Variable2 -0.35594681 0.63096878 -0.2842807 0.09543885 -0.4796104
## Variable3
              0.04107634 -0.32475661 0.1170015 0.45465891 -0.1338026
## Variable4
              0.14017732 \quad 0.43996154 \quad -0.1432722 \quad -0.58902336
                                                              0.2500436
## Variable5
              0.58564298 -0.33488993 -0.3316936 -0.17757922
                                                              0.4537507
## Variable6
              1.00000000 -0.55375759 -0.5352634 -0.10179082 0.6954264
              -0.55375759 1.00000000 -0.2478753 0.02081113 -0.4821698
## Variable7
              -0.53526337 -0.24787528 1.0000000 -0.26895642 -0.3316504
## Variable8
## Variable9
              -0.10179082 0.02081113 -0.2689564 1.00000000 -0.2662986
## Variable10 0.69542644 -0.48216985 -0.3316504 -0.26629860 1.0000000
```

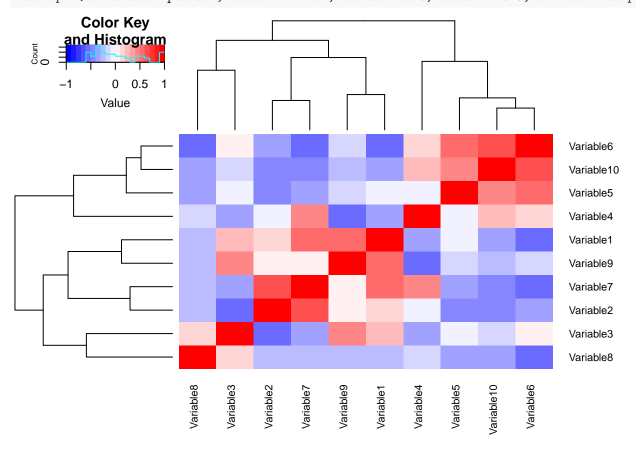
A matrix of correlation as we just obtained can provide a lot of information about the variables. However, for illustration in a paper it is better to use a representation of the correlation matrix called heatmap:

```
#Heatmap plot
library(gplots)

## Attaching package: 'gplots'

## The following object is masked from 'package:stats':

## lowess
heatmap.2(correlation.pearson, trace = 'none', cexRow = 0.8, cexCol = 0.8, col= colorRampPalette(c("blue"))
```



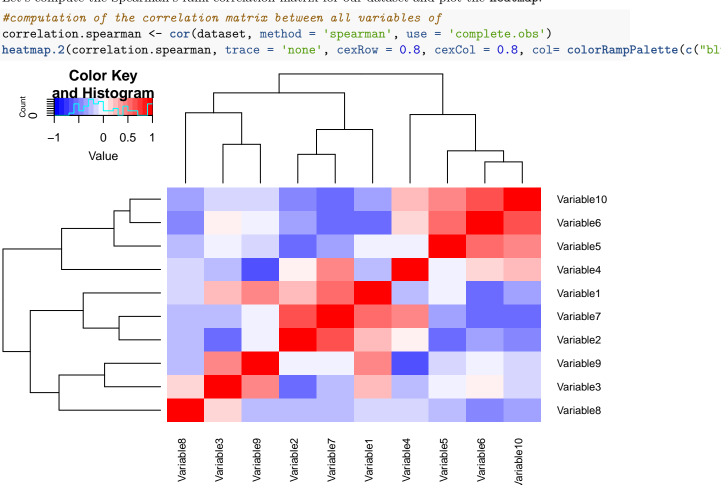
Here, the correlation is represented by a 'heat', blue or 'cold heat' represent negative correlations, red or 'hot heat' represent strong correlations.

The tree produced by the function heatmap on each side of the plot represent the similarity between the

variables based on their correlation matrix. Each variable has a root, all roots are merging to form the base of the tree. The height were the roots of 2 variables are merging represent the dissimilarity between them. The higher they meet the more dissimilar they are.

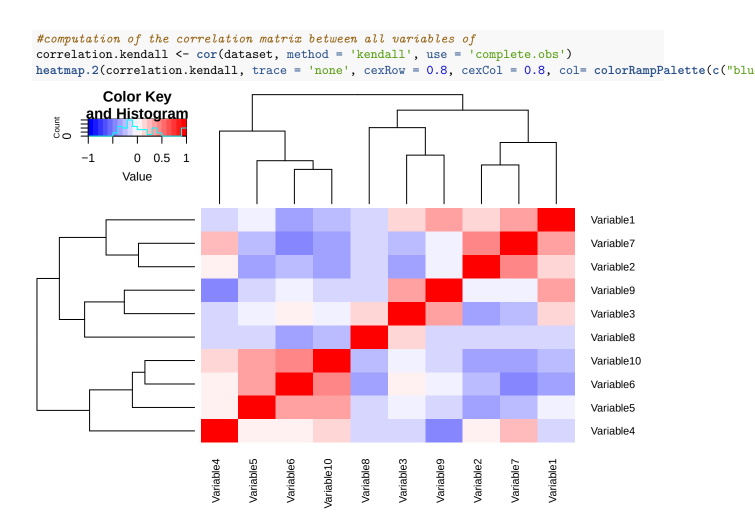
One weakness of the Pearson correlation is that the relationship measured between two variables X and Y is linear, i.e., their relationship is of the form X = aY + b. However, more complex relationship can exist between variables. This is why using different correlation measures call non-parametric is interesting as it can help identify different relationship between variables. However, the interpretation of this correlation can be challenging.

The first non-parametric correlation that we will see is called the Spearman's rank correlation often denoted by ρ . The computation is similar to the Pearson correlation but the values are replaced by their rank. Meaning if we have a vector of values (7,2,5), it will be replaced by (1,3,2) before computing the correlation. Let's compute the Spearman's rank correlation matrix for our dataset and plot the heatmap.



Here as I have only created linearly correlated variables not much difference can be seen between both heatmaps.

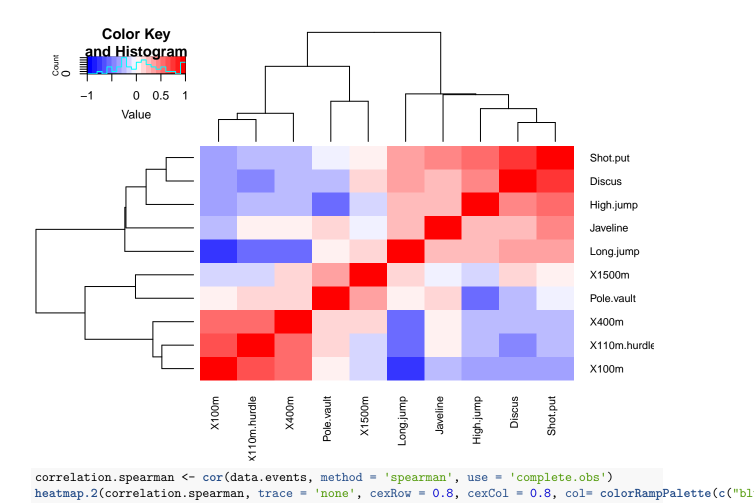
The third correlation is the Kendall's tau represented by τ . Let's consider 2 variables X and Y having 3 observations, (0,1,2) and (3,4,1) respectively. The 3 observations of the 2 variables are defined as (0,3), (1,4), and (2,1). The Kendall's tau computation consist of looking at each pair of observation and counting the number of pair observation concordant and the number of pair discordant. The correlation is then computed as the number of pair concordant minus the number of pair discordant divided by the total number of pairs. This approach has the only advantage to be more interpretable than the Spearman's rank correlation.

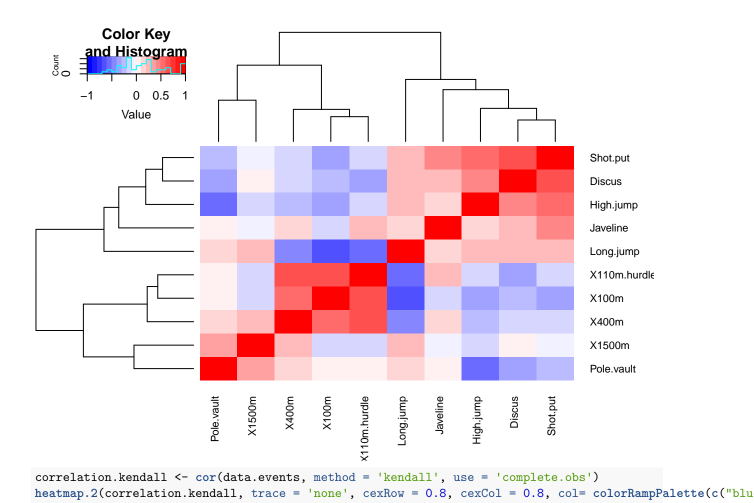


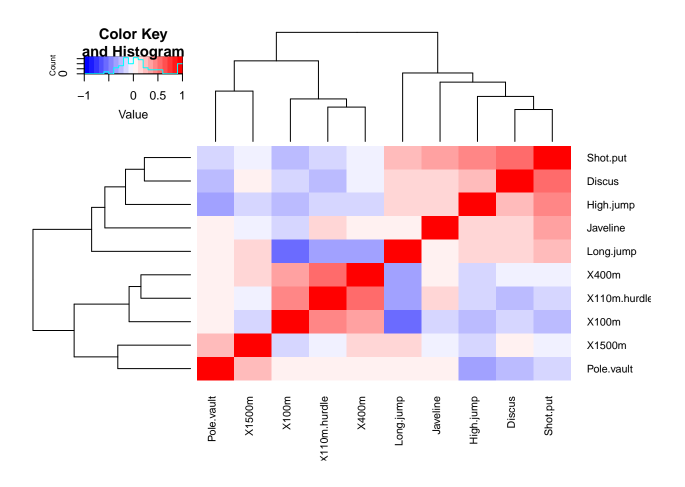
Quick tasks

Compute the correlation matrix between the events of the decathlon in the dataset decathlon2 using the Pearson correlation, the Spearman's rank correlation and Kendall's tau. Display the results on heatmaps.

```
data.events <- decathlon2[,c(1:10)]
correlation.pearson <- cor(data.events, method = 'pearson', use = 'complete.obs')
heatmap.2(correlation.pearson, trace = 'none', cexRow = 0.8, cexCol = 0.8, col= colorRampPalette(c("blu</pre>
```







Principal components analysis

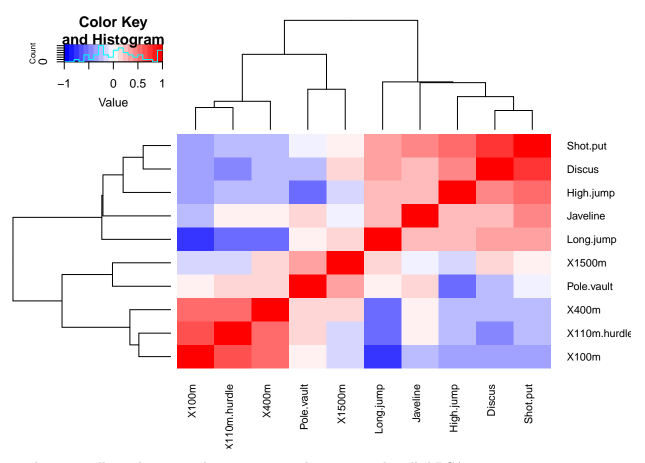
Motivation

After having an overview of the relationship between the variables of the dataset decathlon2, the searcher wants to extract more information and a better visualization from this correlation structure form both samples and variables.

```
library(FactoMineR)
library(gplots)
library(factoextra)

data(decathlon2)

data.events <- decathlon2[,c(1:10)]
correlation.pearson <- cor(data.events, method = 'pearson', use = 'complete.obs')
heatmap.2(correlation.pearson, trace = 'none', cexRow = 0.8, cexCol = 0.8, col= colorRampPalette(c("blue"))</pre>
```



To do so we will use the principal components analysis commonly called PCA.

Working example

We will use the example that we created in the previous section on correlation:

```
library(mvtnorm)
set.seed(352)
sample.size <- 60

correlations <- runif(45, -1, 1)
correlation.matrix <- matrix(0, 10, 10)
correlation.matrix[upper.tri(correlation.matrix, diag=FALSE)] <- correlations
correlation.matrix[lower.tri(correlation.matrix, diag=FALSE)] <- t(correlation.matrix)[lower.tri(corrediag(correlation.matrix) <- 1
standard.deviations <- rep(1,10)
covariance.matrix<- diag(standard.deviations) %*% correlation.matrix %*% diag(standard.deviations)
dataset<-as.data.frame(rmvnorm( sample.size, mean=rep(0,10), covariance.matrix))
colnames(dataset) <- paste('Variable', c(1:10), sep='')</pre>
```

The principle of principal components analysis relies heavily on mathematical algebra that we won't detail in this course. The main idea of principal components analysis is to identify new variables carrying most of the variance existing in your dataset. These new variables are created by linear combinations of the variables available in the dataset and can be used to identify hidden patterns, dimension reduction and the correlated variables in your dataset.

In this example we can see that by redefining the variables and using the principal component 1 (a linear combination of variable x and y) we capture most of the variation.

The principal component analysis consist of finding the principal components and by looking at contribution of each variables on this principal components understanding the structure of the data.

We will perform a principal component analysis on the dataset we simulated. This can be done by using the function PCA of the package FactoMineR.

Note that to perform a PCA, it is important to scale the data. Indeed, the PCA results can be strongly influenced by different variances.

```
pca.analysis <- PCA(dataset, scale.unit = TRUE, graph = FALSE)</pre>
```

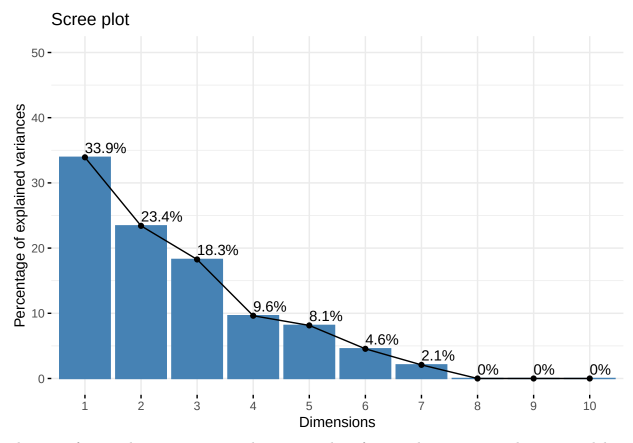
Typically there is a number of principal components equal to the number of variables in your dataset. However, not all of them retain the same amount of variance. The amount of variation retained by each component is called eigenvalue and can help us to determine the number of principal components needed to describe properly the data. By definition the principal component 1 is the one retaining most variance.

```
eigenvalues <- get_eigenvalue(pca.analysis)
eigenvalues</pre>
```

```
eigenvalue variance.percent cumulative.variance.percent
## Dim.1 3.392355e+00
                           3.392355e+01
                                                           33.92355
                                                           57.33613
## Dim.2 2.341258e+00
                           2.341258e+01
## Dim.3 1.825827e+00
                           1.825827e+01
                                                           75.59440
## Dim.4 9.626093e-01
                           9.626093e+00
                                                           85.22049
## Dim.5 8.140583e-01
                           8.140583e+00
                                                           93.36108
## Dim.6 4.551795e-01
                           4.551795e+00
                                                           97.91287
## Dim.7 2.087128e-01
                           2.087128e+00
                                                           100.00000
## Dim.8 1.114076e-31
                           1.114076e-30
                                                           100.00000
## Dim.9 6.117524e-32
                           6.117524e-31
                                                           100.00000
## Dim.10 4.155684e-32
                           4.155684e-31
                                                           100.00000
```

We see that only 5 principal components are needed to capture 93.36% of the total variance present in the dataset. This can be better seen via the scree plot:

```
fviz_eig(pca.analysis, addlabels = TRUE, ylim = c(0, 50))
```



There is unfortunately, no consensus on the exact number of principal components to keep. 3 possibilities can be used to select the number of principal components. The first one is to select the number of principal components to retain a sufficient amount of variance present in the dataset such as 80%. In our case 4 principal components

The second approach is to look at the scree plot to determine the number of components. The number of component is determined at the point, beyond which the remaining eigenvalues are all relatively small and of comparable size. In our case 4 principal components.

Finally the third approach is to look at the eigenvalues of the different principal components and to keep the principal components having a eigenvalue higher than 1. Indeed an eigenvalue higher than 1 indicates that the related principal components have more variance than one of the original variables. In our case 4 principal components.

Now that we have obtain the number of principal components to keep, we can use the principal component analysis as a dimension reduction approach and keep these new variables for statistical models. But we can also investigate the contribution of these variables in each components in order to understand better the relationship between our variables in the dataset.

To do so we need to extract the results for the variables from our PCA.

```
pca.analysis <- PCA(dataset, ncp=4, scale.unit = TRUE, graph = FALSE)
variable.analysis <- get_pca_var(pca.analysis)
variable.analysis</pre>
```

```
## 3 "$cos2" "Cos2 for the variables"
## 4 "$contrib" "contributions of the variables"
```

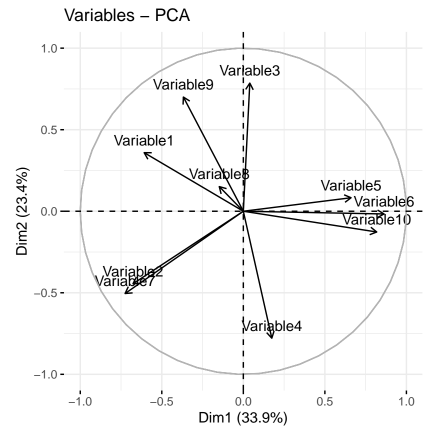
head(variable.analysis\$coord, 4)

```
##
                 Dim.1
                          Dim.2
                                     Dim.3
                                               Dim.4
## Variable1 -0.60818282
                      0.3592227
                                0.47779077
                                           0.3806998
## Variable2 -0.67661752 -0.4446899
                                0.28175127 -0.4878183
## Variable3
           0.3549273
## Variable4
            0.17456857 -0.7784345 -0.03390743
                                           0.4826220
```

The different available results are the coordinates of the variables in the new dimension (important to compute the values of the observation for the new variables) and the contribution of each variable to the different principal components.

With this information, we can create plots to understand the relationship between the variables and the principal components such as the correlation circle.

```
#Plot correlation circle
fviz_pca_var(pca.analysis, col.var = "black", axes = c(1,2))
```



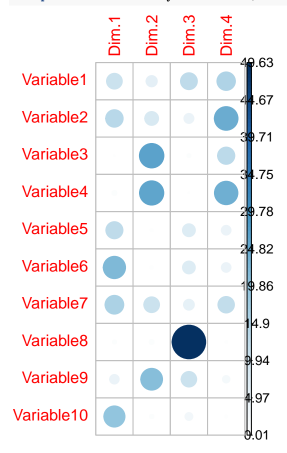
This plot shows the relationship existing between the variables. Positively correlated variables will be grouped together whereas negatively correlated variables will be on the opposite side of the plot origin. The closest the variable is from an axis, the more correlated this variable is with the principal component. Finally, the longer the arrow is the better represented this variable is by the two principal components.

We can see also by plot the amount of contribution that each variable provide to each principal component:

```
library("corrplot")
```

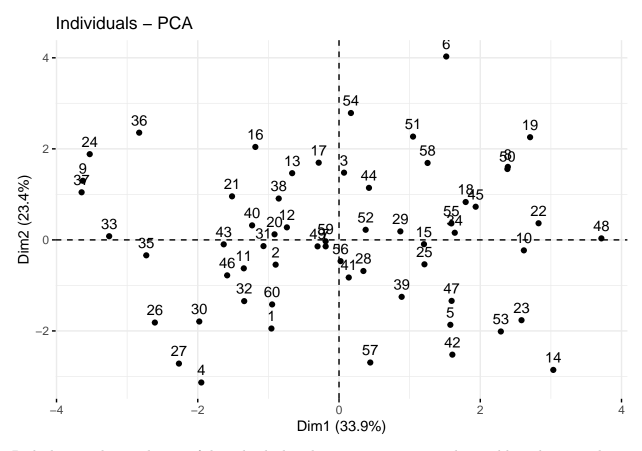
corrplot 0.84 loaded

corrplot(variable.analysis\$contrib, is.corr=FALSE)



Mow that we looked at the relationship existing between the variables and the principal components we can do the same thing for the individuals.

```
#Scatterplot in the 2 first principal components plane
fviz_pca_ind(pca.analysis, col.var = "black", axes = c(1,2))
```



By looking at the coordinates of the individual in the new space we can understand how these samples are clustered. By then looking at the contribution of each variable for the principal components we can then derive conclusion on the relatedness of the individuals based on the original variables.

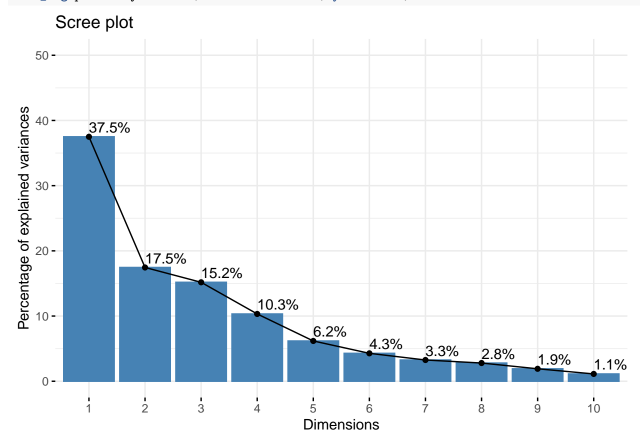
Quick tasks

Perform a principal component analysis on the events of the decathlon2 dataset. Print the eigenvalues table and make a scree plot to determine the number of principal components to retain.

```
data.events <- decathlon2[,c(1:10)]
pca.analysis.deca <- PCA(data.events, scale.unit = TRUE, graph = FALSE)
eigenvalues <- get_eigenvalue(pca.analysis.deca)
eigenvalues</pre>
```

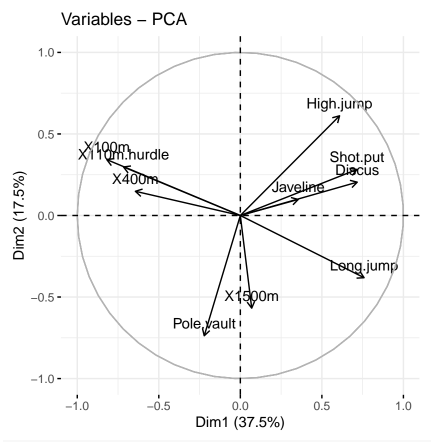
```
##
          eigenvalue variance.percent cumulative.variance.percent
## Dim.1
           3.7499727
                             37.499727
                                                            37.49973
## Dim.2
           1.7451681
                             17.451681
                                                            54.95141
## Dim.3
           1.5178280
                             15.178280
                                                            70.12969
                                                            80.45169
## Dim.4
           1.0322001
                             10.322001
## Dim.5
           0.6178387
                              6.178387
                                                            86.63008
## Dim.6
           0.4282908
                              4.282908
                                                            90.91298
## Dim.7
           0.3259103
                              3.259103
                                                            94.17209
## Dim.8
           0.2793827
                              2.793827
                                                            96.96591
## Dim.9
           0.1911128
                              1.911128
                                                            98.87704
## Dim.10
                                                           100.00000
           0.1122959
                              1.122959
```

fviz_eig(pca.analysis.deca, addlabels = TRUE, ylim = c(0, 50))

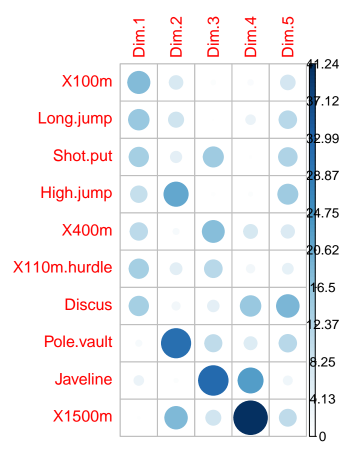


Plot the correlation circle of the first two principal components as well as the contribution plot

```
pca.analysis.deca <- PCA(data.events, ncp=5, scale.unit = TRUE, graph = FALSE)
variable.analysis.deca <- get_pca_var(pca.analysis.deca)
fviz_pca_var(pca.analysis.deca, col.var = "black", axes = c(1,2))</pre>
```

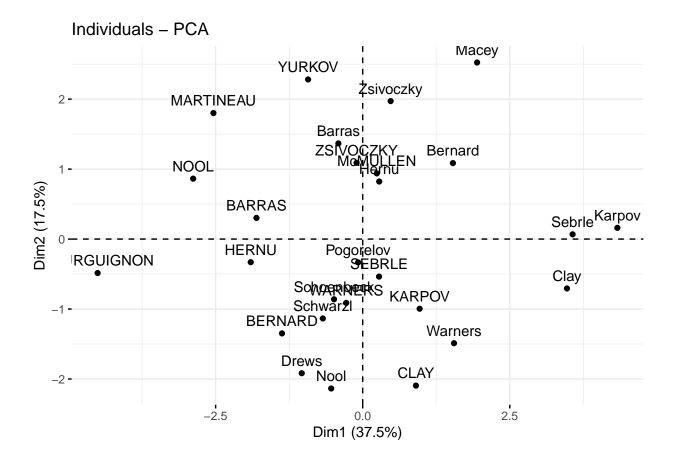


corrplot(variable.analysis.deca\$contrib, is.corr=FALSE)



What can you conclude from the first two principal components? What are they mostly measuring? Plot the representation of the individuals on these two principal components

fviz_pca_ind(pca.analysis.deca, col.var = "black", axes = c(1,2))



Exercises

Exercise 1

Load the dataset mtcars in the R memory using the command data(mtcars).

- 1.1 How many variables and observations are contained in the dataset? Print the 5 first rows of the dataset
- 1.2 Plot the histograms of the continuous numerical variables of the dataset (type ?mtcars to obtain the description of the dataset).
- 1.3 We can clearly distinguish 3 right skewed distribution. Which are they?
- 1.4 The variables vs and am both split the data in 2 categories plot the boxplots of the variable qsec for each group define by vs and am:
- 1.5 Plot boxplots using the command boxplot(qsec ~ vs:am). What is the difference with the plots obtained previously?
- 1.6 compute the mean and standard variation for each continuous variable of the dataset for the 2 different engine types.

Exercise 2

We continue with the same dataset:

2.1 compute the correlation matrix associated for the three different correlations for all variables except the vs and am variables.

2.2 Display the results using heatmaps.

Exercise 3

- 3.1 Perform a pca analysis and plot the resulting scree plot.
- 3.2 display the eigenvalues of the oca analysis. How many principal components would you keep?
- 3.2 Plot the correlation circle for the first two principal components. What is your interpretation of the first principal component?
- 3.3 Display the contribution of each variable on the principal components

Basic statistical tests

Statistical distributions

Motivation

In this section we will describe some of the most commonly used distributions used in statistics to describe the distribution of a random variable. Indeed it is important to learn to recognize some distributions and some of their properties to better use statistical modes and test in your analysis.

Discrete random variables

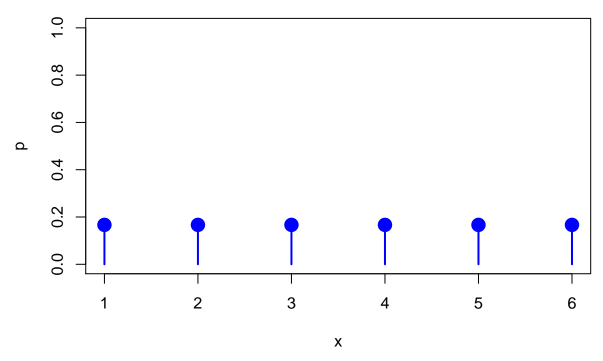
Discrete distribution variables are distributions taking only integer values such as the uniform distribution, the Bernoulli distribution and the Binomial distribution.

The discrete Uniform distribution is a symmetric distribution where a set of finite values have the exact same probabilities to occur. A common example is throwing a fair dice. After throwing the dice, each numbers 1, 2, 3, ..., 6 has the same probability to be obtained, equal 1/6.

