

Statistical Analysis with R

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1 Preface

This manual¹ compactly describes some of the principal statistical methods used in social sciences and other disciplines and how to apply them using R. The objective is to facilitate the implementation of statistical methods using the statistical package R and the graphical interface and development platform RStudio. This manual is accompanied by a self-learning module that runs on RStudio. Please let me know about typos or other comments.

¹The latest version of the manual can be found at <https://github.com/rpc01>.

2 Introduction

2.1 Software

R is a complete statistical analysis open source software. The default installation deploys a large number of data management and statistical capabilities, which still can be extended further by installing or developing optional packages to meet more specific needs.

RStudio is an optional IDE (Interface Development Environment) which runs over R and adds to the R original code-based console an attractive and customizable workspace. RStudio is used for many R users.

The self-learning module provides an interactive environment for autonomous learning and review of the topics covered in this course by working directly, hands-on, with RStudio. This learning tool has been developed with swirl, one of the packages of the extensive R library.

2.2 Installation

Please follow the below links:

1. Install R. Just choose the download for your operating system from <https://www.r-project.org> and follow the instructions to install it. The examples of code included in this manual use mostly the core packages and when using additional packages it is indicated including the command `library()`. If the library is not found in the installation you will have to install it using tools - install packages from the top bar of RStudio.
2. Install RStudio. In this course we will use the open source RStudio Desktop free version which can be downloaded from www.rstudio.com.
3. Optional. Install the course companion *Statistical Analysis with R* which runs on RStudio and uses the library swirl. It is a module which facilitates practicing with the contents covered in the manual². Follow the steps in order:
 - (a) Open RStudio and type into the console: `install.packages("swirl")`
 - (b) Start Swirl. This is the only step that you will repeat every time you want to run the self-learning module. First, you will load the package typing: `library("swirl")` then you will start typing: `swirl()`. Press enter after each line.

²At the time of writing only are available self-learning modules for section 4, Descriptive Statistics, and subsection 5.11, Cluster Analysis of this manual.

- (c) Install the last version of the interactive course *Statistical Analysis with R* from its Github repository typing:
`install_course_github(https://github.com/rpc01/statisticalAnalysisWithR)`

2.3 Using RStudio and R Help

Once you launch RStudio it