```
#plotting probability discrete uniform distribution

x <- c(1:6)
y <- rep(1/6,6)

plot(x,y,type="h",xlim=c(1,6),ylim=c(0,1),lwd=2,col="blue",ylab="p")
points(x,y,pch=16,cex=2,col="blue")</pre>
```

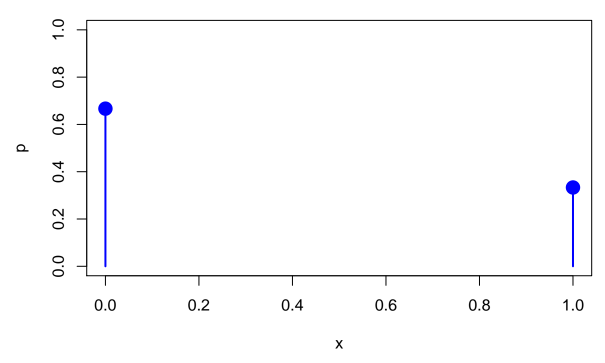


The Bernoulli distribution is a distribution taking as values 0 or 1 with probabilities equal to p or q respectively. With q = 1-p. For example, let's throw a fair dice and check if the number is equal to or lower than 2. The probability of a success (obtaining a value below or equal 2) generally noted 1 is 1/3 while the probability of not succeeding is 2/3. The analysis of the association between covariates and the probability of success is often done via logistic regression, which we will see in Chapter 3.

```
#plotting probability bernoulli distribution

x <- c(0:1)
y <- c(2/3,1/3)

plot(x,y,type="h",xlim=c(0,1),ylim=c(0,1),lwd=2,col="blue",ylab="p")
points(x,y,pch=16,cex=2,col="blue")</pre>
```

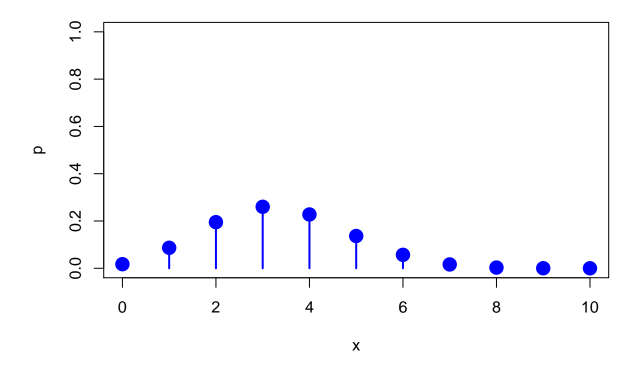


The binomial distribution is describing the results of several trials. The probability describes the chance to obtain k successes on n trials. It is by definition the sum of several Bernoulli. If we go back to the previous dice example we can compute the probability to obtain 4 success in ten trials. This can be done with the functiondbinom. The probability is 0.2276076.

```
#plotting probability binomial distribution

x <- c(0:10)
y <- dbinom( x, 10, 1/3)

plot(x,y,type="h",xlim=c(0,10),ylim=c(0,1),lwd=2,col="blue",ylab="p")
points(x,y,pch=16,cex=2,col="blue")</pre>
```

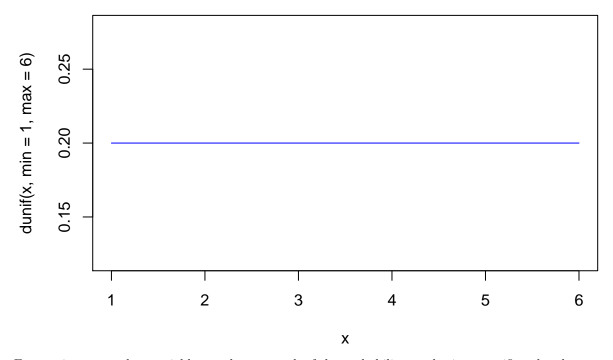


Continuous random variables

A continuous random variable is a random variable with an infinite possibility of values being drawn from it. As we have an infinite possible values, the probability to obtain one value in particular is equal to 0. This is why we do not look at the probability directly but instead we look at the density of drawing values around a specific value in an infinitesimal space around the value. We will now look at some important continuous density distributions.

The continuous uniform distribution has a similar definition than the discrete uniform distribution. The density is the same for each values that can be drawn from it. Therefore, this distribution is defined by the range of the possible values that are drawn from it. Let's plot the density of a continuous uniform distribution between 1 and 6.

```
#Plot the density
curve(dunif(x, min= 1, max= 6), col="blue", from = 1, to = 6)
```

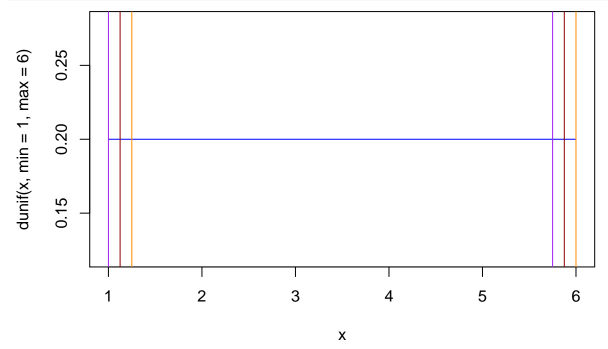


For continuous random variables we do not speak of the probability to obtain a specific value, but we speak of the probability that the value is in a specific range. Let's add the quartiles of the uniform distribution to the plot.

By definition we now that the quartiles split the data into 4 equal parts. So the probability of obtaining a value between 2 consecutive quartiles is 25%. With the same reasoning, the probability of obtaining a value between the first and the third quartile is equal to 50%. Typically, the probability to draw a value from a distribution between a set of 2 values is equal to the area under curve between these two points. By using

the quantiles we can then define an area where a probability to draw a values is equal to 95%.

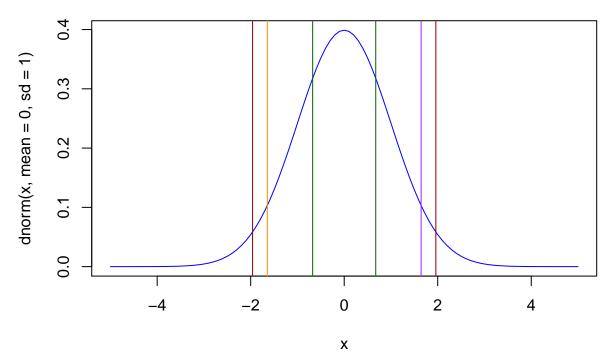
```
#Plot of the density and important quantiles of a continuous uniform distribution
curve(dunif(x, min= 1, max= 6), col="blue", from = 1, to = 6)
abline(v = qunif(c(0.025, 0.975), min= 1, max= 6), col= 'darkred')
abline(v = qunif(c(0, 0.95), min= 1, max= 6), col= 'purple')
abline(v = qunif(c(0.05, 1), min= 1, max= 6), col= 'darkorange')
```



Quantiles are very important in statistical testing at they allow us to draw conclusions such as the confidence intervals or the p-values by looking at the value of the test statistic vs the theoretical distribution of the test.

The most common distribution encountered in the nature is the normal or gaussian distribution. This distribution is symmetrical and is defined by its expectation often denoted μ and it variance σ^2 , i.e. $N(mu, \sigma^2)$. Its values ranges from $-\infty$ to ∞ . Note that it is important to moake the difference between the expectation of the distribution and the sample mean derived from this distribution. Indeed, these two will be equal only if all possible samples are included to compute the sample mean.

```
#Plot of the density and quantiles of a normal distribution
curve(dnorm(x, mean = 0, sd = 1), col = 'blue', from = -5, to = 5)
abline(v = qnorm(c(0.25, 0.75), mean = 0, sd = 1), col = 'darkgreen')
abline(v = qnorm(c(0.025, 0.975), mean = 0, sd = 1), col = 'darkred')
abline(v = qnorm(c(0, 0.95), mean = 0, sd = 1), col = 'purple')
abline(v = qnorm(c(0.05, 1), mean = 0, sd = 1), col = 'darkorange')
```

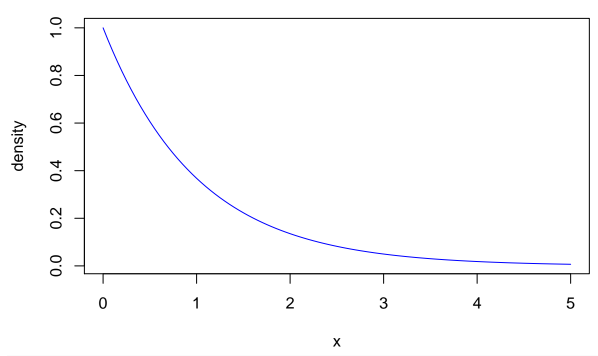


This distribution is very important in statistics due to one of its porperties, the central limit theorem. It states that, if you have a population with mean μ and standard deviation σ and take sufficiently large random samples from the population with replacement, then the distribution of the sample means will be approximately normally distributed. This will hold true regardless of whether the source population is normal or skewed, provided the sample size is sufficiently large (usually n > 30) and the variance of the original distribution is finite.

Let's illustrate the central limit theorem with the gamma distribution that will be seen at another chapter of the course.

```
# Example of the central limit theorem

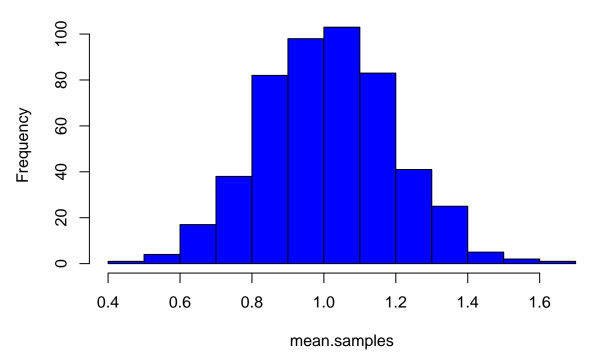
#Plot of the density of a gamma distribution
curve(dgamma(x, shape = 1, rate = 1), from = 0, to = 5, col= 'blue', ylab= 'density')
```



```
#simulation of several samples of size 30 from the same gamma distribution and computation of their mea
mean.samples <- NULL
for (k in 1:500){
   data.sample <- rgamma( 30 , shape = 1, rate = 1)
   mean.samples <- c(mean.samples, mean(data.sample))
}

#Histogram of the mean of the samples
hist(mean.samples, col='blue')</pre>
```

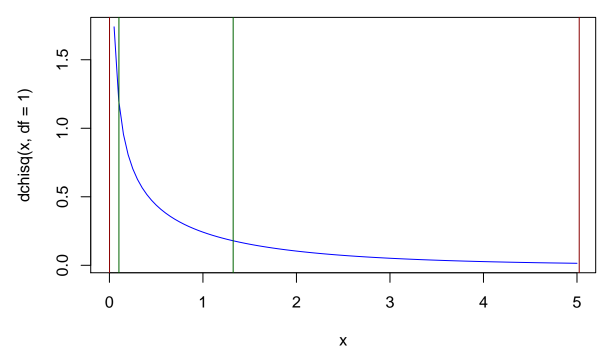
Histogram of mean.samples



As most of the phenomenon are the sum of little random variations it is then logical to see a lot of phenomenon following a gaussian distribution.

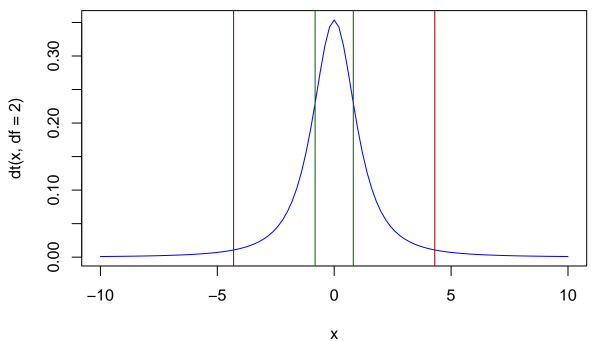
The chi square distribution, noted χ^2 with k degrees of freedom, is defined at the sum of k squared normal distributions having expectancy equal to 0 and variance equal to 1. This distribution is often used for statistical testing.

```
#Plot of the density and quantiles of a chi square distribution with 1 degree of freedom
curve(dchisq(x, df = 1), col = 'blue', from = 0, to = 5)
abline(v = qchisq(c(0.25, 0.75), df = 1), col= 'darkgreen')
abline(v = qchisq(c(0.025, 0.975), df = 1), col= 'darkred')
```



Finally a last important continuous random variable for statistical testing is the Student's t distribution as it is used in the so called Student's t test to compare means and for the obtention of confidence interval. This distribution is defined by its number of degree of freedom k and is the quotient of a normal distribution having expectancy equal to 0 and variance equal to 1 with the square root of a chi square distribution with k degree of freedom. This distribution, as the normal distribution, is symmetrical.

```
#Plot of the density and quantiles of a chi square distribution with 2 degree of freedom curve(dt(x, df = 2), col = 'blue', from = -10, to = 10) abline(v = qt(c(0.25, 0.75), df = 2), col = 'darkgreen') abline(v = qt(c(0.025, 0.975), df = 2), col = 'darkred')
```



This distribution has the same bell shape than the normal distribution. However, this shape is narrower and

becomes more and more narrow as the number of degrees of freedom increases.

Quick Tasks

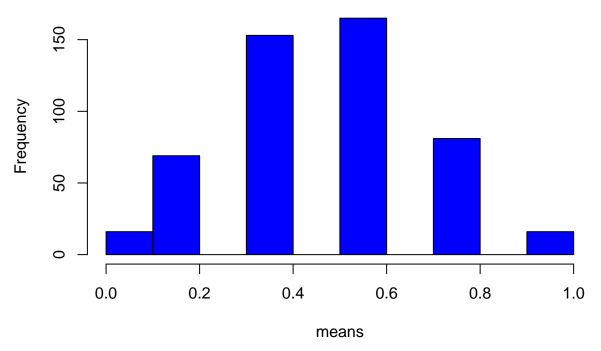
A student of statistics wants to make a small experiment. He throws a coin 50 times and counts the number of times he gets a head. Compute the probability to obtain head 20, 25 and 35 times using the function dbinom. Plot the probability plot resulting from this experiment.

The student creates an experiment protocol where he will throws 5 times the coin and counts the number of times he gets head. He will repeat this experiment 500 times. Simulate each experiment using the rbinom function and compute their mean. Plot the histogram of the 500 means obtained.

c(0:50)

```
means<-NULL
for (k in 1:500){
sample <- rbinom(5, size = 1, prob =0.5)
means <- c(means, mean(sample))
}
hist(means,col="blue")</pre>
```

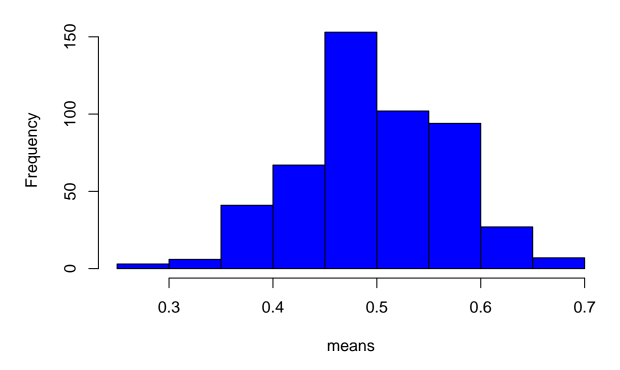
Histogram of means



Not satisfied by his results, the student recruits 9 other students to do the same experiment but this time throwing 50 times the coins for each experiment. Each experiment will be repeated 500 times. Simulate and plot the results.

```
means<-NULL
for (k in 1:500){
sample <- rbinom(50, size = 1, prob =0.5)
means <- c(means, mean(sample))
}
hist(means,col="blue")</pre>
```

Histogram of means



Two-group comparisons

Motivation

We consider again the decathlon2 dataset. After exploring the dataset and understanding its structure, we want to test the performances between the two different events, i.e. the Decastar and the OlympicGames.

```
library(factoextra)
data(decathlon2)
```

When comparing the distribution of a of values from a continuous variable of two groups we commonly compare the mean of the set of values obtained in both groups. Sample means could look different but are the samples really drawn from 2 different distributions?

Working Example

Let us consider a dataset containing two continuous variable that we will call VariableX and VariableY and a categorical variables indicating which groups samples are from.

```
set.seed(352)
sample.size<-30

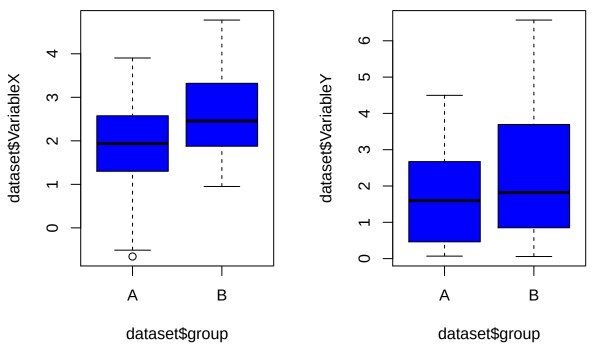
variableX.groupA <- rnorm(sample.size, mean = 2, sd = 1)
variableX.groupB <- rnorm(sample.size, mean = 2.4, sd = 1.1)

variableY.groupA <- rgamma(sample.size, shape = 2, scale = 1)
variableY.groupB <- rgamma(sample.size, shape = 2, scale = 1.1)</pre>
```

```
group <- rep(c('A', 'B'), each = sample.size)
dataset <- data.frame(VariableX = c(variableX.groupA, variableX.groupB), VariableY = c(variableY.groupA</pre>
```

We start by plotting the boxplots for each group of both variables.

```
#Boxplots for variable X and Y
par(mfrow=c(1,2))
boxplot(dataset$VariableX ~ dataset$group, col = 'blue')
boxplot(dataset$VariableY ~ dataset$group, col = 'blue')
```



We can identify by 'eye' differences between the distribution of both group but how can we quantify the differences if there is any? To do so we will use statistical tests.

The most common test to compare two groups is the Student's t test, this test as the same name as the Student's t distribution as it utilizes the same distribution. Let's assume that two sets of values are drawn from distributions with expectancies μ_1 and μ_2 , and variances σ_1^2 and σ_2^2 , respectively. The Student's test statistic is computed as follows:

$$t = \frac{\overline{x_1} - \overline{x_2}}{SEDM}$$

where the standard error of difference of means is:

$$SEDM = \sqrt{SEM_1^2 + SEM_2^2}$$

The square root of the sum of the squared standard errors to mean of each group. The standard error to the mean can itself be computed as the standard deviation divided by the square root of the number of samples.

$$SEM_1 = \frac{\sigma_1}{\sqrt{n}}$$

When using a statistical test two hypotheses are formulated, the null hypothesis H_0 and the alternative hypothesis H_a , respectively. In the case of the Student's t-test, the hypotheses are:

$$H_0: \mu_1 = \mu_2 \text{ and } H_a: \mu_1 \neq \mu_2$$

Note that there can be 3 different alternative hypothesis depending on what we want to test. The test for alternative hypothesis presented above is called the *two sided* Student's t test as we are testing for different

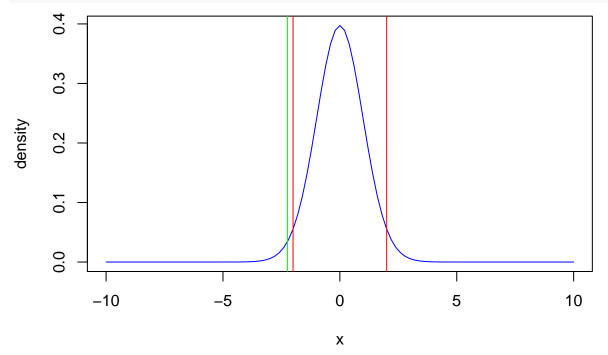
means but we are not making any assumptions on which direction the differences is. Two other alternative can be tested. Namely, $H_a = \mu_1 < \mu_2$, and $H_a = \mu_1 > \mu_2$. Under the null hypothesis the variable t follows a Student's distribution of degrees of freedom equal to n1 + n2 - 2 if the variances σ_1^2 and σ_2^2 are equal. If this is not the case the statistics t can still be approximated as a Student's t distribution with a number of degrees of freedom given by the Welch procedure that we won't detail here.

Let's apply the Student's test to our dataset:

```
#t-test for VariableX
test.variableX <- t.test(dataset$VariableX ~ dataset$group)</pre>
test.variableX
##
##
    Welch Two Sample t-test
##
## data: dataset$VariableX by dataset$group
## t = -2.2435, df = 57.612, p-value = 0.02872
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
   -1.20392326 -0.06848317
## sample estimates:
## mean in group A mean in group B
##
          1.900228
                           2.536431
```

Let's look at the results obtained. We can see that the estimated means are equal to 1.9002279 and 2.5364311 in group A and group B respectively. This lead to an estimated difference in mean of -0.6362032. The computed test statistic is equal to -2.2435031. The number of degrees of freedom estimated is equal to 57.6120941. We can then plot the density distribution of this Student's t random variable.

```
curve(dt(x, df = test.variableX$parameter), col = 'blue', from = -10, to = 10, ylab = 'density')
abline(v = qt(c(0.025, 0.975), df = test.variableX$parameter), col = 'red')
abline(v = test.variableX$statistic, col= 'green')
```



As we can see the value of the statistics is outside the range delimited by the 2.5% quantile and the 97.5% quantile. Therefore it is unlikely that the statistics is drawn from such random variable. It allows us to reject

the null hypothesis and accept the alternative hypothesis with less than 5% chance risk of making a mistake. The p-value obtained from the test is the probability that the statistics is drawn from the hypothetical Student's t distribution under the null hypothesis from quantiles table for example.

An important output information from the output is also the 95% confidence interval. This interval quantifies the uncertainty of the estimate of the mean difference. The 95% confidence interval is a range of values that you can be 95% certain contains the true mean difference between the two populations. As we can see 0 is not in this range allowing us to, once again, reject the null hypothesis of equality of the means. The confidence interval can be computed by using the SEDM and the mean difference obtained in the test. For a level of confidence α of a two sided Student test is give by the formula:

```
high Boundary = mean + t_{1-\frac{\alpha}{2}}*SEDM low Boundary = mean - t_{1-\frac{\alpha}{2}}*SEDM
```

We can see from this formula that two factors can influence the size of the confidence interval. Indeed, the SEDM is a function depending on the standard deviation and the number of samples in each group. On one hand, increasing the number of sample will improve the accuracy of the test and narrow the confidence interval. On the other hand, samples with large standard deviation will lead to larger confidence intervals. We can compute the confidence intervals of the test we made earlier:

```
low.boundary <- test.variableX$estimate[1] - test.variableX$estimate[2] - qt(0.975,df = test.variableX$high.boundary <- test.variableX$estimate[1] - test.variableX$estimate[2] + qt(0.975,df = test.variableX$c(low.boundary, high.boundary)
```

```
## mean in group A mean in group A
## -1.20392326 -0.06848317
```

The Student's t test is a great tool to compare to distributions. However, it has its weaknesses. In case of skewed data, the standard deviation of both compared samples can be relatively different and impact the power of the test to detect relevant differences. In such cases, it is preferable to transform the data using for example a log transformation. For the Student's t test it is requested that the distributions of the samples values are following a normal distribution or to have at least a sample size of 30 in each group (central limit theorem). If this is not the case, it is better to use a non parametric test called Mann-Whitney-Wilcoxon test (or Mann-Whitney U test or Wilcoxon sum rank test).

The Mann-Whitney-Wilcoxon test is non parametric as it does not test a specific parameter to differentiate two populations. Instead we will test if the probability that an observation in group A is superior to an observation from group B is different from the probability that an observation from group B is superior from group A. The main assumption of the Mann-Whitney-Wilcoxon test is that the samples from both groups must be independent so it cannot be used for paired observations. For paired observations we would prefer the Wilcoxon signed-rank test that we won't detail here. If the observations in group A are drawn from a random variable X and the observation in group B are coming from a random variable Y the null and alternative hypotheses are:

$$H_0: P(X > Y) = P(Y > X) \text{ and } H_0: P(X > Y) \neq P(Y > X)$$

To simplify the test relies on comparing the medians of both populations. The statistic of the test called U is computed as follows. All observations are pooled together and ranked based on their values. For example if we have a pooled set of 6 distinct values (2, 4, 4, 4, 1, 5), the ranked version of this set of observations becomes (2, 4, 4, 4, 1, 6). Let's call R_1 . We can then compute U as:

$$\mathbf{U}=R_1$$
 - $\frac{(n_1)(n_1+1)}{2}$

With this statistics U we can then compute the p-value using either a Wilcoxon Rank-Sum Table in case of small samples and approximating U as a normal distribution for large samples:

```
wilcox.test(dataset$VariableY ~ dataset$group, alternative = "two.sided")
```

```
##
## Wilcoxon rank sum test
##
```

```
## data: dataset$VariableY by dataset$group
## W = 358, p-value = 0.1774
## alternative hypothesis: true location shift is not equal to 0
```

We can notice that the output of the Wilcoxon rank-sum test is smaller than for the Student's t test. Indeed as it is a non parametric test it does not provide direct effect size or confidence interval. However, relying only on the p-value is not correct and one would prefer to report also the difference between the median of both samples.

Quick Tasks

Perform the Student's t test and the Wilcoxon rank-sum test to compare the performance for each events and the total number of points between both events in the decathlon2 dataset. Do you see any differences between the test?

```
for (k in colnames(decathlon2)[c(1:10,12)]){
  print(k)
  formula<-as.formula(paste('decathlon2$', k, ' ~ decathlon2$Competition', sep = ''))</pre>
  t.test.deca <- t.test(formula)</pre>
  print(t.test.deca)
  wilcox.test.deca <- wilcox.test(formula, alternative = "two.sided", exact =F)</pre>
  print(wilcox.test.deca)
}
## [1] "X100m"
##
##
   Welch Two Sample t-test
## data: decathlon2$X100m by decathlon2$Competition
## t = 4.1121, df = 22.882, p-value = 0.0004294
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 0.1754888 0.5309947
## sample estimates:
## mean in group Decastar mean in group OlympicG
                                         10.82214
##
                 11.17538
##
##
    Wilcoxon rank sum test with continuity correction
##
##
## data: decathlon2$X100m by decathlon2$Competition
## W = 158, p-value = 0.001247
## alternative hypothesis: true location shift is not equal to 0
## [1] "Long.jump"
##
##
   Welch Two Sample t-test
##
## data: decathlon2$Long.jump by decathlon2$Competition
## t = -2.1271, df = 24.993, p-value = 0.04346
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.444590146 -0.007168095
## sample estimates:
## mean in group Decastar mean in group OlympicG
```

```
7.247692
##
                                        7.473571
##
##
##
  Wilcoxon rank sum test with continuity correction
## data: decathlon2$Long.jump by decathlon2$Competition
## W = 54.5, p-value = 0.08042
## alternative hypothesis: true location shift is not equal to 0
##
## [1] "Shot.put"
## Welch Two Sample t-test
## data: decathlon2$Shot.put by decathlon2$Competition
## t = -2.5529, df = 24.572, p-value = 0.01729
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -1.3415989 -0.1429066
## sample estimates:
## mean in group Decastar mean in group OlympicG
##
                 14.15846
                                        14.90071
##
##
## Wilcoxon rank sum test with continuity correction
##
## data: decathlon2$Shot.put by decathlon2$Competition
## W = 41, p-value = 0.01627
\#\# alternative hypothesis: true location shift is not equal to 0
## [1] "High.jump"
##
## Welch Two Sample t-test
##
## data: decathlon2$High.jump by decathlon2$Competition
## t = -1.0786, df = 24.99, p-value = 0.291
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.11477791 0.03587681
## sample estimates:
## mean in group Decastar mean in group OlympicG
##
                 1.977692
                                        2.017143
##
##
## Wilcoxon rank sum test with continuity correction
## data: decathlon2$High.jump by decathlon2$Competition
## W = 70, p-value = 0.3187
## alternative hypothesis: true location shift is not equal to 0
##
## [1] "X400m"
##
## Welch Two Sample t-test
##
## data: decathlon2$X400m by decathlon2$Competition
```

```
## t = 1.712, df = 24.999, p-value = 0.09927
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.1258454 1.3658454
## sample estimates:
## mean in group Decastar mean in group OlympicG
                    49.63
##
##
##
  Wilcoxon rank sum test with continuity correction
## data: decathlon2$X400m by decathlon2$Competition
## W = 123, p-value = 0.1264
## alternative hypothesis: true location shift is not equal to 0
## [1] "X110m.hurdle"
##
## Welch Two Sample t-test
##
## data: decathlon2$X110m.hurdle by decathlon2$Competition
## t = 2.5613, df = 18.548, p-value = 0.01933
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 0.07762028 0.77787423
## sample estimates:
## mean in group Decastar mean in group OlympicG
##
                 14.71846
                                        14.29071
##
##
## Wilcoxon rank sum test with continuity correction
## data: decathlon2$X110m.hurdle by decathlon2$Competition
## W = 137, p-value = 0.02715
## alternative hypothesis: true location shift is not equal to 0
## [1] "Discus"
##
## Welch Two Sample t-test
## data: decathlon2$Discus by decathlon2$Competition
## t = -0.9115, df = 23.954, p-value = 0.3711
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -3.969244 1.537486
## sample estimates:
## mean in group Decastar mean in group OlympicG
##
                 44.21769
                                        45.43357
##
## Wilcoxon rank sum test with continuity correction
##
## data: decathlon2$Discus by decathlon2$Competition
## W = 77.5, p-value = 0.5281
## alternative hypothesis: true location shift is not equal to 0
```

```
## [1] "Pole.vault"
##
## Welch Two Sample t-test
## data: decathlon2$Pole.vault by decathlon2$Competition
## t = -0.14888, df = 24.971, p-value = 0.8828
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.2249650 0.1946353
## sample estimates:
## mean in group Decastar mean in group OlympicG
                 4.827692
                                        4.842857
##
##
## Wilcoxon rank sum test with continuity correction
##
## data: decathlon2$Pole.vault by decathlon2$Competition
## W = 93, p-value = 0.9417
## alternative hypothesis: true location shift is not equal to 0
##
## [1] "Javeline"
##
## Welch Two Sample t-test
##
## data: decathlon2$Javeline by decathlon2$Competition
## t = -1.3377, df = 24.011, p-value = 0.1935
\#\# alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -6.684946 1.427034
## sample estimates:
## mean in group Decastar mean in group OlympicG
                                        59.58357
##
                 56.95462
##
## Wilcoxon rank sum test with continuity correction
## data: decathlon2$Javeline by decathlon2$Competition
## W = 67, p-value = 0.2541
## alternative hypothesis: true location shift is not equal to 0
## [1] "X1500m"
## Welch Two Sample t-test
## data: decathlon2$X1500m by decathlon2$Competition
## t = 1.8538, df = 18.178, p-value = 0.08007
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.9406291 15.1363434
## sample estimates:
## mean in group Decastar mean in group OlympicG
##
                 282.2000
                                        275.1021
##
```

```
##
##
   Wilcoxon rank sum test with continuity correction
##
## data: decathlon2$X1500m by decathlon2$Competition
## W = 128.5, p-value = 0.07245
## alternative hypothesis: true location shift is not equal to 0
## [1] "Points"
##
##
   Welch Two Sample t-test
##
## data: decathlon2$Points by decathlon2$Competition
## t = -3.8848, df = 24.767, p-value = 0.0006743
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -628.7878 -192.9375
## sample estimates:
## mean in group Decastar mean in group OlympicG
                 7905.923
                                        8316.786
##
##
##
##
   Wilcoxon rank sum test with continuity correction
##
## data: decathlon2$Points by decathlon2$Competition
## W = 19.5, p-value = 0.0005692
## alternative hypothesis: true location shift is not equal to 0
```

Power computations

Motivation

After looking at the results of the group comparison using the decathlon2 dataset the searcher wants to build another study in order to prove that the impact of the material used to build the surface of the stadium can impact the performance of the racing events and the long jump event. While building the protocol of his study the searcher encounters the question: How many sportsmen should he recruit in order to obtain satisfactory testing results?

```
library(factoextra)
data(decathlon2)
```

Working example

To answer this question we need to introduce an important notion in statistical modeling. The power of a statistical test. Let's simulate two variable for 2 different group. To do so we will have a look at two tests:

```
set.seed(352)
#Creation dataset
variableX.groupA <- rnorm(10, mean = 0, sd =1)
variableX.groupB <- rnorm(10, mean = 2, sd =1)
variableY.groupA <- rnorm(10, mean = 0, sd =1)
variableY.groupB <- rnorm(10, mean = 0.3, sd =1)
dataset <- data.frame(VariableX = c(variableX.groupA, variableX.groupB), VariableY = c(variableY.groupA)</pre>
```

We created both Variables to have a different means depending on the group. Now let's test these differences using the Student's t test.

```
t.test(VariableX ~ group, data = dataset)
##
   Welch Two Sample t-test
##
## data: VariableX by group
## t = -2.9165, df = 15.489, p-value = 0.01036
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -2.8904084 -0.4533992
## sample estimates:
## mean in group A mean in group B
        -0.0255522
                         1.6463516
t.test(VariableY ~ group, data = dataset)
##
   Welch Two Sample t-test
##
## data: VariableY by group
## t = -0.55427, df = 17.078, p-value = 0.5866
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -1.220411 0.712452
## sample estimates:
## mean in group A mean in group B
##
        0.07988438
                        0.33386405
```

We can see that even if we simulated both variables from different distributions depending on the group. We ended up with one test detecting significant differences between the groups and the other one not detecting differences. Typically, there is 4 different possible outcomes from a statistical test::

Reality Test results	Non-significant	Significant
H_0 True H_0 False	$1 - \beta$ (True Negative) β (False Negative)	$\begin{array}{c} \alpha \text{ (False Positive)} \\ 1 - \alpha \text{ (True Positive)} \end{array}$

The α represents they type I error, i.e. the probability of finding a significant difference when there is none. This coefficient α is often fixed to 5%. The β coefficient represent the probability to obtain a non-significant test while the alternative hypothesis is true and is called type II error. This is the case when we test the VariableY that we simulated. The power of a statistical test correspond to the values 1- β , and is the probability to reject the null hypothesis appropriately.

The power of a statistical test depends on 4 different parameters, the size of the effect (in the example the effect size is small), the sample size, the variance of the sample, and the significance level of the test α .

If the effect size of the phenomenon we want to study cannot be easily changed, we can influence the three others to obtain a good power:

- Increasing the sample size
- Decreasing the variability of the samples
- Increasing the level alpha, i.e. increasing the chance of false positives

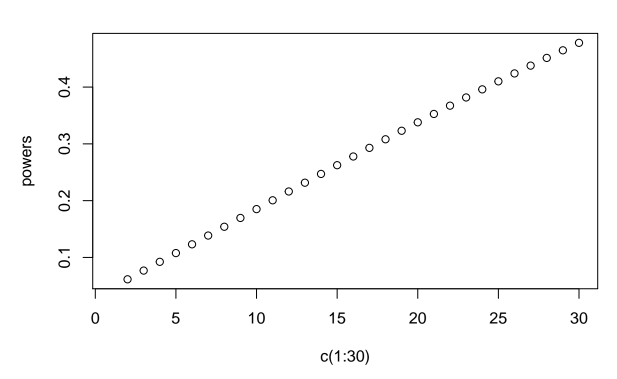
As the latter is not advisable in most studies, the easiest approach to influence the power of the test is to

increase the sample size. Let's compute the power to reject the null hypothesis in our example. To do so we can for example use the function pwr.t.test or power from the R package pwr:

```
library(pwr)
#Power computation VariableX
pwr.t.test(n = 10, d = 2/1, sig.level = 0.05, power = NULL, type = c("two.sample"),
             alternative = c("two.sided"))
##
##
        Two-sample t test power calculation
##
##
                 n = 10
##
                 d = 2
##
         sig.level = 0.05
##
             power = 0.988179
##
       alternative = two.sided
##
## NOTE: n is number in *each* group
#Power computation VariableY
pwr.t.test(n = 10, d = 0.5/1, sig.level = 0.05, power = NULL, type = c("two.sample"),
             alternative = c("two.sided"))
##
##
        Two-sample t test power calculation
##
##
                 n = 10
##
                 d = 0.5
         sig.level = 0.05
##
##
             power = 0.1850957
       alternative = two.sided
##
##
## NOTE: n is number in *each* group
```

We can see that the power to reject the null hypothesis for VariableX is 98.8% but for VariableY it is only 18.5%. As you can see the relation between the power and the effect size is not linear as the effect size is 4 times larger (0.5 to 2) but the power to reject H_0 is more than 5 times larger for VariableX. For simplicity we will not present, here, the exact formula to compute the power as it involves more statistical knowledge than the aim of this course.

We can plot the evolution of the power as a function of the effect size:



We see that even with a sample size of 30 the power to detect the difference between the two groups is only 0.4778965. Instead we can compute what would be the sample size needed for a given a power. The common desired power requested in studies are 80% and 90%. We can compute the sample size needed for both power for the VariableY

```
#computation of sample size for a fixed power of 80%
test.80.percent <- pwr.t.test(n = NULL, d = 0.5/1, sig.level = 0.05, power = 0.8, type = c("two.sample"
             alternative = c("two.sided"))
test.80.percent
##
        Two-sample t test power calculation
##
##
##
                 n = 63.76561
##
                 d = 0.5
         sig.level = 0.05
##
##
             power = 0.8
##
       alternative = two.sided
##
## NOTE: n is number in *each* group
#computation of sample size for a fixed power Of 90%
test.90.percent <- pwr.t.test(n = NULL, d = 0.5/1, sig.level = 0.05, power = 0.9, type = c("two.sample"
             alternative = c("two.sided"))
test.90.percent
##
##
        Two-sample t test power calculation
##
##
                 n = 85.03128
##
                 d = 0.5
##
         sig.level = 0.05
```

##

##

power = 0.9

alternative = two.sided

```
##
## NOTE: n is number in *each* group
```

We can see that we would need 64 samples in each group for a power of 80% and 85 for a power of 90%.

We have seen that in order to determine the proper sample size needed in a study we need to know or to give an estimate of the effect sizes we want to test, the standard deviation existing in the sample and to fix both type I error and the power to detect a true positive. To do so the approach often used is to go back to the literature or to used pilot studies.

Note that for the Student's t test obtaining power and sample sizes is relatively simple due to the exact mathematical formula linking all 4 parameters. But for a lot of test (especially non parametric ones) there are no exact formulas and simulations are needed. Hopefully, there are a lot of function in R that are able to do the simulation and power computations.

Quick Tasks

##

##

##

##

sig.level = 0.05

power = 0.9

alternative = two.sided

Compute the mean difference between the performances in long jump at the OlympicGames and Decastar. Compute the pooled standard deviation of the long jump event. What is the probability to detect a significant difference with a Student's t test in long jump performances within the decathlon2 study? (as we have different sample sizes per group use the function pwr.t2n.test)

```
mean.Decastar <- mean(decathlon2$Long.jump[decathlon2$Competition == 'Decastar'])
mean.OlympicGames <- mean(decathlon2$Long.jump[decathlon2$Competition == 'OlympicG'])
mean.diff <- mean.Decastar - mean.OlympicGames
standard.deviation <- sd(decathlon2$Long.jump)
samplesize.Decastar <- sum(decathlon2$Competition == 'Decastar')
samplesize.OlympicGames <- sum(decathlon2$Competition == 'OlympicG')
power.analysis <- pwr.t2n.test(n1 = samplesize.Decastar, n2 = samplesize.OlympicGames, d = mean.diff/st</pre>
```

pwr.t.test(n = NULL, d = mean.diff/standard.deviation, sig.level = 0.05, power = 0.8, alternative = c("

Compute what would be the needed sample size to obtain a power of 80%? and a power of 90%? (we will consider two group of the same size so here use pwr.t.test)

```
##
##
        Two-sample t test power calculation
##
                 n = 27.65562
##
##
                 d = 0.7673174
##
         sig.level = 0.05
##
             power = 0.8
##
       alternative = two.sided
##
## NOTE: n is number in *each* group
pwr.t.test(n = NULL, d = mean.diff/standard.deviation, sig.level = 0.05, power = 0.9, alternative = c("
##
##
        Two-sample t test power calculation
##
##
                 n = 36.67993
##
                 d = 0.7673174
```

```
## NOTE: n is number in *each* group
```

Finally devise the pooled standard deviation by 2. What would be then the needed sample sizes for a power of 80% and 90%?

```
standard.deviation <- standard.deviation/2
pwr.t.test(n = NULL, d = mean.diff/standard.deviation, sig.level = 0.05, power = 0.8, alternative = c("
##
##
        Two-sample t test power calculation
##
##
                 n = 7.754251
##
                 d = 1.534635
##
         sig.level = 0.05
##
             power = 0.8
##
       alternative = two.sided
##
## NOTE: n is number in *each* group
pwr.t.test(n = NULL, d = mean.diff/standard.deviation, sig.level = 0.05, power = 0.9, alternative = c("
##
##
        Two-sample t test power calculation
##
                 n = 9.989152
##
##
                 d = 1.534635
##
         sig.level = 0.05
             power = 0.9
##
##
       alternative = two.sided
##
## NOTE: n is number in *each* group
```

Exercises

Exercise 1

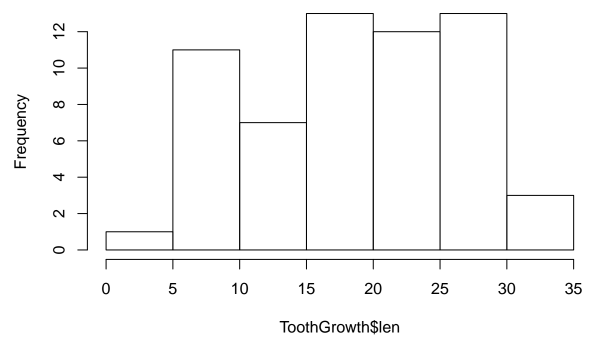
Load the ToothGrowth dataset using the command line data(ToothGrowth). This dataset is the result of an experiment on Guinea Pigs. It looks at the length of odontoblast cells after animals received a dose of vitamin C by one of two delivery methods. Namely, Orange juice (noted OJ) and ascorbic acid noted VC.

Source: Crampton, E. W. (1947). The growth of the odontoblast of the incisor teeth as a criterion of vitamin C intake of the guinea pig. The Journal of Nutrition, 33(5), 491-504. doi: 10.1093/jn/33.5.491.

1.1 Plot the histogram of the length of the odontoblast in the entire dataset.

```
hist(ToothGrowth$len)
```

Histogram of ToothGrowth\$len



- 1.2 Plot the boxplot of the length of the ordotonblast for each method of delivery of the Vitamin C. Compute the quartiles for both distributions.
- 1.3 Compute the mean absolute deviation in each group
- 1.4 Perform a Student't test to compare the mean of the length of the cells in both groups
- 1.5 Extract the value of the t statistics, the standard error and the degree of freedom from the test statistics. Display the results in the sentence:

The t statistics has a value of r, the estimated degrees of freedom of the Student's t distribution are r. The final p.value of the test is r.

- 1.6 Plot the distribution of the Student's t distribution under the null hypothesis
- 1.7 What are the values of the quantiles corresponding to boundaries between significane and non-significance?
- 1.8 what would be the number of Guinea Pigs per group required in order to have a power of 80% to detect a significant difference between both groups? And for a power of 90%?
- 1.9 Perform a Wilcoxon-Mann-Whitney test to compare the distribution of the length of the cells between both delivery methods

Exercise2

- 2.1 Create a plot that represent the power as a function of the sample for an experiment with an estimated difference in mean of 2, a standard deviation of 1.2, and a significance level of 0.05 of a two sided Student's t-test.
- 2.2 Create the same plot for a standard deviation of 2.4.
- 2.2 Create the same plot for a significance threshold of 0.1.

Regression models

Linear regression

Motivation

We have seen in the previous chapter how to compare the distribution between two groups. This gives us an understanding of the relationship between a continuous variable and a binary variable, i.e. a discrete variable with only two possible values. In order to test the association between a continuous variables we need to another approach. For example, does your ability in sprint can influence your pole vault or long jump performances? To answer this kind of question, the best approach is to use linear regression.

```
library(factoextra)

data(decathlon2)
```

Working Example

Let's use the simulated dataset we used in the previous chapter and add one variable VariableZ that we will build using the 2 other variables.

```
set.seed(352)
sample.size<-30

variableX.groupA <- rnorm(sample.size, mean = 2, sd = 1)
variableX.groupB <- rnorm(sample.size, mean = 2.4, sd = 1)
VariableX <- c(variableX.groupA, variableX.groupB)

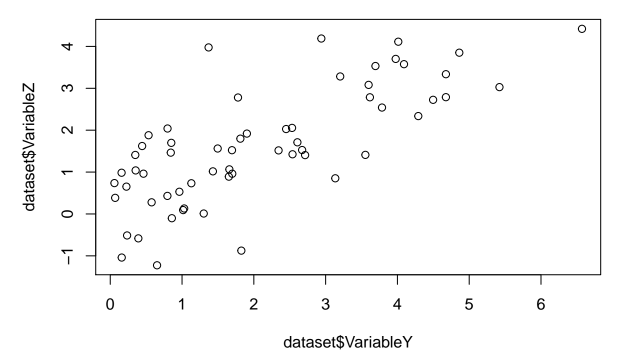
variableY.groupA <- rgamma(sample.size, shape = 2, scale = 1)
variableY.groupB <- rgamma(sample.size, shape = 2, scale = 1.1)
VariableY <- c(variableY.groupA, variableY.groupB)

VariableZ <- runif(1, min = 0, max = 1)*VariableX + runif(1, min = 0, max = 1)*VariableY + rnorm(2*sample group <- rep(c('A', 'B'), each = sample.size)

dataset <- data.frame(VariableX = VariableX, VariableY = VariableY, VariableZ = VariableZ, group = group
What is a linear regression analysis?</pre>
```

Let's make a scatterplot of VariableZ and VariableY.

```
plot(dataset$VariableY, dataset$VariableZ)
```



We can clearly see that as we look at higher values of the VariableY the values of VariableZ are also increasing. Quantifying the effect of a variable x on the variation of a variable of interest y is the aim of a linear regression model. A linear regression model is written as:

$$y_i = \alpha + \beta x_i + \epsilon_i$$

where y is the variable of interest also called outcome and x is the variable we are testing for association with the outcome. ϵ_i are called residuals and represent the part of the outcome that could not be 'explained' by x. Each ϵ_i are supposed independent and drawn from a normal distribution $N(0, \sigma^2)$. The aim of a logistic regression analysis is to estimate the value of α , β and σ^2 . To do so we use the method of least square, i.e., we find the value of α and β that minimizes the sum of squared residuals:

$$SS_{res} = \sum_{i} (y_i - (\alpha + \beta x_i))^2$$

This equation leads to exact formulas for the estimation of the parameters :

$$\hat{\beta} = \frac{\sum_{i} (x_i - \overline{x})(y_i - \overline{y})}{\sum_{i} (x_i - \overline{x})^2}$$

$$\hat{\alpha} = \overline{y} - \hat{\beta}\overline{x}$$

$$\sigma^2 = \frac{SS_{res}}{n-2}$$

Let's do a linear regression of VariableY on VariableZ by using the function lm in R:

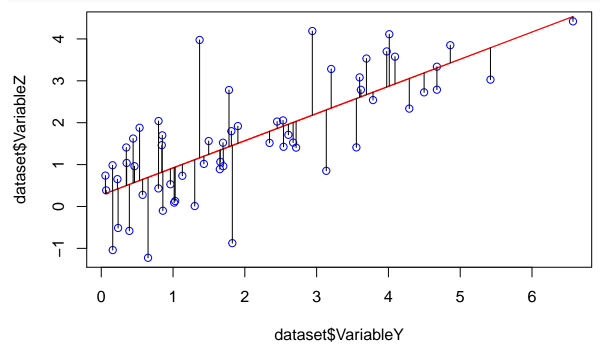
```
regression <- lm(VariableZ ~ VariableY, data = dataset)
regression</pre>
```

```
##
## Call:
## lm(formula = VariableZ ~ VariableY, data = dataset)
##
## Coefficients:
## (Intercept) VariableY
## 0.2723 0.6487
```

The function lm returns 2 values, the intercept which correspond to α and the β coefficient corresponding to the value below the VariableY. The 4beta\$ coefficient can be interpreted as if my variableX increases of 1

my variable will increase of summary(regression) coefficients [2,4]. We can represent the results onto the scatterplot:

```
plot(dataset$VariableY, dataset$VariableZ, col = 'blue')
lines(dataset$VariableY,fitted(regression), col = 'red')
segments(dataset$VariableY,fitted(regression), dataset$VariableY,dataset$VariableZ)
```



In this plot, the blue points are the observed values, the red line is the result of the regression analysis with α being the intercept of the line and β its slope. Finally the black vertical line represent the residuals, i.e. the variation that could not be explained by the linear regression model.

The interest of a linear regression is that we can test if the coefficient β is equal to 0 in order to test for association between the 2 variables. To do so in a univariate linear regression we define the statistics t as:

$$t = \frac{\beta}{se(\beta)}$$

where t (under the assumption that the outcome is normally distributed and under the null hypothesis H_0 : $\beta = 0$) follows a Student's t distribution with n-2 degress of freedom. With this information computing the p-value of the test is the same as for the Sutdent's t test.

For example we can look at the association between VariableX and group using both Student's t test and the linear regression:

```
#Student's t test between VariableX and group
t.test(VariableX ~ group, data = dataset)

##
## Welch Two Sample t-test
##
## data: VariableX by group
## t = -2.293, df = 56.263, p-value = 0.02561
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -1.16872012 -0.07888065
## sample estimates:
## mean in group A mean in group B
```

```
## 1.900228 2.524028
#linear regression between VariableX and group
linear.regression <- lm (VariableX ~ group, data = dataset)
summary(linear.regression)$coefficients</pre>
```

```
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 1.9002279 0.1923664 9.878170 4.879130e-14
## groupB 0.6238004 0.2720472 2.292986 2.549308e-02
```

We see that we obtain exactly the same results. As both approaches are equivalent the sample size and power computation for a linear regression model are the same than for the student's t test.

Note that there is another possible statistical test to apply is the likelihood ratio test. This test comes from a statiscal test called goodness of fit test which allows to test for a significant improvement of the model fit. By comparing the likelihoods, i.e. the probability of observing the set of values based on the statistical model fitted, of the models fitted under the null hypothesis (in our case with only an intercept) and the alternative hypothesis (the model that we fitted). This test will be seen more in detail later in the course. Asymptotically (with the number of samples sufficiently large), both likelihood ratio test and student's t test are equivalent.

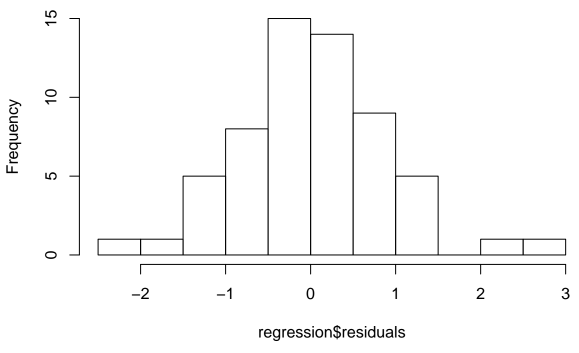
There are four assumptions associated with a linear regression model. First, the outcome has to be a normally ditributed continuous variable. The relationship between the variable x and \overline{y} is linear. The observations have to be independent and the variance of the residuals have to be independent of X (homoscedasticity). To trust the results of the linear regression several steps have to be taken based on these assumptions:

- Checking the distribution of the outcome if the distribution is too far from a gaussian distribution it can be good to apply a transformation to the data. Such as log transformation.
- Checking the distributions of the residuals as they are assumed to be normal.

In our case we simulated the outcome VariableZ as the sum of two gaussian distribution so VariableZ. Instead let's plot the distribution of the residuals:

```
hist(regression$residuals, breaks = 10)
```

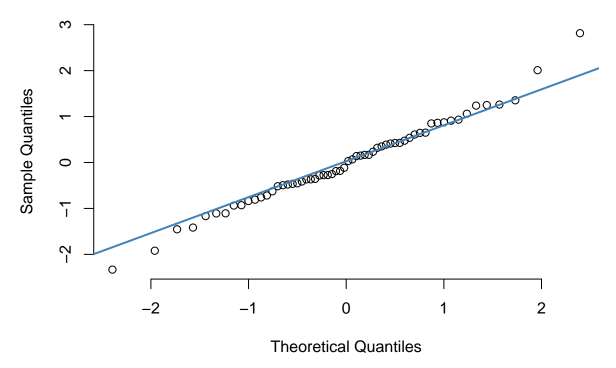
Histogram of regression\$residuals



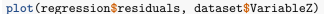
We can see clearly that the residuals distribution is really close from a normal distribution. This clearly indicates that the regression went well. Another plit is commonly used to check if a distribution is following a gaussian distribution. The Q-Q plot. This plot consist of comparing a set of values with the quantiles of a normal distribution:

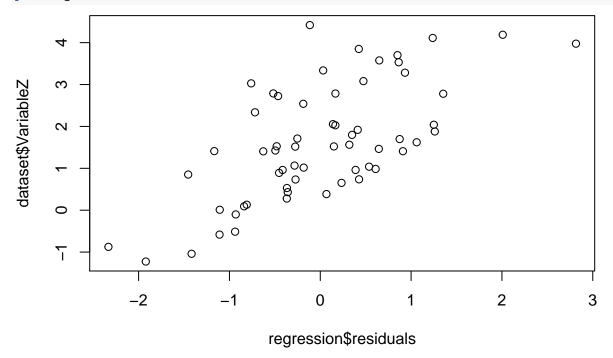
```
qqnorm(regression$residuals, pch = 1, frame = FALSE)
qqline(regression$residuals, col = "steelblue", lwd = 2)
```

Normal Q-Q Plot



The closer the points are from the blue line the closer the set of values is close from a normal distribution. Another interesting graph to plot is the scatter plot of the outcome and the residuals.





We can clearly see a trend in the scatterplot. This trend is interesting, as it means that the residuals of the linear models are associated with the outcome Y. Meaning that the variation not explained by the linear model is not due to randomess and another linear association, independent to VariableY exists. It is easy to

explain in our case as VariableZ was simulated as the weighted sum of VariableX and VariableY.

To obtain a better linear model it might of interest to use linear models with multiple variables such as:

```
z_i = \alpha + \beta_1 y_i + \beta_2 x_i + \epsilon_i
```

let's run the model using the 1m function:

```
multiple.regression <- lm (VariableZ ~ VariableX + VariableY, data=dataset)
multiple.regression</pre>
```

```
##
## Call:
## lm(formula = VariableZ ~ VariableX + VariableY, data = dataset)
##
## Coefficients:
## (Intercept) VariableX VariableY
## -0.2473     0.2658     0.6158
```

And we can check the significance of the β s obtained:

```
summary(multiple.regression)$coefficients
```

```
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.2473115 0.28451895 -0.8692268 3.883686e-01
## VariableX 0.2658086 0.10814688 2.4578481 1.704112e-02
## VariableY 0.6158445 0.07401264 8.3208018 2.020188e-11
```

We can indeed see that both coefficients are correctly identified as significant. We can also note that the coefficient of VariableY is almost not impacted by the addition of VariableX. This is due to the fact that these variables were simulated independently and their correlation should be close to 0. After, computation we obtain a correlation of 0.1804148. With a larger correlation between both variables, the impact on the coefficient obtained on the simple linear regression would have been stronger.

Strong correlation between predictors in the same models can create the so-called multi-collinearity problem that leads to a severe increase of the p.value and therefore a reduction in significance.

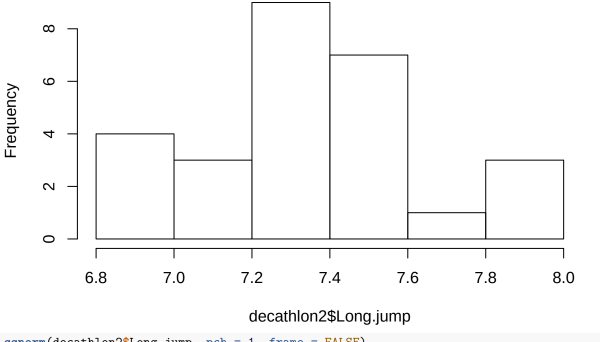
Note that the power and sample size computations for linear regression with multiple predictors becomes rather challenging. Searcher often use rules of thumbs. A commonly encountered rule of thumbs is at least 10-15 samples for each predictors in the model. But this is only a rule of thumbs and as we know, the required sample size is depending on the variation of the outcome of interest as well as the effect size to test.

Quick Task

Check if the variable Long. jump of the decathlon2 dataset follows a normal distribution using a plot.

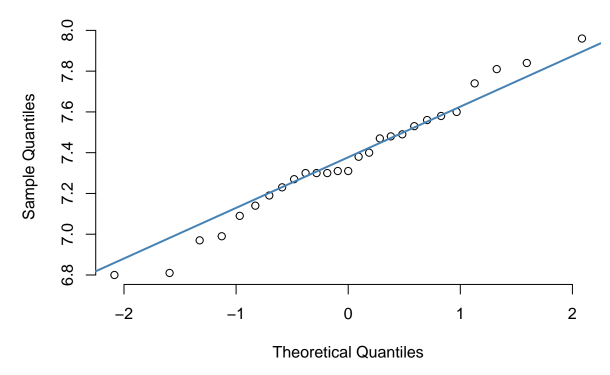
```
hist(decathlon2$Long.jump)
```

Histogram of decathlon2\$Long.jump



```
qqnorm(decathlon2$Long.jump, pch = 1, frame = FALSE)
qqline(decathlon2$Long.jump, col = "steelblue", lwd = 2)
```

Normal Q-Q Plot



Perform a linear regression the check for association between the variables Long.jump and X100m. What is the beta value? Is the resul significant?

```
linear.model <- lm(Long.jump ~ X100m, data = decathlon2)</pre>
linear.model
##
## Call:
## lm(formula = Long.jump ~ X100m, data = decathlon2)
##
## Coefficients:
##
   (Intercept)
                       X100m
       15.8405
                     -0.7711
summary(linear.model)$coefficients
##
                 Estimate Std. Error
                                         t value
                                                      Pr(>|t|)
## (Intercept) 15.8404506 1.5514078 10.210372 2.102907e-10
## X100m
               -0.7710575  0.1410923  -5.464917  1.124878e-05
We know that there is a significant difference in mean for the variable X100m between both competitions.
Add the predictor Competition into the model as well as an interaction effect between Competition and
X100m (add X100m:Competition in the formula). Are the results changing?
linear.model <- lm(Long.jump ~ X100m + Competition + X100m:Competition, data = decathlon2)
linear.model
##
## Call:
## lm(formula = Long.jump ~ X100m + Competition + X100m:Competition,
       data = decathlon2)
##
##
##
   Coefficients:
                                                    X100m
##
                  (Intercept)
##
                      14.1791
                                                   -0.6202
##
         CompetitionOlympicG X100m:CompetitionOlympicG
##
                       6.4655
                                                   -0.5968
summary(linear.model)$coefficients
##
                                Estimate Std. Error
                                                        t value
                                                                    Pr(>|t|)
## (Intercept)
                              14.1791059 2.5883364 5.478077 1.436274e-05
## X100m
                              -0.6202394   0.2315586   -2.678541   1.341569e-02
## CompetitionOlympicG
                               6.4655012 3.9932814 1.619095 1.190593e-01
## X100m:CompetitionOlympicG -0.5968057 0.3640715 -1.639254 1.147685e-01
```

Logistic regression

Motivation

In some studies the response variable only has two possible values. Some examples:

- individuals who have or have not cancer;
- cancer patients who, after treatment, relapse or not;
- from all patients receiving the same treatment, each develops resistance or not.

If analysing data with such a binary response variable, care must be taken, in particular when using regression models. Such a variable can be coded as $\{0,1\}$ in R, for example. However, linear regression is not going to

return predicted values as only 0 or 1. In fact, nothing prevents a linear regression of returning a negative value, or a value larger than 1.

Logistic regression

In order to avoid this and model the data correctly, we need to use a regression model with a *link* function. For a regression model with a response Y and covariates X_1, X_2 , this link function can be written as h, as in the model:

$$E(Y) = h^{-1} (\alpha + \beta_1 X_1 + \beta_2 X_2).$$

The job of the link function is to make sure that the error is modelled correctly, and it yields predicted values in the range desired. For a binary response, the ideal link function is the *logistic* function. It takes any values, positive or negative, and always return a value between 0 and 1. The logistic function has can be written as:

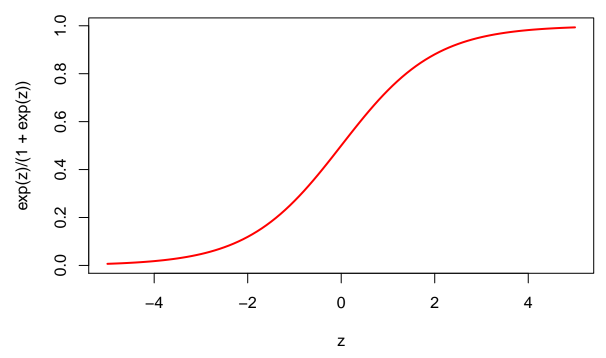
$$f(z) = \frac{e^z}{1 + e^z},$$

for any value of z. The graph of this function for values of z between -5 and 5 is:

```
z \leftarrow seq(from = -5, to = 5, by = 0.1)

plot(z, exp(z)/(1+exp(z)), main = "Logistic function", type = "l", col = "red", lwd = 2)
```

Logistic function



The logistic link function relates the response Y and the linear prediction (which is the function of the covariates, $\alpha + \beta_1 X_1 + \beta_2 X_2$) in the following way:

$$E(Y) = \frac{e^{\alpha + \beta_1 X_1 + \beta_2 X_2}}{1 + e^{\alpha + \beta_1 X_1 + \beta_2 X_2}}.$$

The function can be inverted to write:

$$\log(\frac{E(Y)}{1 - E(Y)}) = \alpha + \beta_1 X_1 + \beta_2 X_2.$$

The glm function

The logistic regression model is fitted by the function glm. This function works in very similar ways to lm for linear regression, such as for example using as main input a formula determining the model to be used. It also may use family and link slots to define which probability distribution is to be used for the errors (family) and which link function is to be used (link). Per family, one link function is used by default. In the case of the logistic model, it is defined family = binomial, and the default link is the logistic.

The response for the logistic model can be given in different ways. We will here use the pair (number dead, number alive) per combination of dose and sex. See the exercises for an example of a binary response.

As with a linear regression, summary and anova are used to obtain an overview of the model fit and test for effects of variables.

Working example

This example is adapted from an example used in Venables and Ripley (1995):

Venables, W. N. and Ripley, B. D. (1995). Modern applied statistics with S-Plus. Springer-Verlag: New York. https://www.springer.com/gp/book/9780387954578

Collet (11=991, p. 75) reports an experiment on the toxicity of the tobacco budworm *Heliothis virescens* to doses of the pyrethroid trans-cypermethrin to which the moths were beginning to show resistance. It is of interest to determine which dose level to choose, so as to guarantee a specific death proportion. The response variable is thus dead or alive per moth. Being a binary variable, a logistic regression is an ideal tool to analyse the data.

Batches of 20 months of each sex were exposed for 3 days to the pyrethroid and the number in each batch which were dead (or knocked down) was recorded. The doses are used in two-fold increases, so it is natural to consider those on the log2-scale. The table of total dead moths per dose and sex is:

sex	d=1	d=2	d=4	d=8	d=16	d=32
male	1	4 2	9	13	18	20
female	0		6	10	12	16

We enter the data into R as follows: the dose is entered in its original scale, then log2-transformed. The total numbers of death moths are entered as a single vector, with all males followed by all females. So, the dose vector needs to be repeated (stacked) to produce entries corresponding to all total death count observations. Finally, we enter the sex by repeating male or female the required number of times.

```
dose <- c(1, 2, 4, 8, 16, 32)
ldose <- log2(dose)
numdead <- c(1, 4, 9, 13, 18, 20, 0, 2, 6, 10, 12, 16)
ldose <- rep(ldose, 2)
sex <- rep(c("male", "female"), each = length(dose))</pre>
```

To define the response as the pair of (number dead, number alive) we do:

```
resp <- cbind(numdead, numalive = 20 - numdead)</pre>
```

Now we fit the logistic model:

```
budworm.lg <- glm(resp ~ sex + ldose, family = binomial)</pre>
summary(budworm.lg)
##
## Call:
## glm(formula = resp ~ sex + ldose, family = binomial)
##
## Deviance Residuals:
##
                                        3Q
        Min
                   1Q
                         Median
                                                 Max
## -1.10540 -0.65343 -0.02225
                                   0.48471
                                             1.42944
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) -3.4732
                            0.4685 -7.413 1.23e-13 ***
                 1.1007
                            0.3558
                                      3.093 0.00198 **
## sexmale
## ldose
                 1.0642
                             0.1311
                                      8.119 4.70e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 124.8756 on 11
                                       degrees of freedom
## Residual deviance:
                                        degrees of freedom
                        6.7571
                                on
                                    9
## AIC: 42.867
##
## Number of Fisher Scoring iterations: 4
There may also be an interaction effect between sex and ldose, meaning that the ldose effect may change
depending on the sex. To check this, we add the interaction term:
budworm.lg.i <- update(budworm.lg, . ~ sex:ldose)</pre>
summary(budworm.lg.i)
##
## Call:
## glm(formula = resp ~ sex:ldose, family = binomial)
##
## Deviance Residuals:
##
        Min
                   1Q
                         Median
                                        3Q
                                                 Max
## -1.45854 -0.29503 -0.05339
                                  0.44888
                                             1.06990
##
## Coefficients:
##
                   Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                    -2.9073
                                0.3893
                                        -7.468 8.12e-14 ***
                                          6.920 4.52e-12 ***
## sexfemale:ldose
                     0.8823
                                 0.1275
## sexmale:ldose
                     1.2893
                                 0.1669
                                          7.723 1.13e-14 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 124.8756
                                        degrees of freedom
                                on 11
## Residual deviance:
                        5.0443 on 9 degrees of freedom
## AIC: 41.155
##
```

```
## Number of Fisher Scoring iterations: 4
```

Here we fitted a new model by using the function update, which added the interaction effect sex:dose to the linear predictor already in budworm.lg. Note that this is the same as fitting the model by calling glm again with the interaction term added.

We can use anova to summarize results for a model, as before. Here we will test for the effect of variables in the model fit budworm.lg. The function anova will perform testing as if variables were added sequentially, one by one. In this case, the effect of sex is tested, subsequently the effect of ldose given that the sex effect is corrected for. Let us try this out:

```
anova(budworm.lg)
```

```
## Analysis of Deviance Table
##
## Model: binomial, link: logit
##
## Response: resp
##
  Terms added sequentially (first to last)
##
##
##
##
         Df Deviance Resid. Df Resid. Dev
## NULL
                             11
                                    124.876
                6.077
                             10
                                    118.799
## sex
          1
## ldose
          1
             112.042
                                      6.757
```

Note that the anova function now did not return a p-value for the covariate effects. When applied with generalized linear model fits, this is the default. This forces the user to choose the appropriate test to run (see help(anova.glm) for details). The suitable test in this case is the chi-square, so we rerun it using:

```
anova(budworm.lg, test = "Chisq")
```

```
## Analysis of Deviance Table
##
## Model: binomial, link: logit
##
## Response: resp
##
## Terms added sequentially (first to last)
##
##
##
         Df Deviance Resid. Df Resid. Dev Pr(>Chi)
## NULL
                            11
                                  124.876
## sex
               6.077
                            10
                                  118.799
                                             0.0137 *
## ldose
             112.042
                             9
                                     6.757
                                             <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

In order to test for the main effects, followed by the interaction, use the entire model fit in one glm call:

```
anova(glm(resp ~ sex + ldose + sex:ldose, family = binomial), test = "Chisq")
```

```
## Analysis of Deviance Table
##
## Model: binomial, link: logit
##
## Response: resp
```

```
##
## Terms added sequentially (first to last)
##
##
##
             Df Deviance Resid. Df Resid. Dev Pr(>Chi)
## NULL
                                  11
                                        124.876
                    6.077
## sex
                                  10
                                        118.799
                                                   0.0137 *
## ldose
              1
                  112.042
                                   9
                                          6.757
                                                   <2e-16 ***
## sex:ldose
              1
                    1.763
                                   8
                                          4.994
                                                   0.1842
## ---
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```

We conclude that both sex and ldose have statistically significant effects, but that the interaction does not.

Quick tasks

The esoph data includes counts of a case-control study of esophageal cancer, according to age (agegp), alcohol consumption (alcgp) and tobacco consumption (tobgp). Fit a logistic regression model to the responses, formed by both case and control counts, using all covariates available. Print a summary of the model fit, and also the ANOVA table.

Fit also a model which includes also an interaction effect between alcohol and tobacco consumption. Print again the model summary and the anova table of the model fit.

Fitted values

It is useful to make a graph of the predicted values. For this, we use the function predict. It needs as input the fitted model, in this case budworm.lg, as well as the data for which you want to make predictions. If no data is given, the fitted values corresponding to the observed data are used. If data is provided, it needs to be given as a data.frame object, containing variables with the same variable names as in the original data.

Let us first extract the fitted values:

```
myfitted <- predict(budworm.lg)
myfitted</pre>
```

```
##
                        2
                                    3
                                                4
                                                            5
                                                                        6
                                                                                    7
             1
## -2.3724119 -1.3081980 -0.2439840
                                        0.8202300
                                                   1.8844439
                                                               2.9486579 -3.4731553
##
            8
                        9
                                   10
                                                           12
                                               11
## -2.4089413 -1.3447274 -0.2805134
                                       0.7837006
```

By default, predict returns values on the scale of the linear predictor, i.e. it returns

$$\hat{\alpha} + \hat{\beta}_1 X_1 + \hat{\beta}_2 X_2,$$

where $\hat{\alpha}, \hat{\beta}_1, \hat{\beta}_2$ are the parameter estimates from the model fit, as given in the column Estimate of the model fit summary:

```
summary(budworm.lg)$coef
```

```
## Estimate Std. Error z value Pr(>|z|)
## (Intercept) -3.473155 0.4685202 -7.413032 1.234445e-13
## sexmale 1.100743 0.3558271 3.093478 1.978249e-03
## ldose 1.064214 0.1310775 8.118971 4.701542e-16
```

While this is useful, it is easier to examine the model fit by comparing the fitted values on the scale of the response. For this, we use

```
myfitted <- predict(budworm.lg, type = "response")
myfitted</pre>
```

```
## 1 2 3 4 5 6 7
## 0.08530076 0.21278854 0.43930479 0.69428515 0.86812073 0.95020002 0.03008577
## 8 9 10 11 12
## 0.08249341 0.20673372 0.43032791 0.68647712 0.86388206
```

These are fitted probabilities of dead moths for each combination of ldose and sex.

Quick tasks

- 1. Continue examining the example of budworms. Fit now the model including the interaction between ldose and sex. Print the ANOVA table with a test for each coefficient. Extract the fitted probabilities on the scale of the response.
- 2. For the **esoph** data analysis, extract the fitted values for the model without and with the interation. Put those together in a matrix.

Exercises

Exercise 1

- 1.1 We go back to the ToothGrowth dataset. We saw that there is a significant difference in the average length of the odontoblast between the two delivery methods of vitamin C. We are now interested in the relationshipt between the dose of vitamin C and the length of odontoblast. Perform a linear regression model to test for association between these 2 variables.
- 1.2 Extract the estimates and corresponding statistics and p-values frol the regression
- 1.3 Create Plots in order to check the assumptions of the model
- 1.3 We can see that the residuals does not really follow a gaussian distribution. Use a log transformation on the length variable and compare the results
- 1.4 The transformation slightly improved the results. From the previous exercises we have observed an association between the delivery method and the length of the cells. Ass the delivery method in the linear regression:
- 1.5 It could be relevant to add a possible interaction between the dose given and the delivery method. Add an interaction effect in the model:
- 1.6 What do you conclude from this analysis?

Exercise 2

2.1. Return to the budworms example. Using the model fitted including the main effects of sex and ldose as well as their interaction, extract the fitted probabilities. Make a graph of observed and expected probabilities, and include a legend. Make sure to distinguish the groups, for example by using different plotting symbols for sex and different colours depending on the dose. Note that the observed probabilities are given by numdead/20.

Exercise 3

Analyse now the birthwt data from the package MASS, to investigate factors that may affect the chance of low birth weight.

- 3.1 Fit a logistic regression model to the variable representing low weight, using smoke as covariate in the model. Note that the response variable now is binary, not counts. Examine the model fit to check if it represents a good approximation of the data.
- 3.2 Extract fitted probabilities and plot those together with the observed data. Note that the latter is now simply a binary variable.
- 3.3 Fit now a logistic regression model to the variable representing low weight, using all covariates in the model. Examine the model fit to check if it represents a good approximation of the data. Extract fitted probabilities and plot those together with the observed data. Note that the latter is now simply a binary variable.
- 3.4 We think that the model can be improved further, but it is not immediately clear which variables should be included, and in which order. For this, we can use the function step, which performs stepwise regression. This involves comparing model fits by using the Akaike Information Criterion (AIC). A statistical test helps deciding whether the AIC value indicates a statistically significant improvement between model fits. Try to improve the basic model which includes smoke and lwt, by adding variables age, ptl, ht and ftv one at a time. Check the help file for step, and call the function using as object the fitted model with smoke and lwt, and as scope the formula for the model including these two, plus the extra covariates.

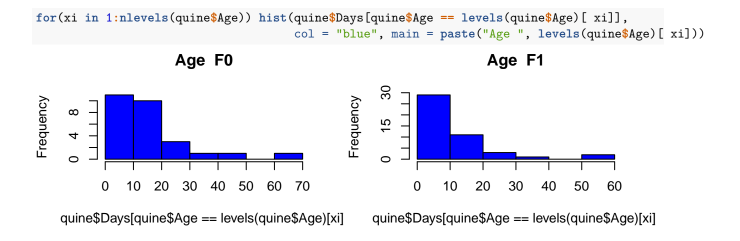
More tests and count data

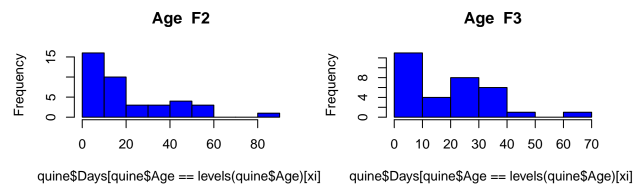
Comparing more than 2 groups

Motivation

Consider the quine data on absenteeism from school in an Australian region. Per child involved in the study, the data includes the number of days absent from school in that year, as well as the age group (in 4 categories). One question is: does the number of absent days change depending on the school year?

```
library(MASS)
data(quine)
str(quine)
                    146 obs. of 5 variables:
## 'data.frame':
## $ Eth : Factor w/ 2 levels "A", "N": 1 1 1 1 1 1 1 1 1 1 ...
## $ Sex : Factor w/ 2 levels "F", "M": 2 2 2 2 2 2 2 2 2 2 ...
## $ Age : Factor w/ 4 levels "F0", "F1", "F2",...: 1 1 1 1 1 1 1 1 2 2 ...
## $ Lrn : Factor w/ 2 levels "AL", "SL": 2 2 2 1 1 1 1 1 2 2 ...
## $ Days: int 2 11 14 5 5 13 20 22 6 6 ...
# Examine how many observations you have per age group
table(quine$Age)
## F0 F1 F2 F3
## 27 46 40 33
# Now display the data for `Days` per age group
par(mfrow = c(2, 2))
```





You could try to solve this by applying a Student's-t test, or a Wilcoxon test, to compare each pair of Age levels. This would involve 6 tests.

In such situations, it would be better to compare the means of the groups under study in one go. What we wish to know is: is there at least one age group that does not have the same mean number of absent days as the others?

To do this, we need to better understand what we understand by "having the same mean".

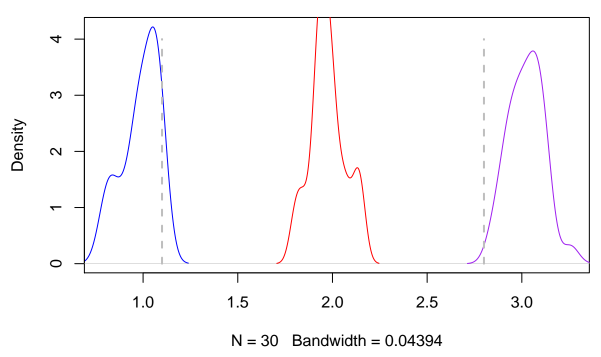
Working examples

Consider the following data: a continuous variable is observed for cases within groups X, Y, Z. The variable follows a normal distribution with a mean depending on the group: 1, 2 and 3 for groups X, Y, Z respectively. Say that we have 100 observations for each group, and that the empirical densities of the values per group aree:

```
set.seed(53412)
ssize <- 30
x <- rnorm(ssize, mean = 1, sd = 0.1)
y <- rnorm(ssize, mean = 2, sd = 0.1)
z <- rnorm(ssize, mean = 3, sd = 0.1)
mylims <- range(c(x, y, z))
plot(density(x), xlim = mylims, col = "blue")
lines(density(y), col = "red")
lines(density(z), col = "purple")
segments(1.1, 0, 1.1, 4, lty = "dashed", lwd = 2, col = "gray")</pre>
```

```
segments(2.8, 0, 2.8, 4, lty = "dashed", lwd = 2, col = "gray")
```

density.default(x = x)



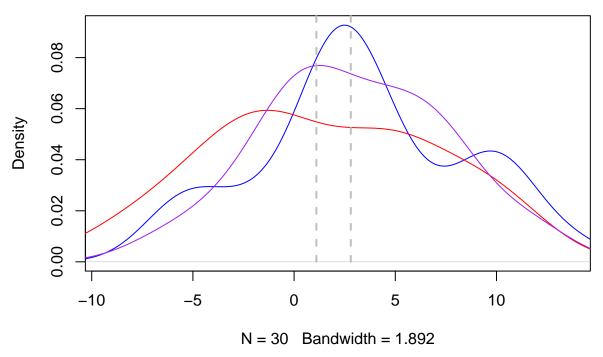
```
data1 <- data.frame(x, y, z)
```

We can clearly separate observations arising from these 3 groups. Indeed, for example if we knew that a value of 1.1 was observed for one of these variables, we would guess that it corresponded to a case in group X. Similarly, if we knew that a value of 2.8 had been observed, we would safely guess it to correspond to a case in group Z.

Now let us simulate the data again for the same setup:

```
set.seed(36912)
x <- rnorm(ssize, mean = 1, sd = 5)
y <- rnorm(ssize, mean = 2, sd = 5)
z <- rnorm(ssize, mean = 3, sd = 5)
mylims <- range(c(x, y, z))
plot(density(x), xlim = mylims, col = "blue")
lines(density(y), col = "red")
lines(density(z), col = "purple")
segments(1.1, 0, 1.1, 4, lty = "dashed", lwd = 2, col = "gray")
segments(2.8, 0, 2.8, 4, lty = "dashed", lwd = 2, col = "gray")</pre>
```

density.default(x = x)



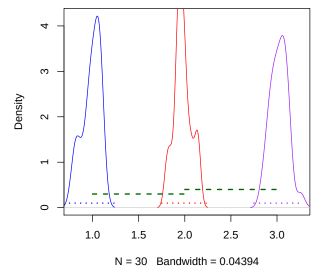
```
data2 <- data.frame(x, y, z)</pre>
```

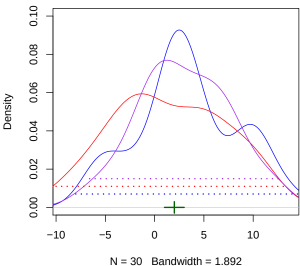
In this case, it is difficult to distinguish data from these 3 groups. Indeed, we could now not guess to which group a value of 1.1 would correspond: it can be easily generated by cases in either X or Y, and even by some in Z. The same is true for 2.8. Note that the group means are the same as before: 1, 2 and 3. However, now the variance within each group is much larger than for the first data.

What this means is that: observations from these 3 groups are different from each other if they can be distinguished from each other, and that can only be done if the variance *between* groups is larger than the variance *within* groups.

```
par(mfrow = c(1, 2))
mydata <- data1
mylims <- range(c(mydata$x, mydata$y, mydata$z))</pre>
plot(density(mydata$x), xlim = mylims, col = "blue", main = "")
lines(density(mydata$y), col = "red")
lines(density(mydata$z), col = "purple")
segments(0.75, 0.1, 1.28, 0.1, lty = "dotted", lwd = 2, col = "blue")
segments(1.75, 0.1, 2.28, 0.1, lty = "dotted", lwd = 2, col = "red")
segments(2.75, 0.1, 3.28, 0.1, lty = "dotted", lwd = 2, col = "purple")
segments(1, 0.3, 2, 0.3, lty = "dashed", lwd = 2, col = "darkgreen")
segments(2, 0.4, 3, 0.4, lty = "dashed", lwd = 2, col = "darkgreen")
mydata <- data2
mylims <- range(c(mydata$x, mydata$y, mydata$z))</pre>
plot(density(mydata$x), xlim = mylims, col = "blue", main = "", ylim = c(0, 0.1))
lines(density(mydata$y), col = "red")
lines(density(mydata$z), col = "purple")
segments(-8, 0.007, 15, 0.007, lty = "dotted", lwd = 2, col = "blue")
segments(-10, 0.011, 13.5, 0.011, lty = "dotted", lwd = 2, col = "red")
```

```
segments(-6.6, 0.015, 12.8, 0.015, lty = "dotted", lwd = 2, col = "purple")
segments(1, 0.00, 3, 0.0, lty = "solid", lwd = 2, col = "darkgreen")
segments(2, -0.003, 2, 0.003, lwd = 2, col = "darkgreen")
```





Thus, in order to know if observations corresponding to the different groups can be distinguished or not, we need to consider how large the variance between groups (in green) is, relative to the variances within groups (in their respective colours). This is what the F test does, via the statistic F = variance between groups / variance within groups. If the variance between groups is large compared to the variance within groups, as with the first dataset, observations can be distinguished between groups. Otherwise, observations from different groups overlap, as in the second dataset. The F test is applied assuming that the groups cannot be distinguished, meaning that the variances within and between groups are similar. When observations from the different groups cannot be distinguished, the value of the F-test statistic is around 1. When observations can be distinguished between groups, the value of the F-test statistic will be relatively large, in particular larger than 1. How large it needs to be depends on the number of samples and the number of groups being compared, determining the degrees of freedom used in the F-test statistic.

The F test is the one used in ANOVA (which stands for ANalysis Of VAriance - now you know why!).

Let us apply this to the first dataset we generated. To do this, we need to fit a linear regression to the data with the observed variable as response and the group as explanatory factor. Then we compute the ANOVA table of the resulting fit:

```
group <- factor(rep(c("X", "Y", "Z"), each = ssize))
mydata <- data1
myvar <- c(mydata$x, mydata$y, mydata$z)
myfit <- lm(myvar ~ group)
anova(myfit)</pre>
```

Examine the ANOVA table. The column "Mean Sq" actually displays the estimated variances. The first row corresponds to group so it is the variance between groups X, Y, Z. The second row corresponds to

Residuals, and this is the variance within groups - or the residual variance not explained by group. By dividing the variance between groups by the one within, the F-test statistic is obtained (note that the values printed are rounded off). As we expected, the F-test statistic is very large and its corresponding p-value is very small, in this case.

Now for the second dataset:

```
mydata <- data2
myvar <- c(mydata$x, mydata$y, mydata$z)
myfit <- lm(myvar ~ group)
anova(myfit)

## Analysis of Variance Table
##
## Response: myvar
## Df Sum Sq Mean Sq F value Pr(>F)
## group 2 61.75 30.875 1.176 0.3134
## Residuals 87 2284.17 26.255
```

The F test within the ANOVA table relies on the data arising from a normal distribution. If that is not the case, the alternative is to use the Kruskal-Wallis test, which can be thought of as the nonparametric version of the F test - much in the same way as the Wilcoxon test is the nonparametric version of the Student's-t test. Two alternative syntaxes for the Kruskal-Wallis test are:

```
kt <- kruskal.test(myvar, group)
kt

##
## Kruskal-Wallis rank sum test
##
## data: myvar and group
## Kruskal-Wallis chi-squared = 2.0227, df = 2, p-value = 0.3637
kt2 <- kruskal.test(myvar ~ group)
kt2

##
## Kruskal-Wallis rank sum test
##
## Kruskal-Wallis rank sum test
##
## Kruskal-Wallis chi-squared = 2.0227, df = 2, p-value = 0.3637</pre>
```

The Kruskal-Wallis test is based on ranks of the data. In this case, it is more powerful to find differences between groups than the F test from ANOVA.

Quick tasks

Let us now consider the quine data again. Use both the F test as well as the Kruskal-Wallis test to decide if the number of days of absence differs between age groups.

```
myfit <- lm(Days ~ Age, data = quine)
anova(myfit)

## Analysis of Variance Table
##
## Response: Days
## Df Sum Sq Mean Sq F value Pr(>F)
## Age 3 2535 845.04 3.3547 0.02074 *
```

```
## Residuals 142 35769 251.90
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
kruskal.test(Days ~ Age, data = quine)

##
## Kruskal-Wallis rank sum test
##
## data: Days by Age
## Kruskal-Wallis chi-squared = 7.6347, df = 3, p-value = 0.0542
```

Testing independence in 2x2 tables

Motivation

Consider again the quine data on absenteeism from school in an Australian region. Per child involved in the study, the data includes the number of students with average or slow learning speed Lrn, as well as Sex. One question is: do students of a given sex learn faster than students of the other sex? In other words: is there evidence for association between learning speed and sex? This problem is different from the of comparing two groups with a Student's-t test, because here both variables (Lrn and Sex) are grouping variables, whilst for the Student's-t test one variable is continuous and the other is a grouping variable.

We can examine the data to check for evidence of this association. This can be done by computing the number of cases within each class of learning speed Lrn and, of those, how many are of each Sex. We can also display the results as a barplot.

```
library(MASS)
data(quine)
str(quine)
## 'data.frame':
                    146 obs. of 5 variables:
   $ Eth : Factor w/ 2 levels "A", "N": 1 1 1 1 1 1 1 1 1 1 ...
## $ Sex : Factor w/ 2 levels "F", "M": 2 2 2 2 2 2 2 2 2 2 ...
## $ Age : Factor w/ 4 levels "F0", "F1", "F2", ...: 1 1 1 1 1 1 1 1 2 2 ...
## $ Lrn : Factor w/ 2 levels "AL", "SL": 2 2 2 1 1 1 1 1 2 2 ...
## $ Days: int 2 11 14 5 5 13 20 22 6 6 ...
# Examine how many observations you have per age group
table(quine$Lrn, quine$Sex)
##
##
         F M
##
     AL 40 43
     SL 40 23
##
# Now display the data in this table as a barplot
plot(quine$Lrn ~ quine$Sex)
```



How do we decide, on the basis of the number of cases for each learning speed and sex, whether or not the two display association?

Working examples

1 7 29

```
set.seed(55498)
n <- 100
x1 <- rbinom(n, prob = 0.3, size = 1)
y1 <- rbinom(n, prob = 0.2, size = 1)

x2 <- x1
y2 <- rbinom(n, prob = 0.8*x2, size = 1)</pre>
```

Let us consider studies involving per case the observation of two variables, X and Y, each one having values 0, 1. A table of the observed results is, in study 1:

```
table(x1, y1)
##
      у1
## x1
       0 1
##
     0 53 11
##
     1 27 9
and in study 2:
table(x2, y2)
      у2
##
        0
## x2
           1
##
     0 64
          0
```

Looking at the tables it is difficult to decide whether or not the variables X and Y are independent or not. In

fact, what sort of table would we expect to obtain, if X and Y were independent?

General setup

Consider the generic problem where two grouping variables X, Y are observed, each with groups labelled by either 0 or 1. If X and Y are independent, we would expect that the number of cases with X=0 to be the same, regardless of the value of Y. The estimated probability of a case having X=0 is just the proportion of cases with X=0 in the study - say px0. As a consequence, the probability of a case having X=1 is 1-px0. The same holds for the number of cases with Y=0. We will represent by py0 the probability that Y=0, so 1-py0 represents the probability that of a case having Y=1. If a total of Y=1 is Y=1 is Y=1 is Y=1 in the expected number of cases having Y=1 is Y=1 is Y=1 in the expected number of cases having Y=1 is Y=1 in the expected number of cases having Y=1 is Y=1 in the expected number of cases having Y=1 is Y=1 in the expected number of cases having Y=1 is Y=1 in the expected number of cases having Y=1 is Y=1 in the expected number of cases having Y=1 is Y=1 in the expected number of cases having Y=1 is Y=1 in the expected number of cases having Y=1 is Y=1 in the expected number of cases having Y=1 is Y=10.

If X and Y are independent, it follows the number of cases with both X=0 and Y=0 is expected to be N px0 py0, obtained from the product of the separate probabilities of each value of X and Y, in the study. For an overview, here is the table of expected frequencies in terms of these probabilities is:

Frequencies	Y=0	Y=1	Total
X=0	N px0 py0	N px0 (1-py0)	N px0
X=1	N (1-px0) py0	N (1-px0) (1-py0)	N(1-px0)
Total	N py0	N (1-py0)	N

So, if X and Y are independent, we know what to expect. Tests for independence of X and Y compare observed tables with this expected table. The expected table is constructed given the total counts for X and Y per possible value, so the row and column totals. These are called the *marginal frequencies* of X and Y. So tests for independence consider the row and column totals as fixed.

Note that, given the row and column totals, we need only fix one of the entries in the table, and all others are subsequently determined.

Working examples (cont)

For the first example, the row totals are 64 and 36 for X=0 and X=1 respectively, whilst the column totals are 80 and 20 for Y=0 and Y=1 respectively. This means that the empirical probabilities px0, py0 are respectively 0.64 and 0.8.

If X and Y are independent, the expected table of frequencies given these marginal frequencies is:

Frequencies	Y=0	Y=1	Total
X=0	N 0.64 * 0.8	N 0.64 * 0.2	N 0.64
X=1	N 0.36 * 0.8	N 0.36 * 0.2	N 0.36
Total	N 0.8	N 0.2	N

where N in this case equals 100.

The table formed by the observed marginal frequencies, but not taking into account the actual observations, is

Frequencies	Y=0	Y=1	Total
X=0	x0y0	x0y1	64
X=1	x1y0	x1y1	36
Total	80	20	100

where x0y0 and x0y1 represent the counts on the row corresponding to X=0, for Y=0 and Y=1 respectively. Similarly, x1y0 and x1y1 represent the counts on the row corresponding to X=1, for Y=0 and Y=1 respectively.

So, it is possible to observe any table of this form, so long as the counts x0 y0, x0 y1, x1 y0 and x1 y1 are such that the marginal frequencies are observed as given.

As we pointed out, we need only choose a value for one of the entries, and that determines all the other entries in the table. For example, say we set x0 y0 = 1. Then the table becomes:

Frequencies	Y=0	Y=1	Total
X=0	1	64 -1	64
X=1	80 -1	36 - 80 + 1	36
Total	80	20	100

Check that the same happens, regardless of the entry chosen. Fix a value for x1 y0 and replace it in the table, and check which values you get for the remaining entries.

The chi-square test

One commonly used test for independence in 2x2 tables is the chi-square test. It involves comparing the observed counts in the table with what would be expected, if X and Y were independent. Given the marginal frequencies, one one entry in the table is needed to determine the entire table.

The test compares the observed (O) and expected (E) counts of one entry (say x0 y0) by means of the test statistic:

$$\chi = \frac{(O-E)^2}{E^2}$$

The null and alternative hypotheses are

 H_0 : X and Y are independent vs. H_a : X and Y are not independent

Under the null hypothesis, χ follows asymptotically a chi-square distribution with 1 degree of freedom - this because, given the marginal frequencies, the table is determined once one entry is fixed.

The term asymptotically means that the distribution of χ can be approximated by the chi-square with 1 d.f., under H_0 , if the sample size is large enough. The concept of 'large enough' may differ depending on the researcher. Many researchers see this as requiring at least 5 observations in all entries of the table. For others, this means that all expected frequencies must be at least 5. The function chisq.test in R will give you a warning if the table used does not warrant the use of the chi-square test. If indeed the frequencies are too small to guarantee that the distribution of chi can be approximated by the chi-square, Fisher's exact test can be used (see below).

Working examples (cont)

We now apply the test to the working examples we had. For the first table we have:

table(x1, y1)

```
## y1
## x1 0 1
## 0 53 11
## 1 27 9
```

```
chisq.test(table(x1, y1))
##
   Pearson's Chi-squared test with Yates' continuity correction
##
##
## data: table(x1, y1)
## X-squared = 0.45844, df = 1, p-value = 0.4984
Now for the second table:
myt <- table(x2, y2)
chisq.test(myt)
##
##
   Pearson's Chi-squared test with Yates' continuity correction
##
## data: myt
## X-squared = 68.754, df = 1, p-value < 2.2e-16
Note that the test result can be saved and separate slots of the test can be extracted. Here we extract the
test statistic, the degrees of freedom and the p-value:
my.chtest <- chisq.test(myt)</pre>
my.chtest$statistic
## X-squared
## 68.75379
my.chtest$p.value
## [1] 1.11556e-16
my.chtest$parameter # this is the slot containing the p-value -- see ?chisq.test
## df
## 1
```

Fisher's exact test

The Fisher's exact test is another test that can be used to check whether or not two binary variables X and Y are independent. It calculates the probability of each possible 2x2 table of being observed, given the marginal frequencies. Then it adds up the probability of all tables at least as extreme as the one observed to yield a p-value. As it does not require approximations to compute the p-value, it is referred to as the 'exact' test, in contrast to the chi-square test with an approximated chi-square distribution. Yielding an exact p-value, it can be applied to all datasets.

Working examples (cont)

p-value = 0.4361

We will apply it now to the two tables of the examples. For the first example:

```
##
## Fisher's Exact Test for Count Data
##
## data: table(x1, y1)
```

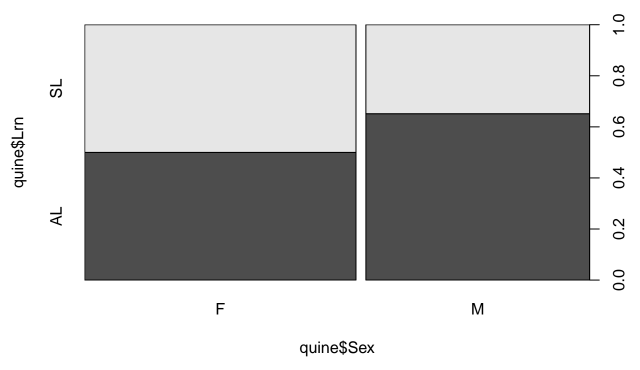
```
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 0.5169743 4.8487080
## sample estimates:
## odds ratio
##
     1.598236
Now for the second example:
fisher.test(table(x2, y2))
   Fisher's Exact Test for Count Data
##
##
## data: table(x2, y2)
## p-value < 2.2e-16
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 48.98022
                   Tnf
## sample estimates:
## odds ratio
##
          Tnf
As for the chi-square test result, slots of the test can be extracted. For example, we extract the p-value:
fisher.test(table(x2, y2))$p.value
## [1] 6.726116e-19
```

Quick tasks

1. Use both the Fisher's exact test as well as the chi-square test to decide if there is evidence of association between learning speed and sex in the quine data. Extract the p-value of each test, and add the p-value to the barplot of the data.

```
fisher.test(quine$Lrn, quine$Sex)
##
   Fisher's Exact Test for Count Data
##
## data: quine$Lrn and quine$Sex
## p-value = 0.09285
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 0.258569 1.099752
## sample estimates:
## odds ratio
## 0.5371981
myf.test <- fisher.test(quine$Lrn, quine$Sex)</pre>
mych.test <- chisq.test(quine$Lrn, quine$Sex)</pre>
plot(quine$Lrn ~ quine$Sex, main = paste("Chi-square p=", round(mych.test$p.value, 2),
                                          "Fisher ex p=", round(myf.test$p.value, 2)))
```

Chi-square p= 0.09 Fisher ex p= 0.09



2. Now consider two other datasets involving variables X and Y, with the cross-tabulations of the variables given below - first for study 1 1:

```
table(x1, y1)
```

```
## y1
## x1 0 1
## 0 53 11
## 1 27 9
```

Now for study 2:

```
table(x2, y2)
```

```
## y2
## x2 0 1
## 0 64 0
## 1 7 29
```

Try to use both the chi-square test and Fisher's exact test to investigate whether or not the data supports the null hypothesis of independence between ${\tt X}$ and ${\tt Y}$.

```
myt1 <- table(x1, y1)
chisq.test(myt1)

##

## Pearson's Chi-squared test with Yates' continuity correction
##

## data: myt1
## X-squared = 0.45844, df = 1, p-value = 0.4984

fisher.test(myt1)</pre>
```

```
##
  Fisher's Exact Test for Count Data
##
##
## data: myt1
## p-value = 0.4361
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 0.5169743 4.8487080
## sample estimates:
## odds ratio
     1.598236
myt2 \leftarrow table(x2, y2)
chisq.test(myt2)
##
   Pearson's Chi-squared test with Yates' continuity correction
##
## data: myt2
## X-squared = 68.754, df = 1, p-value < 2.2e-16
fisher.test(myt2)
##
##
    Fisher's Exact Test for Count Data
##
## data: myt2
## p-value < 2.2e-16
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 48.98022
## sample estimates:
## odds ratio
##
          Inf
```

Testing independence in nx2 tables

Motivation

Consider now the relationship between learning speed Lrn and age Age in the quine data, from the package MASS:

```
library(MASS)
table(quine$Age, quine$Lrn)

##

## AL SL

## F0 19 8

## F1 15 31

## F2 16 24

## F3 33 0
```

As before, we would like to know if data for the two variables involved, Age and Lrn, suggests that they are independent or not. This table has now more than 2 rows, so Fisher's exact test is not trivial to apply. However, the chi-square test can be used.

Working examples

```
set.seed(55498)
n <- 100
x1 <- rbinom(n, prob = 0.3, size = 2)
y1 <- rbinom(n, prob = 0.2, size = 1)

x2 <- x1
y2 <- rbinom(n, prob = 0.8*x2, size = 1)</pre>
```

```
## Warning in rbinom(n, prob = 0.8 * x2, size = 1): NAs produced
```

Let us consider studies involving per case the observation of two variables, X and Y, each one having values 0, 1. A table of the observed results is, in study 1:

```
table(x1, y1)

## y1
## x1 0 1
## 0 38 7
## 1 33 10
## 2 9 3
and in study 2:
table(x2, y2)
```

```
## y2
## x2 0 1
## 0 45 0
## 1 8 35
## 2 0 0
```

Looking at the tables it is difficult to decide whether or not the variables X and Y are independent or not. In fact, what sort of table would we expect to obtain, if X and Y were independent?

General setup

We will extend the setup previously used for 2x2 tables to the case where the table has more than 2 rows. As before, we assume that we are studying two grouping variables X and Y, where X may assume n values and Y may assume 2.

Say that X may assume 3 values, namely 0, 1 and 2. Then, analogously to the 2x2 table, the nx2 table in terms of empirical probabilities is:

Frequencies	Y=0	Y=1	Total
X=0	N px0 py0	N px0 (1-py0)	N pxO
X=1	N px1 py0	N px1 (1-py0)	N px1
X=2	N (1-px0-px1) py0	N (1-px0-px1) (1-py0)	N(1-px0-px1)
Total	N pyO	N (1-py0)	N

Tests for independence of X and Y compare observed tables with this expected table. The expected table is constructed given the total counts for X and Y per possible value, the marginal frequencies of X and Y. So tests for independence consider the row and column totals as fixed.

Note that, given the row and column totals, we need only fix one of the entries in the table, and all others

are subsequently determined.

Chi-square test for nx2 tables

The extension of the chi-square test to nx2 tables is straightforward. We will briefly give the details here.

As before, the marginal frequencies of the observed table are considered as fixed. Using the same reasoning as for 2x2 tables, the probability of each entry in the table is equal to the product of the marginal probabilities.

For each row, since the marginal frequency (total row count) is given, it suffices to fix one of the entries, and the other one is determined. In addition, once one but the last row has one entry fixed, the last row is determined based on the total column counts. This means that, for the first n-1 rows, it suffices to fix one entry to determine the row, and this determines the last row. For this reason, the chi-square test for a nx2 table has n-1 degrees of freedom.

Working examples (cont)

Analogously to the development for 2x2 tables, if X and Y are independent in this 3x2 table example, the expected table of frequencies given these marginal frequencies is, for study 1:

Frequencies	Y=0	Y=1	Total
X=0	N 0.45 * 0.8	N 0.45 * 0.2	N 0.45
X=1	N 0.43 * 0.2	N 0.43 * 0.2	$N \ 0.43$
X=2	N 0.12 * 0.8	N 0.12 * 0.2	N 0.12
Total	N 0.8	N 0.2	N

where N in this case equals 100.

The table formed by the observed marginal frequencies, but not taking into account the actual observations, is

Frequencies	Y=0	Y=1	Total
$\overline{\mathbf{X}=0}$	х0у0	x0y1	45
X=1	x1y0	x1y1	43
X=2	x2y0	x2y1	12
Total	80	20	100

where x0y0 and x0y1 represent the counts on the row corresponding to X=0, for Y=0 and Y=1 respectively. Similarly, x1y0 and x1y1 represent the counts on the row corresponding to X=1, for Y=0 and Y=1 respectively. Finally, x2y0 and x2y1 represent the counts on the row corresponding to X=2, for Y=0 and Y=1 respectively.

So, it is possible to observe any table of this form, so long as the counts x0y0, x0y1, x1y0, x1y1, x2y0 and x2y1, are such that the marginal frequencies are observed as given.

As we pointed out, we need only choose a value for one of the entries per row, and that determines the other entry in the row - except for the last row, which is determined if the first n-1 are filled in. For example, say we set all entries on the first column, for all but the last row, equal to 1. Then the table becomes:

Frequencies	Y=0	Y=1	Total
X=0	1	45 -1	45
X=1	1	43 -1	43
X=2	80 -2	12 - 80 + 2	12

Frequencies	Y=0	Y=1	Total
Total	80	20	100

You may wish to check for yourself that the same happens, regardless of the entries chosen to be fixed.

Quick tasks

1. Use the chi-square test to decide if there is evidence of association between learning speed and age in the quine data. Extract the p-value of the test, and add the p-value to the barplot of the data.

```
mych.test <- chisq.test(quine$Lrn, quine$Age)

plot(quine$Lrn - quine$Age, main = paste("Chi-square p=", round(mych.test$p.value, 2) ) )

Chi-square p= 0

Fo F1 F2 F3

quine$Age
```

2. Now consider two other datasets involving variables X and Y, with the cross-tabulations of the variables given below - first for study 1:

```
table(x1, y1)

## y1
## x1 0 1
## 0 38 7
## 1 33 10
## 2 9 3

Now for study 2:
table(x2, y2)
## y2
```

```
## x2 0 1
## 0 45 0
## 1 8 35
## 2 0 0
```

Use a test to investigate whether or not the data supports the null hypothesis of independence between ${\tt X}$ and ${\tt Y}$.

```
myt1 <- table(x1, y1)
chisq.test(myt1)
## Warning in chisq.test(myt1): Chi-squared approximation may be incorrect
##
   Pearson's Chi-squared test
##
##
## data: myt1
## X-squared = 1.0279, df = 2, p-value = 0.5981
myt2 \leftarrow table(x2, y2)
chisq.test(myt2)
## Warning in chisq.test(myt2): Chi-squared approximation may be incorrect
    Pearson's Chi-squared test
##
##
## data: myt2
## X-squared = NaN, df = 2, p-value = NA
```

Testing symmetry in 2x2 tables

Motivation

Patients suspected of having lung cancer get a scan to check if the scan includes anomalies. The scan is analysed by a radiologist, but this may take a few days.

A computer program has been developed to analyse the images and detect anomalies. This is quicker than the radiologist, as the computer can be left continuously running. Researchers want to know if the computer program detects more anomalies than the radiologist or not. How can we decide on this?

Note: any anomaly detected is a positive result so, for the purpose of this example, there are no anomalies not worth being detected (false positives).

Working example

The data consists of two variables: the result of the radiologist res.radio and the result of the computer program res.comp. Each variable has one entry per scan, indicating by the value 1 if an anomaly was found in the scan, and by the value 0 otherwise.

To assess how often the two tests give similar results and how often they do not, we make a table of these variables:

```
table(res.radio, res.comp)

## res.comp

## res.radio no anomaly anomaly
## no anomaly 45 12
```

```
## anomaly 7 36
```

Now we need to decide if the difference observed is large enough to suggest one test detects more anomalies than the other.

General setup

In general, two variables X and Y are observed, each corresponding to one test test: pass or fail. A typical table observed is

Frequencies	Y=fail	Y=pass
X=fail	a	b
X=pass	С	d

The counts a and d correspond to cases where both tests yield the same result: in a cases both tests indicate a fail, and in d cases both tests indicate a pass. So, to decide on whether one of the two tests is better than the other, only counts b and c are informative.

If the tests disagree in a random manner, without a trend towards one yield a pass more often than the other, we expect to see similar values for b and c, relative to all results that disagree b+c. If test X more often yields a pass than test Y, we would expect to see c relatively larger than b, compared with b+c.

The statistical test to answer this question is McNemar's test. It is implemented in R via the mcnemar.test function. The test computes the probability of observing one of the discordant counts (b, say), or a more extreme value, out of a total b+c counts, at random. For that, typically an approximation of the true binomial distribution to the chi-square (with 1 degree of freedom) is used. In this course, we will use this function for our illustrations.

Working example (cont)

In the study with the scans evaluated by a radiologist and a computer program, we had:

```
table(res.radio, res.comp)
##
               res.comp
## res.radio
                no anomaly anomaly
##
     no anomaly
                         45
                                 12
     anomaly
                                 36
Now applying McNemar's test to the data:
mcnemar.test(table(res.radio, res.comp))
##
##
   McNemar's Chi-squared test with continuity correction
##
## data: table(res.radio, res.comp)
## McNemar's chi-squared = 0.84211, df = 1, p-value = 0.3588
```

Quick tasks

1. A study of twins submits each twin to the same test. Twin 1 is just told to complete the test, whilst twin 2 receives extra explanation before completing the test. The study then evaluates whether or not

the explanation helps, i.e., twin 2 performs better than twin 1 in the test. The table of the test results is:

```
table(Twin1, Twin2)
```

```
## Twin2
## Twin1 fail pass
## fail 66 34
## pass 13 87
```

Use a test to decide whether twin 2 performs better than twin 1.

```
mcnemar.test(table(Twin1, Twin2))
```

```
##
## McNemar's Chi-squared test with continuity correction
##
## data: table(Twin1, Twin2)
## McNemar's chi-squared = 8.5106, df = 1, p-value = 0.003531
```

Notes

An exact version of the test makes use of the binomial distribution directly. An implementation of this version is also available in R, for example via the exact2x2 package from Michael Fay, which can be found via the link below:

https://cran.r-project.org/web/packages/exact2x2

Exercises

1. Patients who regular have migraine often also have photophobia. A study was setup to evaluate whether a new treatment helps decrease the frequency of photophobia occurence. Patients first answer a questionnaire, indicating whether they had at least one migraine with photophobia in the last 6 months (photophobia = 1) or not (photophobia = 0). Six months after the new treatment is completed, they fill in the questionnaire again, indicating whether they had at least one migraine with photophobia in the 6 months after treatment or not. The observed table is as follows:

table(BeforeTreatment, AfterTreatment)

```
## AfterTreatment
## BeforeTreatment noPhoto Photo
## noPhoto 155 58
## Photo 39 148
```

Perform a test to decide whether the results after treatment lead to more often, or less often, photophobia than before treatment.

2. A case-control study

A study has been put together to better understand whether the exposure to environmental factors is associated with a rare type of cancer or not. To do this, cases of this cancer are included and, per case, a matching control is selected, which is similar to its corresponding case in all important covariates. The data consists thus of pairs of individuals, one being the case and one its matching control. Perform a test to decide whether exposure occurs more often in one of the groups (case or control).

```
## Controls
## Cases noExp Exp
## noExp 9 10
```

```
## Exp 36 245
```

Relative risk and odds ratio

Motivation

Researchers want to better understand the relationship between smoking and stroke. For this, they have collected data of people who have suffered a stroke in the last 5 years, and from people with similar demographic variables who have never had a stroke. They have then asked these people whether or not they smoked cigarettes regularly, or had done in the past for at least a year, in the 5 years prior to the stroke these people were considered smokers. All others were considered non-smokers.

This study thus involves cases and controls, and the latter are chosen so as to have a similar profile as that of cases, but there is no 1-1 matching between cases and controls - so no pairs (case, control). The data consists of two variables: smoke indicating whether an individual is categorized as a smoker (yes) or not (no); and stroke indicating whether an individual has had a stroke (1) or not (0).

The observed data can be summarized by tabulating smoke and stroke:

```
table(stroke, smoke)
```

```
## smoke
## stroke no yes
## 0 107 26
## 1 43 74
```

Some of the questions researchers want to address are:

- is the risk of having a stroke different between smokers and non-smokers?
- are the odds of having a stroke larger for smokers, compared with non-smokers?

The first question relates to the risk of having a stroke given that someone is a smoker, relative to the risk of having a stroke given that the person is not a smoker. This is what we call a *relative risk*. In this context, *risk* has the meaning of probability.

The second question relates to odds, the relation between the number of people who get a stroke and the number of people who do not, given their smoking status. The comparison between the two odds is made via the *odds ratio*.

We will see these concepts in more detail in the following sections.

Relative risk

Consider a generic table observed as

Frequencies	Y =0	Y=1	Total
Cases Controls Total	a	b	a+b
	c	d	c+d
	a+c	b+d	N

where the patient status (case or control) is the variable of interest, and Y represents the explanatory grouping variable. There is interest in studying the association between the patient status and the explanatory variable.

The risk of being a case, given that Y=0, is calculated by a/(a+c). Similarly, the risk of being a case, given that Y=1, is calculated by b/(b+d). So, the relative risk is the ratio between these two risks, or

```
(a/b)*(b+d)/(a+c).
```

For the example of stroke and smoking above, we have cases on row 2 and controls on row 1. In this case, the table is

Frequencies	Y =0	Y =1	Total
Controls	107	26	133
Cases	43	74	117
Total	150	100	250

Then the risk of stroke amongst non-smokers is 43 divided by 150, which gives

```
myt <- table(stroke, smoke)</pre>
r.stroke.smoke0 <- myt[2, 1]/colSums(myt)[1]</pre>
r.stroke.smoke0
##
           nο
## 0.2866667
Similarly, the risk of stroke amongst smokers is 74 divided by 100, which gives
r.stroke.smoke1 <- myt[2, 2]/colSums(myt)[2]</pre>
r.stroke.smoke1
## yes
## 0.74
The relative risk of having a stroke in smokers compared to non-smokers is then 0.74 divided by 0.29, yielding
rr <- r.stroke.smoke1/r.stroke.smoke0
round(rr, 4)
##
      yes
```

So, the chance of someone having a stroke is 2.58 larger amongst smokers than amongst non-smokers.

Quick tasks

2.371795

2.5814

Researchers are looking at the relationship between colon cancer and BMI. For this, they put together a prospective study, where patients are followed up. Those who developed colon cancer also had their BMI at time of diagnosis recorded. As reference, they used the individuals who did not develop colon cancer during the study. The resulting table of the explanatory variable bmi (categorized as either high or normal-low) and the response colon.cancer (categorized as 1 for cases and 0 for controls) is:

```
## bmi
## colon.cancer normal-low high
## 0 185 115
## 1 65 185
```

Compute the relative risk of developing colon cancer given high BMI, compared to that given normal or low BMI.

```
myt <- table(colon.cancer, bmi)
rr.colon <- (myt[2, 2]/colSums(myt)[2])/(myt[2, 1]/colSums(myt)[1])
rr.colon
## high</pre>
```

Odds ratio

Consider again the generic table observed as

Frequencies	Y =0	Y =1	Total
Cases	a	b	a+b
Controls	c	d	c+d
Total	a+c	b+d	N

The odds of being a case amongst individuals with Y=1 is b/d, whilst the odds of being a case amongst individuals with Y=0 is a/c. Then the odds ratio of being a case, of individuals with Y=1 compared to that of individuals with Y=0, is the ratio (b/d)/(a/c), qhich is the same as (bc)/(ad).

For the example of stroke and smoking above, we have the odds of having a stroke amongst non-smokers being 65 divided by 185. Similarly, the odds of having a stroke amongst smokers is 185 divided by 115. This leads to the following odds: 0.35 amongst non-smokers and 1.61 amongst smokers. Then the odds ratio of having a stroke, between smokers and non-smokers, is 1.61 divided by 0.35, which gives 4.58.

```
myt <- table(stroke, smoke)
or <- (myt[2, 2] * myt[1, 1]) / (myt[1, 2] * myt[2, 1])</pre>
```

Note again that, in this case, cases are on row 2 and controls on row 1.

So, the odds of someone having a stroke is 7.08 larger amongst smokers than amongst non-smokers.

Quick tasks

Compute now the odds ratio of developing colon cancer given high BMI, compared to that given normal or low BMI.

```
myt <- table(colon.cancer, bmi)
or.colon <- (myt[2, 2]/myt[1, 2])/(myt[2, 1]/myt[1, 1])
or.colon</pre>
```

[1] 4.578595

Relative risk vs. odds ratio

In principle, both relative risk and odds ratio are statistics that can be computed, given any 2x2 tables. However, their usefulness, and interpretation, depends on the experimental design used.

The relative risk is evaluated assuming that the relative frequencies a/(a+c) and b/(b+d) are good approximations for the probabilities of being a case, given Y=0 and Y=1, respectively. Since calculations are conditional on the marginal frequencies a+c and b+d, these must represent well the frequencies of Y=0 and Y=1 in the general population, in the sense that (b+d)/(a+c+b+d) is a good estimate of the proportion of individuals with Y=1 in the population under study.

In contrast, the odds ratio does not make use directly of the observed relative frequencies. So it needs not rely on representative (relative) marginal frequencies.

Typically relative marginal frequencies can only be expected in the context of prospective studies. Retrospective studies, especially case-control sudies, do not satisfy this assumption. For this reason, odds ratios are often used in the context of case-control studies, whilst relative risks can only be reliably interpreted in prospective (cohort) studies.

In some cases, the computed relative risk and odds ratio yield similar values. For example, if the marginal frequency a+c is approximately equal to c, and the marginal frequency b+d is approximately equal to d, then it is easy to see that the two statistics will yield similar results. One situation where both $a+c\approx c$ and $b+d\approx d$ occur is when the response involves rare cases, so that both a and c are small, and the marginal frequencies of both Y=0 as well as Y=1 are large enough.

Logistic regression models (cont)

The concepts of relative risk and odds ratio have been introduced above, and we computed them on the basis of 2x2 tables. This helps with understanding what they mean. However, 2x2 tables can only represent the relationship between a response (having a stroke or not) and one explanatory (grouping) variable (smoking habit). Most studies involve more variables, and thus more general methods to compute them are needed.

To compute the odds ratio, a logistic regression can be used, allowing for the inclusion of more explanatory variables. Here we will check that this is the case. Indeed, if we fit a logistic regression to the stroke and smoking habit example, we get:

```
lgfit <- glm(stroke ~ smoke, family = binomial)
summary(lgfit)</pre>
```

```
##
## Call:
## glm(formula = stroke ~ smoke, family = binomial)
##
## Deviance Residuals:
##
      Min
               1Q
                   Median
                                3Q
                                       Max
## -1.641 -0.822
                  -0.822
                             0.776
                                     1.581
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
                             0.1806
                                     -5.049 4.44e-07 ***
##
                -0.9116
  (Intercept)
##
  smokeyes
                  1.9576
                             0.2908
                                      6.731 1.68e-11 ***
##
                     '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## (Dispersion parameter for binomial family taken to be 1)
##
                                       degrees of freedom
##
       Null deviance: 345.55
                               on 249
## Residual deviance: 294.35
                               on 248
                                       degrees of freedom
  AIC: 298.35
##
## Number of Fisher Scoring iterations: 4
sfit <- summary(lgfit)$coef</pre>
```

The coefficient of the smoke variable (corresponding to smoke=yes) in the model is 1.96. It turns out that this is the same as the logarithm of the odds ratio:

```
log(or)
```

```
## [1] 1.957597
```

Alternatively, we compute exp(beta), where beta is the coefficient of smoke=yes estimated by the logistic model. This gives 7.08, which is the same as the odds ratio 7.08 computed from the table.

```
exp(sfit[2, 1])
```

```
## [1] 7.08229
or
```

```
## [1] 7.08229
```

1.399669 2.542158

So, the log-odds ratio corresponds to the coefficient of a binary variable from a logistic regression. If more explanatory variables are included in the logistic regression, the computed log-odds ratio yields a measurement that is corrected for the effects of the remaining variables.

As we already know, a confidence interval for a coefficient of the fitted regression can be computed. We can use the function confint on the model fit object to do that:

```
confint(lgfit, 2)

## Waiting for profiling to be done...

## 2.5 % 97.5 %
```

This is the 95% confidence interval for the logarithm of the odds ratio. Typically there is interest in testing whether the odds ratio is equal to 1 (in which case the odds are the same in both groups). On this scale, this test is equivalent to testing whether the log-odds ratio is equal to 0. The above confidence interval does not include 0, indicating that the null hypothesis that the odds ratio is equal to 1 can be rejected, with a significance level of 0.05 (because it is a 95% confidence interval).

An approximate confidence interval for the odds ratio is then

```
exp(confint(lgfit, 2))
## Waiting for profiling to be done...
## 2.5 % 97.5 %
## 4.053859 12.707069
```

Poisson regression models

While odds ratio can be computed by fitting a logistic regression, relative risk can be computed by fitting a Poisson regression.

For some background: the logistic regression assumes that the response observations can be modelled by a binomial distribution, whilst the Poisson regression assumes that the response observations can be modelled by a Poisson distribution. The binomial distribution models counts for a given total, whilst the Poisson distribution models counts without conditioning on a total. So, while both regression models are suitable for a response variable that represents counts, in the logistic regression computations are conditional on the total number of cases and controls observed, which is not the case with the Poisson regression.

We now fit a Poisson regression to the stroke and smoking habits table:

```
lgfit <- glm(stroke ~ smoke, family = poisson)
summary(lgfit)

##
## Call:
## glm(formula = stroke ~ smoke, family = poisson)</pre>
```

```
## Min 1Q Median 3Q Max
## -1.2166 -0.7572 -0.7572 0.2867 1.0355
##
```

Coefficients:

Deviance Residuals:

##

```
##
               Estimate Std. Error z value Pr(>|z|)
                -1.2494
                                     -8.193 2.54e-16 ***
                             0.1525
## (Intercept)
                             0.1917
  smokeyes
                  0.9483
                                      4.946 7.59e-07 ***
##
##
  Signif. codes:
                     '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
   (Dispersion parameter for poisson family taken to be 1)
##
##
##
       Null deviance: 177.67
                               on 249
                                        degrees of freedom
## Residual deviance: 152.01
                               on 248
                                        degrees of freedom
  AIC: 390.01
## Number of Fisher Scoring iterations: 5
sfit <- summary(lgfit)$coef</pre>
```

The coefficient of the smoke variable (corresponding to smoke=yes) in the fitted Poisson regression is 0.95. It turns out that this is the same as the logarithm of the relative risk:

```
log(rr)
```

```
## yes
## 0.9483301
```

Alternatively, we can compute the relative risk via exp(beta), where beta is the estimated coefficient of smoke=yes in the Poisson regression. This yields 2.58, the same as the computed relative risk from the table 2.58.

```
exp(sfit[2, 1])

## [1] 2.581395

rr

## yes

## 2.581395
```

Similarly to what was done for the odds ratio, we can now build a confidence interval for the coefficient of the fitted regression. We use the function **confint** on the model fit object to do that:

```
confint(lgfit, 2)

## Waiting for profiling to be done...

## 2.5 % 97.5 %

## 0.5776317 1.3315650
```

This is the 95% confidence interval for the logarithm of the relative risk. Typically there is interest in testing whether the relative risk is equal to 1 (in which case the risk the same in both groups). On this scale, this test is equivalent to testing whether the log-relative risk is equal to 0. The above confidence interval does not include 0, indicating that the null hypothesis that the relative risk is equal to 1 can be rejected, with a significance level of 0.05 (because it is a 95% confidence interval).

Quick tasks

Using regression models, recompute the relative risk and odds ratio of having colon cancer, amongst individuals with high BMI relative to individuals with normal or low BMI. Note: before you have computed relative risk and odds ratio using the counts table. For this task, you need the data consisting of the variables recording

one value per individual, which can be found in file data_colon_RR_OR.txt. Add the computed values to the text below. Compare results you found with those computed using the tables in previous exercises.

```
data.colon <- read.delim("data_colon_RR_OR.txt")

lgfit <- glm(colon.cancer ~ bmi, family = binomial)
log.or <- summary(lgfit)$coef[2, 1]
or <- exp(log.or)

pfit <- glm(colon.cancer ~ bmi, family = poisson)
log.rr <- summary(pfit)$coef[2, 1]
rr <- exp(log.rr)</pre>
```

Using data from the colon cancer study, we found that the relative risk of developing colon cancer, amongst individuals with high BMI compared to those with normal or low BMI, was 2.37. For the same data, the odds ratio of colon cancer, amongst individuals with high BMI compared to those with normal or low BMI, was 4.58.

Power analysis for count tables

Motivation

We want to know how many samples are needed if we were to repeat the study of learning speed and association with sex, using as pilot the data the quine data. Let us first review the data and make the table we will use for analyses:

```
library(MASS)
# Examine how many observations you have per age group
table(quine$Lrn, quine$Sex)

##
## F M
## AL 40 43
## SL 40 23
mytable <- table(quine$Lrn, quine$Sex)</pre>
```

The aim is to test if there is a statistically significant difference between the proportions of females, between the groups defined by learning speed. We will test this by using either Fisher's exact test or the chi-square test. Subsequently, we wish to know by how much the sample size would have to change, to modify the test's conclusions.

Fisher's exact test

Fisher's exact test for independence of the marginal counts in this table gives:

```
fisher.test(mytable)
```

```
##
## Fisher's Exact Test for Count Data
##
## data: mytable
## p-value = 0.09285
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 0.258569 1.099752
```

```
## sample estimates:
## odds ratio
## 0.5371981
```

So the test is not significant if we use $\alpha=0.05$. Now we will evaluate the power of the test to find the difference between proportions to be significant, given the current table. For this, we make use of the function power.fisher.test from the package statmod, which is automatically installed when you install R. As there is no closed form for the distribution of the Fisher's exact test statistic under the alternative, the power cannot be evaluated on the basis of a mathematical formula. Instead, the function simulates data according to given parameters, and yields the estimated power.

As inputs, the function expects the proportions of females in the two groups (p1 and p2), the sample sizes in the two groups (the row sums), the significance level ($\alpha = 0.05$ is the default), the number of simulations and the alternative (one or two-sided). Below we use the function to compute the power for the table of learning speed and sex from the quine data:

```
library(statmod)
power.fisher.test(p1 = 40/83, p2 = 40/63, n1 = 83, n2 = 63)
```

[1] 0.49

The estimated power is lower than 0.8, the level typically desired for studies. We could now increase the sample size and check if the desired power of 0.8 would be achieved. Note that we keep the proportions p1 and p2 the same, so that the effect size is the same as for the original data. We also keep the relative proportions in groups 1 and 2 the same, i.e., n1/n2 is kept fixed. We can for example multiply both sample sizes by 2:

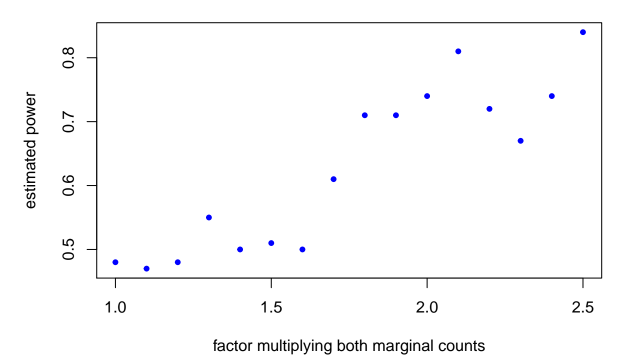
```
power.fisher.test(p1 = 40/83, p2 = 40/63, n1 = 83*2, n2 = 63*2)
```

[1] 0.72

This yields a number closer to 0.8. Let us now define a vector of values which we will multiply both sample sizes by, and compute the power for each value:

Now we make a graph of the estimated power, as a function of the multiplying factor:

Power Fisher exact test



We used here a relatively small number of simulations, the default 100. This of course yields variability between estimates, meaning they are not always increasing.

So, a power of at least 0.80 is achieved for the last few multiplying factors, corresponding to minimum sample sizes required of:

```
min(sr[ powerv >= 0.8 ]*83)

## [1] 174.3

for group 1, and

min(sr[ powerv >= 0.8 ]*63)

## [1] 132.3

for group 2. This would yield a total sample size for a new study of:

min(sr[ powerv >= 0.8 ]*83) + min(sr[ powerv >= 0.8 ]*63)

## [1] 306.6
```

Note that Fisher's exact test makes no distinction between rows and columns, in the sense that the test statistic is the same, if we invert rows and columns. So, conclusions here would have been the same if we had used as inputs the proportions of learning speeds per sex, and the column sums, instead.

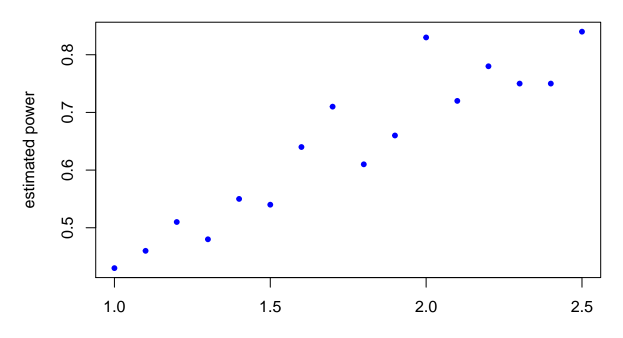
Quick tasks

Compute the power for a series of multiplying factors for the column sums of the learning speed and sex table of the quine data, considering now the problem of comparing proportions of average learners per group. Plot the estimated powers and determine the minimum sample size corresponding to 0.80 power.

```
sr <- seq(from = 1, to = 2.5, by = 0.1)
powerv <- NULL</pre>
```

The graph of the estimated power, as a function of the multiplying factor:

Power Fisher exact test



factor multiplying both marginal counts

So, a power of at least 0.80 is achieved for the last few multiplying factors, corresponding to minimum sample sizes required of:

```
min(sr[ powerv >= 0.8 ]*80)

## [1] 160
for group 1, and
min(sr[ powerv >= 0.8 ]*66)

## [1] 132
for group 2.
```

Chi-square test

The power for a chi-square test can be computed by using the function pwr.chisq.test of the package pwr. This function, as others from the same package, enables us to give all arguments but one, and the function will automatically determine the last argument using all the given ones. Here we will focus on determing the sample size, given the remaining arguments.

We will use in for the same example as above, where we studied the association between learning speed and sex, using the quine data. Let us first use the chi-square test to test for the association between the two

variables, and save the observed test statistic:

chisq.stat <- chisq.test(mytable)\$stat</pre>

```
chisq.test(mytable)

##

## Pearson's Chi-squared test with Yates' continuity correction

##

## data: mytable

## X-squared = 2.7949, df = 1, p-value = 0.09456
```

Based upon this test result, we conclude that there is no statistically significant association between learning speed and sex, with a significance level of 0.05. How large would the sample size need to be, in order to yield a statistically significant result?

We already knew that a chi-square test statistic for a 2x2 table has, under the null hypothesis of no association between rows and columns, asymptotically a chi-square distribution with 1 d.f.. The power we may state as 0.8, and the significance level remains 0.05.

We need also to state which effect size is required. This would be the effect size observed in the current data. It is equal to the square root of the observed test statistic, divided by the sample size of the current study. We have already computed the observed test statistic, and the sample size of the current study can be computed by sum(mytable). Then we have:

```
eff.size <- sqrt(chisq.stat/sum(mytable))</pre>
```

Now load the package required and compute the minimum sample size:

```
library(pwr)
pwr.chisq.test(w = eff.size, df = 1, power = 0.8)
```

```
##
##
        Chi squared power calculation
##
##
                  w = 0.1383592
                 N = 410.0061
##
##
                 df = 1
         sig.level = 0.05
##
##
             power = 0.8
##
## NOTE: N is the number of observations
```

Quick tasks

Using again the learning speed and sex example of the quine data, compute the sample size required to obtain a significant chi-square test if the significance level desired was 0.025, instead of 0.05, while keeping all other parameters the same.

```
pwr.chisq.test(w = eff.size, df = 1, power = 0.8, sig.level = 0.025)
```

```
##

## Chi squared power calculation

##

## w = 0.1383592

## N = 496.5209

## df = 1

## sig.level = 0.025
```

```
## power = 0.8
##
## NOTE: N is the number of observations
```

Odds ratio

The minimum sample size required to find a specific odds ratio as different from 1 is the same as the one required to find the log-odds ratio as different from 0. Thus, the power analysis required is the same as the one for a coefficient from the logistic regression.

Notes

We have used a choice of functions available in R to explain the concept of minimum sample size, and illustrate how it varies according to various parameters, for the chi-square and Fisher's exact tests. However, there are other choices of functions, which may be more convenient for a particular application.

To estimate the minimum sample size required for McNemar's test of symmetry, we suggest using the exact2x2 package. We refer to the vignette of the McNemar's test functions implemented in the package for more details:

https://cran.r-project.org/web/packages/exact2x2/vignettes/exactMcNemar.pdf

Exercises

Exercise 1

A study has looked at the relationship between aspirin use and heart attacks via a randomized clinical trial. The aim is to test whether aspirin taken regularly reduces mortality from cardiovascular disease. Study participants did not know if they used aspirin or a placebo. The table below summarizes some of their findings, according to fatal miocardial infarcts (FMI) and non-fatal ones (NFMI).

	FMI	NFMI	No attack
Placebo Aspirin	18 5	171 99	10845 10933

Source: example 2.2.4 (pp. 16-17) of Agresti, A. (1990) Categorical data analsis. Wiley, New York.

1.1. Compute the proportions of heart attack depending on the drug used. Hint: After entering the data, add up the columns corresponding to heart attack and use prop.table. Save the result as an object. Write a small bit of text where you indicate the proportions found.

The study found that there were r of those using aspirin, and r of those using placebo.

- 1.2 Compute the relative risk of heart attacks in the placebo group, compared with the aspirin group.
- 1.3 Compute the sample odds ratio of having a heart attack in the placebo group.
- 1.4 Write a short piece of text where the computed relative risk and odds ratio are included, with inline code.
- 1.5 Test the hypothesis that the use of aspirin, compared to placebo, has no effect on the chance of a heart attack, against the hypothesis that aspirin modifies the chance, without making an assumption about in which direction the modification may occur.
- 1.6 Extract the odds ratio estimates from 1.3 and 1.6, and include them as inline code in the text below.

The odds ratio estimated by us from the table was r, and the one estimated by Fishers exact test wasr'.

Exercise 2

A study looked at the association between job satisfaction and income, gathering the data in the table below.

	Very dissatisf.	Little dissatisf.	Moderately satisf.	Very satisf.
<6,000	20	24	80	82
6,000-15,000	22	38	104	125
15,000-25,000	13	28	81	113
>25,000	7	18	54	92

Source: example 2.3.2 (pp. 20-21) of Agresti, A. (1990) Categorical data analsis. Wiley, New York.

- 2.1 Enter the data into R by making a matrix. Assign meaningful row and columns names. Print out the matrix to check that it has been entered correctly.
- 2.2 Add the counts from the dissatisfaction columns up, keeping the split between income levels. Do the same for the satisfaction columns. Make a new table (or a matrix) with these two new columns.
- 2.3 Use the chi-square test to compare the spread of people across income classes between satisfied and dissatisfied groups. Print out the test result and save the p-value in an object.
- 2.4 Complete the text below with the values of this example using inline code.

This study recorded job satisfaction from r people across r income classes. A chi-square test for the count distribution across income classes yielded a p-value of r.

- 2.5 Compute the total sample size that would be required, in order to find the observed effect size as significant with a significance level $\alpha = 0.05$ and power 0.8. Note that the number of degrees of freedom is equal to the number of rows in the table, minus 1.
- 2.6 Compute log-odds ratios for each income class, compared with the lowest income class.
- 2.7 Complete the following text with inline code.

This study found that odds of being dissatisfied, compared with being satisfied, is for the income class >25,000 r that for the income class of <6,000.

Exercise 3

The table below gives the grade into which left and right eye of the same person were classified. Use an appropriate test to check if there is evidence that eye grades of one side are typically better than the other side, against the null hypothesis that the better grade is observed at random for the left and for the right eye.

	Left eye graded best	Left eye graded worst
Right eye graded best	3532	700
Right eye graded worst	597	2648

- 3.1 Enter the data as a matrix and define row and column names. Check that your definition is correct.
- 3.2 Use an appropriate test to check if there is evidence that eye grades of one side are typically better than the other side, against the null hypothesis that the better grade is observed at random for the left and for the right eye.

Exercise 4

A study compared radiation therapy with surgery in treating cancer of the larynx. The data is given below.

	Cancer controlled	Cancer not controlled
Surgery	21	2
Radiation therapy	15	3

- 4.1 Enter the data as a matrix, and define row as well as column names.
- 4.2 Use a test to check if there is evidence that the response (cancer controlled or not) is independent of therapy.
- 4.3 Compute the power of the test given the current data to find a difference between the two observed proportions.
- 4.4 Evaluate the power for larger sample sizes, keeping the relative sample size for the first and second groups the same. Make a graph of results, with the total expected sample size on the x-axis and the power on the y-axis.

Survival data analysis

Introduction

Motivation

A migraine study wants to evaluate the time to recovery given a new drug (D2), compared to a commonly used drug (D1). Patients with migraine are invited to take part in the study, where they are randomized into two groups, each given one of the drugs. In each group, participants must take the drug they are given shortly after a new migraine episode starts. They record the time of taking the drug, and the time the migraine has stopped. In the course of the study, some patients could not record the time of recovery as they fell asleep before it happened. In such cases, all patients recorded that they had recovered by the time they woke up.

The response in this case involves the event observed, recovered or not after a single dose of the assigned drug, as well as the time to event. The explanatory variable is the drug received. Of course, it is possible to analyse the association between each of the two variables with the drug used separately from each other, but doing this ignores important issues. Firstly, the event observed (recovered or not) may be related to the time to event. Secondly, there is interest in drawing conclusions for the combination of event as well as time-to-event variable, instead of for them separately. So, there is interest in considering the two variables as a single response. Indeed, a model that takes both event as well as time-to-event into account at the same time has more power to find effects than one that analyses the two variables separately. In addition, it yields conclusions about effects on both variables at the same time.

Another important aspect is that, for some participants, the precise time at which recovery happened was not recorded, but it is known to have been after a certain time point. This means that the time-to-event is not precisely known, but known to be at least a certain value, for these participants. Such observations are said to be *censored*.

We will now see how to analyse such a response variable, formed by a combination of an event variable with the time-to-event, where sometimes the time-to-event is not precisely known.

Working example

Consider the migraine study with the two drugs. The time-to-event is in this case the time (in minutes) between taking the assigned drug and the event. The event can be recovery (coded as 1) or fail (coded as 0), where the latter corresponds cases where the participant falls asleep without having recovered beforehand. Note that the time-to-event in case of a fail event is recorded as the time between taking the drug and the last moment the participant is known to not having recovered, which is when the participant fell asleep. As such, the time-to-event is at least as long as the time recorded, since migraine recovery may have recovered at any time while the participant slept.

```
set.seed(39562) # (493756)
ss <- 400
fdrug <- factor(rep(1:2, each = ss/2), labels = c("D1", "D2"))
event <- c(rbinom(ss/2, 1, prob = .7), rbinom(ss/2, 1, prob = .75))
tte <- round(rgamma(ss, shape = 30, rate = 1+event/4))
migraine <- data.frame(event, time = tte, drug = fdrug)
write.table(migraine, file = "migraine_data.txt", sep = "\t", row.names = FALSE)</pre>
```

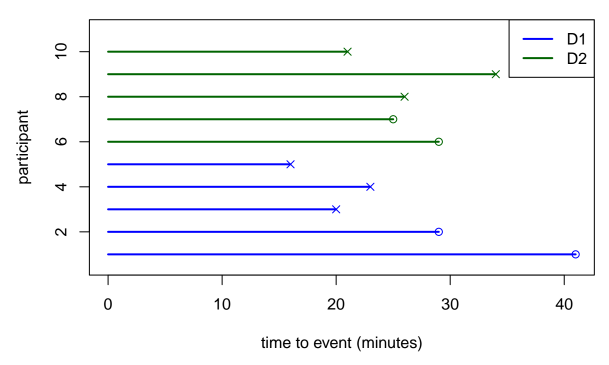
The observed data for a subset of 10 participants is as follows:

```
mydata <- rbind(migraine[1:5, ], migraine[(ss/2+1):(ss/2+5), ])
mydata</pre>
```

```
##
        event time drug
## 1
            0
                 41
                       D1
## 2
            0
                 29
                       D1
## 3
            1
                 20
                       D1
## 4
            1
                 23
                       D1
## 5
            1
                 16
                       D1
## 201
            0
                 29
                      D2
## 202
            0
                 25
                       D2
## 203
                 26
            1
                       D2
## 204
            1
                 34
                       D2
## 205
            1
                 21
                       D2
```

The data can be visualized in the following way:

Data for 10 participants



In the graph, the time point 0 represents the moment the participant took the drug for that migraine event, and the horizontal line indicates the period during which the participant's migraine is known to have persisted. The right-hand point where the line ends includes a symbol representing the event observed: x indicates a recovery event, and o indicates a fail event.

Kaplan-Meier curve

One of the most commonly used methods to analyse survival data is the Kaplan-Meier curve. It combines information from an event variable with the time-to-event to yield a probability of survival per time point. Originally called the product limit estimator, it is a result of ideas proposed independently by E. Kaplan and P. Meier for publication by the Journal of the American Statistical Association, which by suggestion of the journal's editor was combined into a single article published in 1958.

Here we will present an intuitive view of how the Kaplan-Meier curve is computed, and how it can be interpreted. More formal introductions are widely available, for example from chapter 12 of Peter Dalgaard's book.

The Kaplan-Meier curve essentially yields an estimate of the probability of survival beyond each time point. By "survival" here we mean a time-to-event longer than the time point considered. A graph is produced using the estimated probabilities for multiple time points.

Such a probability of survival computed for each time point essentially is the proportion of individuals at risk of having an event just before the time point subtracted from the number of events at the time point, relative to all individuals. Note that the proportion of individuals at risk changes only when an event is observed, and the total number of individuals typically stays fixed. So, the Kaplan-Meier probability of survival estimates only changes at the time points where events are observed.

Censored observations (in the sense defined above) belong to the group of all individuals at risk during the entire duration of the study, since we essentially do not know anything about their time-to-event, except that it was larger than a certain time.

Computing the Kaplan-Meier curve does not require any assumptions about the data distribution (event or time-to-event), apart from the facts that event is a grouping variable (as it defines one of a set of known events, per individual) and time-to-event is a non-negative variable (as it represents time). This means it is a non-parametric method, and it makes it widely applicable.

Working example (cont)

We can illustrate this with the subset of the migraine study, where we first consider participants from a single group, thus making use of the same drug:

```
mydata <- migraine[1:10, ] #
mydata1 <- mydata[, -3]
mydata1</pre>
```

```
##
       event time
## 1
           0
                41
                29
## 2
            0
## 3
            1
                20
## 4
            1
                23
## 5
            1
                16
## 6
            1
                15
## 7
            Λ
                23
## 8
            0
                31
## 9
                22
            1
## 10
                28
```

We can easily see how to obtain the estimated probability that the recovery takes longer than a given time, per time point. For any time point prior to the first event occurring, the probability of recovery taking longer than that time point is the number of participants at risk at that time point divided by the total of participants, so it equals 1. Subsequently, for each time point shortly before an event (recovery) is observed, the number of participants at risk decreases by the number of events, which equals 1. We illustrate this by displaying the computed survival probabilities below for the subset of the migraine data:

```
mydatas <- mydata1[order(mydata1$time), ]
nrisk <- nrow(mydatas) - cumsum(mydatas$event)[-c(3, 5)]
prob.surv <- nrisk/rep(nrow(mydatas), length(nrisk))
prob.surv</pre>
```

```
## [1] 0.9 0.8 0.6 0.5 0.4 0.4 0.4 0.4
```

The first two time points correspond to 1 event, but on the third time point 2 events are observed, explaining the larger drop.

Note that, for the time points corresponding to censored time-to-event observations, the probability did not change, as no event was observed. For those, no computation is needed.

Kaplan-Meier curves yield estimates for the survival probabilities. Confidence intervals are often obtained via what is called Greenwood's formula. As the Kaplan-Meier curve, these are nonparametric estimates since they do not rely on any assumption about the data distribution.

Note that estimates of both the Kaplan-Meier curve as well as of the confidence intervals on the right-hand side of the plot typically are associated with high variability. Therefore, care must be taken when interpreting results for the part of the curve with few individuals at risk. More details about this can be found in chapter 12 of Peter Dalgaard's book.

Survival data analysis in R

In R, classic survival data analysis methods are available via the package survival. Let us start by loading it.

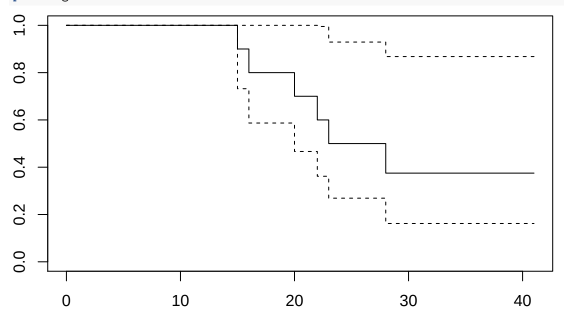
library(survival)

A survival data analysis starts by the creation of the response object, which combines the event variable with the time-to-event. In the migraine study example, these are variables event and time. The response object is created by the function Surv():

```
migr.surv <- Surv(time = mydata1$time, event = mydata1$event)
```

With the response object created, it is straightforward to plot the Kaplan-Meier curve yielded by this object:

```
plot(migr.surv)
```



The computation of the Kaplan-Meier curve is actually done using the function **survfit**. It requires as input a formula object, with the **Surv** object as response and for example a grouping variable as covariate. In this case we will consider no grouping, so we use 1 to represent a model with just the intercept:

```
svfit <- survfit( migr.surv ~ 1)</pre>
```

The summary of survfit yields the computed survival probabilities. In particular, we can now check that the computed probabilities we obtained above are correct:

```
summary( svfit )
```

```
## Call: survfit(formula = migr.surv ~ 1)
##
##
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
                             0.900
                                    0.0949
                                                    0.732
                                                                  1.000
##
      15
              10
                        1
               9
##
      16
                        1
                             0.800
                                    0.1265
                                                    0.587
                                                                  1.000
##
      20
               8
                        1
                             0.700
                                    0.1449
                                                    0.467
                                                                  1.000
               7
      22
                                                    0.362
                                                                  0.995
##
                        1
                             0.600
                                    0.1549
##
      23
               6
                        1
                             0.500
                                    0.1581
                                                    0.269
                                                                  0.929
##
      28
                             0.375 0.1606
                                                    0.162
                                                                  0.868
                        1
```

Quick tasks

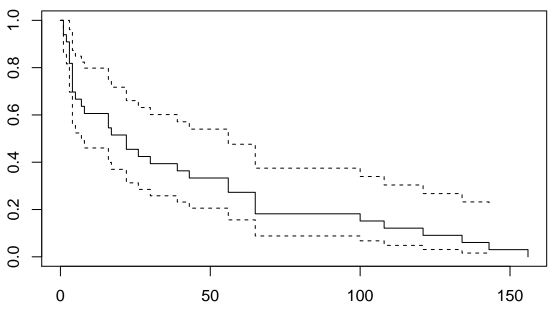
1. The dataset leuk from the library MASS contains time to death for 33 leukemia patients. Consider the event of interest to be death. To explore overall survival times (time to death) after leukemia diagnosis, plot the Kaplan-Meier curve of the data.

```
library(MASS)
names(leuk)
```

```
## [1] "wbc" "ag" "time"

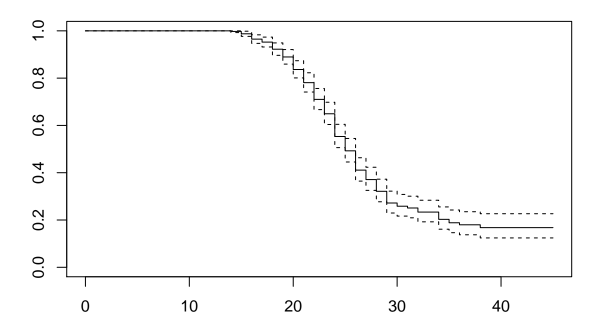
# Note: all patients had the event (death), so there is no censoring
# In such cases, the survival object can be created in two ways: either make a vector
# with one entry per patient, and all entries equal to 1 (indicating an event),
# or else use as input for `Surv()` only the time variable. Then by default the
# function will assume that all patients had an event (see `help(Surv)` for more details)

leuk.surv <- Surv(leuk$time, rep(1, nrow(leuk)))
# Alternative
leuk.surv <- Surv(leuk$time)</pre>
```



2. Plot the Kaplan-Meier curve for the entire migraine data.

```
migr.surv <- Surv(time = migraine$time, event = migraine$event)
plot(migr.surv)</pre>
```



The log-rank test

Typically there is interest in studying the association between a survival response and a covariate, say a grouping variable. This was the case in the migraine working example, where each group made use of a different drug to treat migraine. There is then interest in comparing the survival functions of the two groups.

We can compute the Kaplan-Meier estimators per group, but how do we compare the two obtained curves? This can be done using the log-rank test.

Let us assume that the covariate splits individuals in the study into two groups, and that Kaplan-Meier estimates of the survival distributions were computed per group. The log-rank test helps us test the null hypothesis that two survival distributions are statistically the same, against the alternative hypothesis that they are different. The concept of "statistically the same" seems rather vague. What we mean by it is: under the null hypothesis, differences between the two estimated survival distributions arise at random. This can be seen as comparing the number of observed events/no events per interval (defined by the time between two consecutive events) between the two groups, in what essentially is a 2x2 table:

Frequencies	Group 2, no event	Group 2, event	Total
Group 1, no event	y11	y12	y1.
Group 1, event	y21	y22	y2.
Total	y.1	y.2	N

For any given interval, this comparison can be done by means of a Fisher's exact test (which we saw in chapter 4). The log-rank test results if, per interval, the observed statistic is compared to the expected, across all intervals. The distribution of the resulting test statistic can be approximated by the chi-square with 1 degree of freedom, the same that we used for 2x2 tables when the individual frequencies were large enough.

Note that the log-rank test assumes that the censoring is unrelated to the response variables, so both to event as well as to time-to-event. In fact, many of the survival data analysis methods make this assumption. It is therefore crucial to verify that this assumption holds in any data being analysed. This cannot be done by a statistical test, but you could for example

In R, this can be done by using the **survfit** function. The argument is a formula object, defining the comparison to be performed.

Working example (cont)

Let us compare the obtained survival curves between groups using drugs D1 and D2 in a small subset of the migraine example. First select the subset:

```
mydata <- rbind(migraine[1:10, ], migraine[(ss/2+1):(ss/2+10), ])</pre>
```

Then create the response, the Surv object:

```
migr.surv <- Surv(time = mydata$time, event = mydata$event)
```

Now use survfit to compute the Kaplan-Meier per group. This function uses as input a formula, defining the Surv response and the grouping variable drug indicating the drug used per individual:

```
svfit <- survfit(migr.surv ~ mydata$drug)</pre>
```

The object yielded by survfit is of class survfit. We can print out a summary of the object:

```
print(survfit)
```

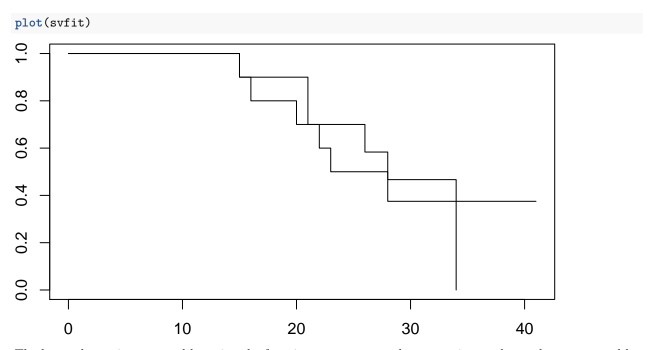
```
## function (formula, ...)
## {
## UseMethod("survfit")
## }
## <bytecode: 0x55fdb473ee38>
## <environment: namespace:survival>
```

and prind the Kaplan-Meier survival probabilities estimated for the two groups:

```
summary(svfit)
```

```
## Call: survfit(formula = migr.surv ~ mydata$drug)
##
##
                    mydata$drug=D1
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
##
      15
             10
                       1
                            0.900 0.0949
                                                   0.732
                                                                 1.000
##
      16
              9
                            0.800 0.1265
                                                   0.587
                                                                 1.000
                       1
##
      20
              8
                       1
                            0.700 0.1449
                                                   0.467
                                                                 1.000
##
              7
      22
                       1
                            0.600 0.1549
                                                   0.362
                                                                 0.995
##
      23
              6
                       1
                            0.500 0.1581
                                                   0.269
                                                                 0.929
##
      28
                            0.375 0.1606
                                                   0.162
                                                                 0.868
                       1
##
##
                    mydata$drug=D2
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
##
             10
                       1
                            0.900 0.0949
                                                   0.732
                                                                 1.000
      15
              9
                       2
                                                                 1.000
##
      21
                            0.700 0.1449
                                                   0.467
##
      26
              6
                       1
                            0.583 0.1610
                                                   0.340
                                                                 1.000
##
      28
              5
                            0.467
                                    0.1658
                                                   0.233
                                                                 0.936
                       1
##
      34
               1
                            0.000
                                       NaN
                                                      NA
                                                                    NA
```

Making a graph of the Kaplan-Meier curves is also straightforward:



The log-rank test is computed by using the function **survdiff** on the comparison to be made, represented by the formula involving the **Surv** object as response, and the grouping variable on the right-hand side:

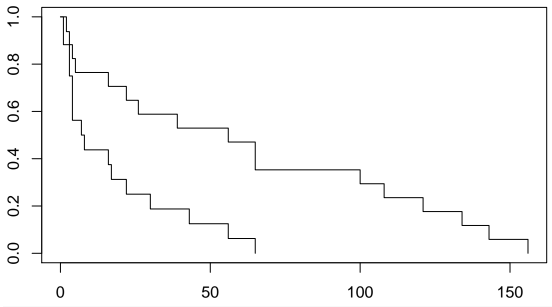
```
survdiff(migr.surv ~ mydata$drug)
## survdiff(formula = migr.surv ~ mydata$drug)
##
                   N Observed Expected (O-E)^2/E (O-E)^2/V
##
## mydata$drug=D1 10
                             6
                                    5.6
                                           0.0292
                                                     0.0572
                            6
                                    6.4
## mydata$drug=D2 10
                                           0.0255
                                                     0.0572
    Chisq= 0.1 on 1 degrees of freedom, p= 0.8
```

Quick tasks

1. Consider again the leukemia data in dataset leuk. We will now compare survival probabilities between the groups given by ag, representing the test result for the presence of Auer rods and/or significant granulation of leukaemic cells.

Plot the Kaplan-Meier survival curves separately per group defined by the AG test. Compute the log-rank test to compare the two survival distributions between the two groups.

```
# Create the Surv object
leuk.surv <- Surv(leuk$time) # no event variable as all patients experience the event (death)
# Define the comparison to be made
leuk.fit <- survfit(leuk.surv ~ leuk$ag)
# Make a plot if the survival distributions for the two groups
plot(leuk.fit)</pre>
```



Test whether the difference is statistically significant using the log-rank test
survdiff(leuk.surv ~ leuk\$ag)

```
survdiff(formula = leuk.surv ~ leuk$ag)
##
##
                     N Observed Expected (O-E)^2/E (O-E)^2/V
## leuk$ag=absent
                    16
                             16
                                      9.3
                                               4.83
                                                          8.45
                             17
                                     23.7
   leuk$ag=present 17
                                               1.90
                                                          8.45
##
##
    Chisq= 8.4 on 1 degrees of freedom, p= 0.004
```

Regression models for survival data

Motivation

We have seen that we can estimate survival probability curves by using the Kaplan-Meier method, and that we can test if they are intrinsically the same using the log-rank test. This test is used for comparing the survival distribution curves between two groups. However, the effect of variables with more groups, or of continuous variables, and that of many variables at the same time, cannot be (directly) tested by means of the log-rank test.

What we want to do is to use a framework like that of regression models, to be able to study the effect on the survival response of multiple variables, and of multiple types of variables, at the same time. Typically regression models only consider one response at the time. So what needs to be done to consider these two variables, in a way so that survival probabilities can be estimated?

The first step is to understand what we want to model. Say that, instead of the survival response, we observe a rate λ_i at which patients die, for samples indexed by $i = 1, \ldots, n$. We will model this using regression as a linear function of a covariate X_i . So we could write

$$\lambda_i = \alpha + \beta X_i, \ i = 1, \dots, n.$$

Since λ_i is a rate, it is always non-negative. The model above could, however, yield estimates that are negative. To avoid this, we use a logarithmic link, as in:

$$\log(\lambda_i) = \alpha + \beta X_i, \ i = 1, \dots, n,$$

or, equivalently,

$$\lambda_i = \exp(\alpha + \beta X_i), \ i = 1, \dots, n.$$

This model arises naturally if patients die at an exponential rate, where the exponential is a probability distribution. Other distributions can be used instead of the exponential, but the basic idea remains to explain the rate at which events are observed by a log-linear function of the covariates. This part of the model represents the contribution of covariates.

A general formulation for modelling survival data is as follows: the hazard h(t) of events at any time t is a product of a baseline hazard $h_0(t)$, common to all observations and varying with t, multiplied by a part that depends only on the covariates and is fixed in time. That is:

$$h_i(t) = h_0(t) \exp(\alpha + \beta X_i), i = 1, \dots, n,$$

where the hazard $h_i(t)$ represents the probability of an event being observed at t for observation i, given that i has not had an event up until t. In this formulation, the baseline hazard function describes how events are generated over time, considering all observations. Then the effect of a covariate just represents a multiplicative effect on the baseline hazard for the entire study period.

Such a formulation is based on a few asumptions:

- the covariate effect is fixed in time;
- there is a common baseline hazard yielding events for all observations, which is independent of the covariates;

The latter implies that

$$h_i(t)/h_j(t) = \frac{h_0(t)\exp(\alpha + \beta X_i)}{h_0(t)\exp(\alpha + \beta X_j)} = \exp(\beta(X_i - X_j)),$$

which means that the hazard ratio for two observations at the same time point t does not depend on t, only depending on their covariates. This means that these observations have proportional hazards.

Most regression models for survival data make use of this assumption, modelling the baseline hazard as the only function of time, multiplied by a log-linear function of the covariates. Classes of models arise depending on how the baseline hazard $h_0(t)$ is modelled.

Parametric models for the baseline hazard include the exponential, as used above, the gamma, the log-normal and the Weibull. These can be fitted with the R function **survreg**. While they may be of interest, in practice non-parametric hazard functions are more commonly used, as they are more flexible. For this reason, we will here focus on survival regression models with non-parametric hazards.

Cox proportional-hazards model

Perhaps the best known approach for estimating a non-parametric baseline hazard function in the context of a proportional-hazards model is the one proposed by D. R. Cox in 1972. The focus of fitting shifts from estimating the baseline hazard to estimating the proportional factors, the coefficients of the covariates in the model.

Two important assumptions of the proportional hazards model remain: the effect of covariates does not vary with time, and the proportionality of hazards.

Working example

To illustrate methods we will use the dataset Melanoma. Check what it contains by reading its help file, and examining the structure of the data.

```
# ?Melanoma
library(survival)
library(MASS)
str(Melanoma)
```

```
205 obs. of 7 variables:
  'data.frame':
##
   $ time
              : int 10 30 35 99 185 204 210 232 232 279 ...
##
   $ status
              : int
                    3 3 2 3 1 1 1 3 1 1 ...
##
              : int 1 1 1 0 1 1 1 0 1 0 ...
   $ sex
##
   $ age
              : int 76 56 41 71 52 28 77 60 49 68 ...
              : int 1972 1968 1977 1968 1965 1971 1972 1974 1968 1971 ...
##
   $ year
                     6.76 0.65 1.34 2.9 12.08 ...
   $ thickness: num
              : int 1000111111...
  $ ulcer
```

In this example, time is the survival time in days, and status is an indicator of the patient's status by the end of the study:

- status = 1: "dead from malignant melanoma";
- status = 2: "alive on January 1, 1978";
- status = 3: dead from other causes"

We will first create a Surv object having only the event dead from melanoma, and both alive as well as dead due to other causes together as censoring. For this, we use:

```
survm <- Surv(Melanoma$time, Melanoma$status == 1)</pre>
```

So, in this case the endpoint variable used in Surv is a logical vector, equal to TRUE if status = 1, and FALSE otherwise.

Now we want to know if sex affects the survival probability of melanoma. To do this, we compute the Cox regression using the Surv object defined above, and sex as a covariate:

```
mel.cox <- coxph(Surv(time, status == 1) ~ sex, data = Melanoma)
summary(mel.cox)</pre>
```

```
## Call:
## coxph(formula = Surv(time, status == 1) ~ sex, data = Melanoma)
##
    n= 205, number of events= 57
##
##
##
         coef exp(coef) se(coef)
                                     z Pr(>|z|)
## sex 0.6622
                 1.9390
                          0.2651 2.498
                                        0.0125 *
##
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
       exp(coef) exp(-coef) lower .95 upper .95
## sex
           1.939
                     0.5157
                                1.153
                                           3.26
##
## Concordance= 0.59 (se = 0.034)
## Likelihood ratio test= 6.15 on 1 df,
                                           p=0.01
## Wald test
                        = 6.24
                                on 1 df,
                                           p=0.01
## Score (logrank) test = 6.47 on 1 df,
                                           p=0.01
```

Interpreting the summary output is similar to that of a fitted regression model: you have a table of coefficients

with estimates, standard errors and a p-value for the test that the coefficient is equal to zero (or not). The end of the summary lists various tests that evaluate the goodness-of-fit of the model, so indicate whether or not the model represents well the data.

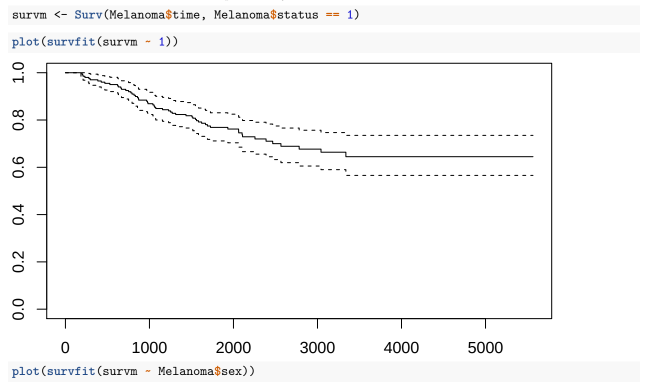
The choice of group of interest will define the sign of the coefficient of sex in the Cox model fit. In this example, the event of interest is death and the group corresponding to sex=0 (females) is taken by the intercept, representing the baseline. Then the coefficient estimated for sex corresponds to the difference in log-hazard for individuals with sex=1 (males) compared to those with sex=0. In this particular case, the computed coefficient is positive, indicating that male patients have a worse prognosis (higher progression probability to death, taking both time and event into account) than females. So, in this example, survival probability actually means progression to death, as the event of interest is death.

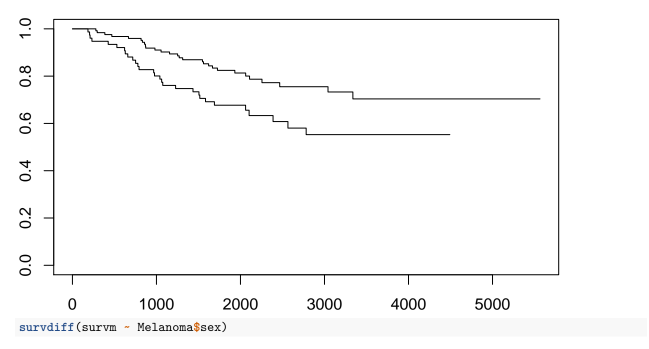
Note also that, if the group coding is inverted, with the second group now being assigned to the intercept, then the estimated coefficient of the grouping variable is the same as the one with the original coding, multiplied by -1.

The summary of the fitted model has one additional part, compared to other regression models: a table of coefficients and confidence intervals on the exponential scale. This is given to make it easier to interpret the coefficients on the scale of the survival probability.

Quick tasks

1. Make Kaplan-Meier curves to the Melanoma data using the same Surv object defined above, i.e. with both censoring categories combined into a single one. Do this both for the entire data together, as well as for the observations according to gender. Compute the log-rank test to check whether there is evidence that the two survival probability curves are different.





```
## Call:
## survdiff(formula = survm ~ Melanoma$sex)
##
##
                    N Observed Expected (0-E)^2/E (0-E)^2/V
                                    37.1
## Melanoma$sex=0 126
                             28
                                              2.25
                                                         6.47
                             29
## Melanoma$sex=1 79
                                    19.9
                                              4.21
                                                         6.47
##
   Chisq= 6.5 on 1 degrees of freedom, p= 0.01
```

\$ time: int 65 156 100 134 16 108 121 4 39 143 ...

2. Let us consider again the leuk example with overall survival data of leukemia cases, of the package MASS.

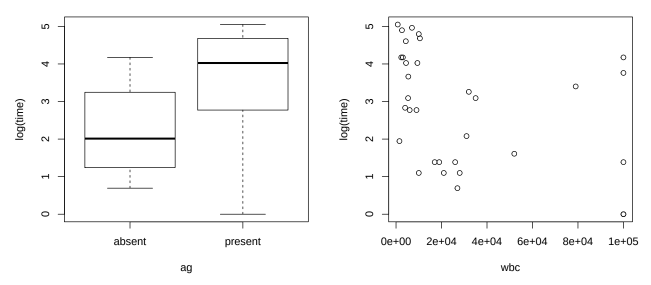
```
library(MASS)
library(survival)
str(leuk)

## 'data.frame': 33 obs. of 3 variables:
## $ wbc : int 2300 750 4300 2600 6000 10500 10000 17000 5400 7000 ...
## $ ag : Factor w/ 2 levels "absent", "present": 2 2 2 2 2 2 2 2 2 2 ...
```

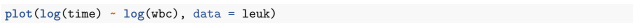
We define again the response as a function of time only, since all patients experience an event (death). Subsequently, we fit a survival regression of the survival response as a function of ag and of white blood cell count wbc. We will use Cox regression for this.

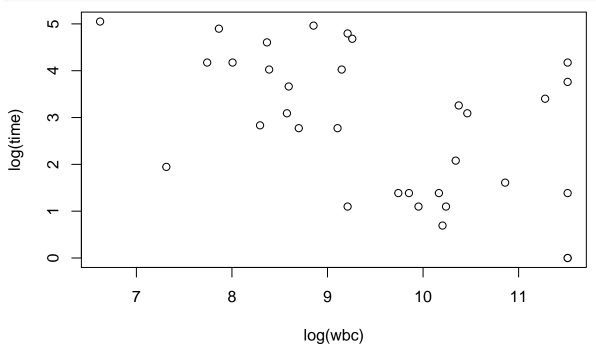
First, let us examine the nature of the relationship between time and the covariates ag and wbc. For this, we make a graph. We use time on the log-scale, as this is equivalent to what happens in the regression model:

```
par(mfrow = c(1, 2))
plot(log(time) ~ ag + wbc, data = leuk)
```



The covariate wbc has many small values and only a few large ones. This is characteristic of a variable on an exponential scale. So we take its logarithm and produce the plot again:





This relationship seems to be closer to linear. So, we will include wbc in the model with a log-transformation.

A Cox regression is fitted via the function coxph which, as regression functions lm and glm, expect a formula as input. The formula relates the Surv response to the covariates, in this case ag and log(wbc):

```
leuk.cox <- coxph(Surv(time) ~ ag + log(wbc), data = leuk)
summary(leuk.cox)

## Call:
## coxph(formula = Surv(time) ~ ag + log(wbc), data = leuk)
##
## n= 33, number of events= 33
##</pre>
```

```
coef exp(coef) se(coef)
                                              z Pr(>|z|)
## agpresent -1.0691
                        0.3433
                                 0.4293 -2.490 0.01276 *
                        1.4444
## log(wbc)
              0.3677
                                 0.1360 2.703 0.00687 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
             exp(coef) exp(-coef) lower .95 upper .95
## agpresent
                0.3433
                           2.9126
                                      0.148
                                                0.7964
## log(wbc)
                1.4444
                           0.6923
                                       1.106
                                                1.8857
##
## Concordance= 0.726 (se = 0.047)
## Likelihood ratio test= 15.64 on 2 df,
                                             p = 4e - 04
                                            p=5e-04
## Wald test
                        = 15.06 on 2 df,
## Score (logrank) test = 16.49 on 2 df,
                                            p = 3e - 04
  3. Consider again the Melanoma data, and now define sex as 1 for females and 0 for males. Fit the Cox
    regression model again and compare results with those when sex was used with the original coding.
sex1 <- 1 - Melanoma$sex
mel.cox1 <- coxph(survm ~ sex1)</pre>
summary(mel.cox1)
## Call:
## coxph(formula = survm ~ sex1)
##
##
    n= 205, number of events= 57
##
##
           coef exp(coef) se(coef)
                                        z Pr(>|z|)
## sex1 -0.6622
                   0.5157 0.2651 -2.498
                                            0.0125 *
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
        exp(coef) exp(-coef) lower .95 upper .95
                                0.3067
                       1.939
## sex1
           0.5157
                                           0.8672
##
## Concordance= 0.59 (se = 0.034)
## Likelihood ratio test= 6.15 on 1 df,
                                            p=0.01
                        = 6.24 on 1 df,
## Wald test
                                            p=0.01
## Score (logrank) test = 6.47 on 1 df,
                                            p=0.01
# With the original coding we had:
summary(mel.cox)
## coxph(formula = Surv(time, status == 1) ~ sex, data = Melanoma)
##
##
     n= 205, number of events= 57
##
         coef exp(coef) se(coef)
                                     z Pr(>|z|)
##
## sex 0.6622
                 1.9390
                          0.2651 2.498
                                        0.0125 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
       exp(coef) exp(-coef) lower .95 upper .95
           1.939
                     0.5157
                                1.153
## sex
```

##

Group-specific baseline hazards

The Cox model is easily extended to allow different baseline hazards for different groups. For example, in the leuk data there could be interest in allowing for different event rates in time, depending on the ag group of the patient - patients termed AG positive display Auer rods and/or significant granulation of their leukemic cells, which may affect the overall event rate. Different hazards according to ag group can be fitted by including the argument strata as ag in the model, as in:

```
leuk.coxs <- coxph(Surv(time) ~ strata(ag) + log(wbc), data = leuk)
summary(leuk.coxs)</pre>
```

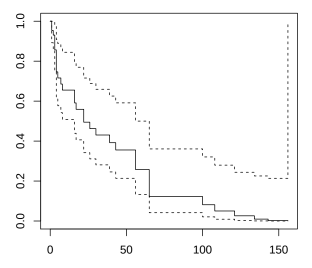
```
## Call:
## coxph(formula = Surv(time) ~ strata(ag) + log(wbc), data = leuk)
##
##
    n= 33, number of events= 33
##
##
             coef exp(coef) se(coef)
                                       z Pr(>|z|)
## log(wbc) 0.3906
                    ## --
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
           exp(coef) exp(-coef) lower .95 upper .95
              1.478
                        0.6767
## log(wbc)
                                  1.117
                                           1.955
##
## Concordance= 0.688 (se = 0.067)
## Likelihood ratio test= 7.78 on 1 df,
                                       p=0.005
                      = 7.5 on 1 df,
                                       0.006
## Wald test
## Score (logrank) test = 7.92 on 1 df,
                                       p=0.005
```

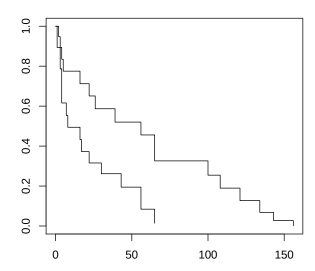
Plots of the estimated survival distributions can be produced by using as input survfit applied to the fitted Cox model:

```
par(mfrow = c(1, 2))
plot(survfit(leuk.cox), main = "AG as covariate")
plot(survfit(leuk.coxs), main = "AG defining strata")
```



AG defining strata





So, from the first model we obtained a single survival distribution, plotted making use of the reference or average value of each covariate. The stratified Cox regression plot yielded two survival distributions, one per stratum (AG group).

Quick tasks

Consider again the Melanoma data, and now fit the Cox proportional-hazards model stratified by sex. Check the model fit.

```
mel.coxs <- coxph(survm ~ strata(Melanoma$sex))</pre>
summary(mel.coxs)
## Call: coxph(formula = survm ~ strata(Melanoma$sex))
##
## Null model
     log likelihood= -241.2087
##
##
     n = 205
# With the original model with a sex effect, we had:
summary(mel.cox)
## Call:
  coxph(formula = Surv(time, status == 1) ~ sex, data = Melanoma)
##
##
##
     n= 205, number of events= 57
##
##
         coef exp(coef) se(coef)
                                      z Pr(>|z|)
  sex 0.6622
                 1.9390
                           0.2651 2.498
                                          0.0125 *
##
##
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
  Signif. codes:
##
##
       exp(coef) exp(-coef) lower .95 upper .95
           1.939
                      0.5157
                                 1.153
                                            3.26
## sex
##
## Concordance= 0.59 (se = 0.034)
## Likelihood ratio test= 6.15
                                 on 1 df,
                                            p=0.01
## Wald test
                         = 6.24 on 1 df,
                                            p=0.01
```

```
## Score (logrank) test = 6.47 on 1 df, p=0.01
```

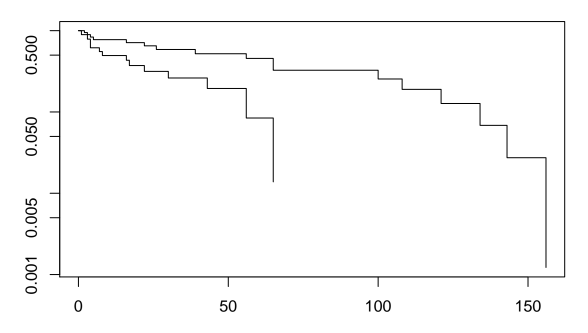
The proportional hazards assumption

By fitting a separate baseline hazard per group, we can compare the group-specific baseline hazards. This helps with checking the proportional-hazards assumption. Above we have made a graph of the separate hazards, on the scale of the survival probability. We can remake this plot on the scale of the log-probability. If the proportional hazards assumption holds, this should yield relatively parallel proportional hazards.

In the leuk example, this can be done as follows:

```
plot(survfit(leuk.coxs), main = "Leukemia data, stratified by AG", log = TRUE)
```

Leukemia data, stratified by AG

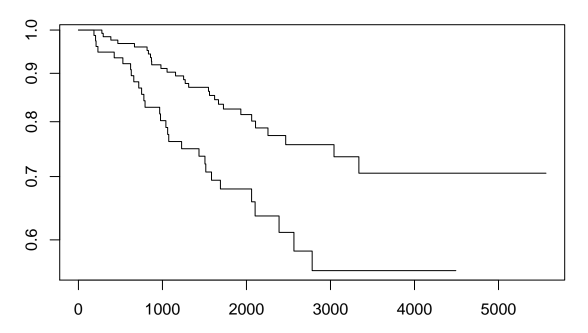


Quick tasks

Plot the survival probability distribution estimated using the stratified model, and decide whether the proportional hazards assumption holds.

```
plot(survfit(mel.coxs), main = "Melanoma data, stratified by sex", log = TRUE)
```

Melanoma data, stratified by sex



Power analysis for survival data

Motivation

Researchers want to start a study to find if progression is related to a chromosomal aberration sometimes found in cancer patients. Here we mean by *progression* the combination of time from end-of-treatment to relapse, as well as the event, which can be relapse or end-of-study, the latter representing censoring. In addition, individuals are recorded as having the aberration or not. From a previous study, progression data and information about the chromosomal aberration are available. Now they want to know how many patients are needed in their new study, designed to test if this chromosomal aberration affects progression or not.

In addition to the chromosomal aberration, other covariates will be taken into account when studying progression, such as age and BMI. For this reason, the researchers decided that they will use a Cox proportional-hazards model to analyse the data and test for the effect of the chromosomal aberration on progression to relapse. The aim of the study is to test whether or not prognosis is the same in the two groups, against the alternative hypothesis that prognosis is different between the two groups.

Since both pilot data are available and the analysis model has been chosen, a power analysis and/or sample size determination is possible.

Power and sample size for Cox regression

As for other types of power analysis and minimum sample size calculations, we can use a formula that relates power, significance level and some statistics from the pilot study to perform calculations. The formula is:

$$N = \frac{[z(1 - \alpha/2) + z(\text{power})]^2}{p(1 - p)\beta_1^2 P\{\text{event}\}}$$

where

• α is the desired significance level;

- the power is equal to 1β , where β is the type-II error;
- $z(\alpha)$ is the quantile function corresponding to the standard normal distribution, yielding the value x that leaves α probability that a value is observed smaller than, or equal to, x, as implemented by qnorm;
- p represents the proportion of cases with the chromosomal aberration in the pilot study or, if known, in this population of patients;
- β_1 represents the log-hazard ratio for the chromosomal aberration effect estimated from a Cox proportional-hazards model for the pilot data;
- $P\{\text{event}\}\$ represents the probability of events in the patient population, in this case estimated by the proportion of relapses (the event of interest);
- N is the sample size.

As before, we can input values for some of these values, and evaluate the remaining ones.

The above formula is built assuming that the test is double-sided, i.e. that the null hypothesis that prognosis does not differ between the two groups is tested against the alternative that the prognosis is different between the two groups. If the alternative is that prognosis is *better* in one group than the other, then the formula to be used is mostly the same, except that $1 - \alpha/2$ is replaced by $1 - \alpha$.

Note that the way the groups are defined (which group corresponds to the intercept) does not matter in this context. This is because the coefficient of the grouping variable β_1 is squared in the model, so its sign does not affect the results.

You do not have to learn this formula by heart. But we will practice using it, so that you better understand what the components mean, and how they can be used when designing experiments.

Working example

Let us consider again the Melanoma data. For these data, the event of interest is status=1, and all other events are considered together as censoring. We define below the Surv object and fit the Cox model, as before:

```
library(survival)
library (MASS)
str(Melanoma)
   'data.frame':
                    205 obs. of 7 variables:
##
                      10 30 35 99 185 204 210 232 232 279 ...
##
               : int
##
                      3 3 2 3 1 1 1 3 1 1 ...
   $ status
               : int
   $ sex
               : int
                      1 1 1 0 1 1 1 0 1 0 ...
##
               : int
                      76 56 41 71 52 28 77 60 49 68 ...
##
                      1972 1968 1977 1968 1965 1971 1972 1974 1968 1971 ...
   $ year
               : int
##
   $ thickness: num
                      6.76 0.65 1.34 2.9 12.08 ...
               : int 1000111111...
   $ ulcer
survm <- Surv(Melanoma$time, Melanoma$status == 1)</pre>
mel.cox <- coxph(Surv(time, status == 1) ~ sex, data = Melanoma)
mel.cox
## Call:
## coxph(formula = Surv(time, status == 1) ~ sex, data = Melanoma)
##
##
         coef exp(coef) se(coef)
                                      z
                          0.2651 2.498 0.0125
## sex 0.6622
                 1.9390
##
## Likelihood ratio test=6.15 on 1 df, p=0.01314
```

```
## n= 205, number of events= 57
```

The coefficient for sex, corresponding to the log-hazard ratio, is statistically significant from 0 at the significance level of $\alpha = 0.05$, but not if we use $\alpha = 0.005$. Let us evaluate what the sample size should be to detect a similar effect size on the log-hazard ratio, if $\alpha = 0.005$ was to be used.

Minimum sample size

Below we write a function to compute the sample size, given the desired power to find a log-hazard ratio from a Cox proportional-hazards model fit. This is written with a binary grouping in mind.

```
# qet.ssize.surv
# Function to compute the sample size required to find a log-hazard ratio using Cox regression,
# where the ratio is computed between two groups
# Inputs
# beta: power, so 1-prob. type-II error
# alpha: desired significance level
# p1: proportion of individuals in group 1 - it does not matter which group is taken as group 1,
      since this enters the formula via p1(1-p1)
# b1: log hazard ratio between the two groups
  It corresponds to the beta coefficient in the cox ph regression
# pevents: prop events
# Output:
# the sample size required
get.ssize.surv <- function(beta, alpha=0.05, p1=0.5, b1=0.5, pevents=1)</pre>
 num <- ( qnorm(1-alpha/2,lower.tail=FALSE) + qnorm(beta, lower.tail=FALSE) ) ^2</pre>
 den <- p1*(1-p1)*(b1^2)*pevents
 n <- num/den
  n
}
```

This function takes as arguments:

- beta: power, so 1-probability of the type-II error. This must be a value between 0 and 1;
- alpha: the desired significance level. This must be a value between 0 and 1;
- p1: proportion of individuals in group 1 it does not matter which group is taken as group 1, since this enters the formula via p1(1-p1). This must be a value between 0 and 1;
- b1: log hazard ratio between the two groups. It corresponds to the beta coefficient computed by the Cox proportional-hazards regression.
- pevents: proportion of events in the data at hand.

Working example (cont)

Let us extract the quantities needed to perform a power calculation. The proportion of events is the proportion of individuals with status=1 in the data:

```
p.event <- mean(Melanoma$status == 1)</pre>
```

The log-hazard ratio estimated by the Cox model is stored in the slot coefficients which, in this case, has a single entry:

```
b1 <- mel.cox$coefficients
```

We also need to compute the proportion of individuals in group 1 (which is arbitrarily chosen). The grouping variable sex has values:

```
table(Melanoma$sex)

##
## 0 1
## 126 79

so its mean is equal to the proportion of individuals in group 1:
p1 <- mean(Melanoma$sex)</pre>
```

Now we can run compute the sample size required, for power = 0.8, say, to detect an effect with a significance level of 0.005.

```
get.ssize.surv(beta = 0.8, alpha = 0.005, p1 = p1, b1 = b1, pevents = p.event)
## sex
## 461.0044
```

Quick tasks

1. Consider again the leukemia data, for which a Cox model was fitted to understand how ag and log(wbc) affect prognosis. Compute the sample size required to find the same effect size for ag as in this data, using a significance level of $\alpha = 0.005$.

```
leuk.cox <- coxph(Surv(time) ~ ag + log(wbc), data = leuk)
# Check the coefficients
leuk.cox$coefficients

## agpresent log(wbc)
## -1.0690504 0.3676989

# We would like to extract the first element of this vector
b1 <- leuk.cox$coefficients[1]
p.event <- 1 # all have an event
p1 <- mean(leuk$ag == "present")
get.ssize.surv(beta = 0.8, alpha = 0.005, p1 = p1, b1 = b1, pevents = p.event)

## agpresent
## 46.63675</pre>
```

Reference

Hsieh, F. Y., and P. W. Lavori. 2000. Sample-size calculations for the Cox proportional hazards regression modelwith nonbinary covariates. Controlled Clinical Trials21: 552–560. DOI: 10.1016/s0197-2456(00)00104-5

Notes

Formulae for power analysis and minimum sample size determination based upon the log-rank test also exist. However, these are applicable to the very specific case where a two-group comparison is made, and no

covariates are used. For this reason, we have decided to present here the solution for the Cox proportional-hazard case only. We should note that the log-rank calculations are of course useful when proportional-hazards cannot be guaranteed, in which case analyses must rely on nonparametric tests such as the log-rank.

Formulae for this sort of problem are also available for more complex designs, such as those from clinical trials. Many of those, as well as the one for the log-rank test, have been implemented in R via the package powerSurvEpi. Their help file is available via the link below:

https://cran.r-project.org/web/packages/powerSurvEpi/powerSurvEpi.pdf

In addition, Stata has many of such formulae implemented. You may wish to check those, or the references in their documentation:

https://www.stata.com/manuals/pss-2powercox.pdf#pss-2powercoxhsiehlavori2000

If using Stata, please make sure you save the scripts used to running analyses!

Other issues

We have just seen an introduction to survival data analysis. It is good to be aware of the things that, in the interest of time, we did not see during this introduction.

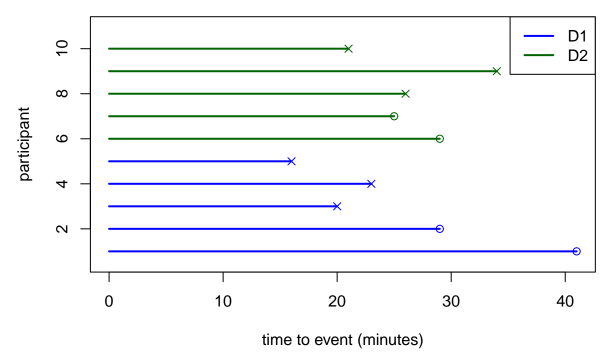
Other types of censoring

Right-, left- and interval censoring

In our examples here we have had censored observations. These were observations from which we knew the starting points, but not the time to event - only that the event did not happen until a certain time point. Recall the graoh:

```
set.seed(39562) # (493756)
ss <- 400
fdrug \leftarrow factor(rep(1:2, each = ss/2), labels = c("D1", "D2"))
event <-c(rbinom(ss/2, 1, prob = .7), rbinom(ss/2, 1, prob = .75))
tte <- round(rgamma(ss, shape = 30, rate = 1+event/4))
migraine <- data.frame(event, time = tte, drug = fdrug)</pre>
mydata <- rbind(migraine[1:5, ], migraine[(ss/2+1):(ss/2+5), ])</pre>
myxlim <- c(0, max(mydata$time))</pre>
mypch <- rep(4, nrow(mydata))</pre>
mypch[ mydata$event == 0 ] <- 1</pre>
plot(1, 1, xlim = myxlim, ylim = c(0.5, nrow(mydata)+1), col = "white",
     main = "Data for 10 participants", xlab = "time to event (minutes)", ylab = "participant")
mcol <- rep(c("blue", "darkgreen"), each = nrow(mydata)/2)</pre>
for(xi in 1:nrow(mydata)) segments(0, xi, mydata$time[xi], xi, col = mcol[xi], lwd = 2)
for(xi in 1:nrow(mydata)) points(mydata$time[xi], xi, col = mcol[xi], pch = mypch[ xi ])
legend("topright", legend = c("D1", "D2"), lty = "solid", lwd = 2, col = c("blue", "darkgreen") )
```

Data for 10 participants



This type of censoring is referred to as *right-censoring*, because in the graph the right-hand side of the follow-up is incomplete.

In other cases, the starting time is unknown. This can be the case when the patient enters the study some time after disease onset, and the exact date of disease onset is unknown. This type of censoring is referred to as *left-censoring*, because in the graph the left-hand side of the follow-up would be incomplete. Of course, there are studies where both types of censoring occur.

In R, Surv can be defined for left-, right- and interval censoring. Typically Cox regression can be used to analyse data with these different censoring types, so long as the censoring is independent of the event probabilities. However, the Kaplan-Meier method cannot be used directly with left-censored data.

Type I and type II censoring

Most progression studies record time to event, and censoring then occurs when either the study is completed before an individual has had an event, or else the individual has been lost to follow-up due to a reason unrelated to the event. In such cases, this is referred to as $type\ I$ censoring. Kaplan-Meier and Cox regression can be used to analyse such data, as we have seen.

In some cases, a trial is stopped after a certain number of events is reached. As individuals who did not experience an event before the end of the trial are censored, these may correspond to those with longer time-to-event observations. So, the censoring process and the event probability are not independent. This is referred to as *type II* censoring. To analyse data from such studies other models are required than the ones seen here.

Competing risks

Let us look again at the Melanoma data:

```
# ?Melanoma
library(survival)
library(MASS)
str(Melanoma)
```

```
'data.frame':
                   205 obs. of 7 variables:
               : int 10 30 35 99 185 204 210 232 232 279 ...
##
   $ time
   $ status
               : int
                     3 3 2 3 1 1 1 3 1 1 ...
                     1 1 1 0 1 1 1 0 1 0 ...
##
   $ sex
               : int
                     76 56 41 71 52 28 77 60 49 68 ...
   $ age
                     1972 1968 1977 1968 1965 1971 1972 1974 1968 1971 ...
##
   $ year
               : int
                     6.76 0.65 1.34 2.9 12.08 ...
   $ thickness: num
               : int 1000111111...
   $ ulcer
```

In this example, time is the survival time in days, and status is an indicator of the patient's status by the end of the study:

- status = 1: "dead from malignant melanoma";
- status = 2: "alive on January 1, 1978";
- status = 3: dead from other causes"

sex 0.6622

1.9390

0.2651 2.498

We previously created a Surv object having only the event dead from melanoma, and both alive as well as dead due to other causes together as censoring. For this, we used:

```
survm <- Surv(Melanoma$time, Melanoma$status == 1)</pre>
plot(survfit(survm ~ Melanoma$sex))
\infty
9
0
       0
                  1000
                              2000
                                           3000
                                                       4000
                                                                    5000
mel.cox <- coxph(Surv(time, status == 1) ~ sex, data = Melanoma)
summary(mel.cox)
## Call:
## coxph(formula = Surv(time, status == 1) ~ sex, data = Melanoma)
##
##
     n= 205, number of events= 57
##
         coef exp(coef) se(coef)
                                      z Pr(>|z|)
```

0.0125 *

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
      exp(coef) exp(-coef) lower .95 upper .95
## sex
          1.939
                    0.5157
                               1.153
                                          3.26
##
## Concordance= 0.59 (se = 0.034)
## Likelihood ratio test= 6.15
                               on 1 df,
                                          p=0.01
## Wald test
                       = 6.24
                               on 1 df.
                                          p=0.01
## Score (logrank) test = 6.47
                               on 1 df,
                                          p=0.01
```

So, in this case the endpoint variable used in Surv is a logical vector, equal to TRUE if status = 1, and FALSE otherwise. However, by considering censoring together with death due to other causes, the probability of survival is biased.

Specifically, the assumption that observations are censored independently of the event probability is violated. This is because individuals who are censored are still at risk, whilst those who died due to other causes are not. As a result, the probability of an event becomes inflated. That is the case with both Kaplan-Meier as well as standard Cox regression

To analyse survival data that includes competing risks, typically people use the Fine and Gray model. There is a large body of research done in this area, so more information can be obtained from various good sources. Here I suggest the following article if you wish to read more about the problem and some solutions:

https://academic.oup.com/ndt/article/28/11/2670/1823847

The Fine and Gray model is available in R via the CRAN package cmprsk:

https://cran.r-project.org/web/packages/cmprsk/cmprsk.pdf

Exercises

Exercise 1

Consider again the leukemia data in dataset leuk. We will now compare survival probabilities between the groups given by ag, representing the test result for the presence of Auer rods and/or significant granulation of leukaemic cells.

- 1.1 Plot the Kaplan-Meier survival curves separately per group defined by the AG test. Compute the log-rank test and add the test result to the graph's title.
- 1.2 By default the curves are plotted with the same colour. Add colours to the plot by using the slot col in the plot call. Note that curves are plotted using the order of the factor levels in the grouping variable, which is also the order used by print and summary in this example D1 comes first, followed by D2. Add a legend to the plot indicating which group corresponds to which colour.
- 1.3 Note that in this case no confidence interval was displayed. Indeed, only when there is a single curve are confidence intervals displayed by default. Display confidence intervals in the plot, using the option conf.int = TRUE.

Colours defined for the groups we now used also for the confidence interval curves.

- 1.4 Use the log-rank test to compare the survival probabilities of the two groups defined by ag. Save the p-value corresponding to the test as an object.
- 1.5 Plot the Kaplan-Meier curves separately per group, and add the p-value to the plot title. Use different colours to display the curves of different groups.
- 1.6 Now fit a Cox regression model to the Surv response using log(wbc) as covariate, and another using both log(wbc) and ag as covariates. Save the fitted models as objects.

- 1.7 The model is fitted using methods similar to those used for generalized linear models. In particular, we can here also use ANOVA to compare models that are nested. Use anova to compare these two model fits. Can you conclude that the model with both log(wbc) and ag as covariates yields a better fit than the model with only log(wbc)?
- 1.8 Let us compare survival probabilities estimated by Kaplan-Meier with those estimated by the Cox model. Make the graph of the Kaplan-Meier curves separately per ag group, as well as of the Cox model's baseline hazards estimated separately, without correcting for log(wbc). How do the survival probabilities estimated by Kaplan-Meier compare with those estimated by Cox regression?
- 1.9 Now repeat this plot, using the Cox model with separate baseline hazards depending on the group defined by ag as before, and correcting for the effect of log(wbc). How do these results compare with those from the previous exercise?

Exercise 2

Read in the migraine data again. Check that the Kaplan-Meier curve produced by using the survfit object with just an intercept is the same as the one produced by plotting the Surv object directly.

Exercise 3

Researchers want to start a study to find if progression is related to a chromosomal aberration sometimes found in cancer patients. Here we mean by *progression* the combination of time from end-of-treatment to relapse, as well as the event, which can be relapse or end-of-study, the latter representing censoring. In addition, individuals are recorded as having the aberration or not. From a previous study, progression data and information about the chromosomal aberration are available. Now they want to know how many patients are needed in their new study, designed to test if this chromosomal aberration affects progression or not.

In addition to the chromosomal aberration, other covariates will be taken into account when studying progression, such as the study. For this reason, the researchers decided that they will use a Cox proportional-hazards model to analyse the data and test for the effect of the chromosomal aberration on progression to relapse.

- 3.1. Read the data in from file data_progression_2groups.txt.
- 3.2. Now explore the data. Check which variables it contains, compute the number of events equal to yes, and the proportion of them.
- 3.3. Create a numeric variable representing the event (=1 if an event ocurred, =0 otherwise). The number of days of progression-free survival is given in the variable pfs. Create a Surv object and make a Kaplan-Meier curve of the data.
- 3.4. Check how many cases there are in each group, and make separate Kaplan-Meier plots for the groups.
- 3.5 Perform a log-rank test to compare the progression data between the groups.
- 3.6 Now fit a Cox proportional-hazards model to estimate the difference in log-hazards between the two groups. Check if there is evidence that the proportional hazards assumption holds.
- 3.7 Now use these results to determine how many samples would be required, in order to have the power of 0.80 to find an effect of the same size of the current effect, with a significance level of 0.01. Do not forget to load the function to estimate the sample size using the available parameters.
- 3.8 Now use these results to determine how many samples would be required, in order to have the power of 0.80 to find an effect of half the size of the current effect, with a significance level of 0.01.
- 3.9 Make a graph of the sample size necessary to find an effect size of values between the current log-hazard ratio and half of its size, using all other parameters the same as before.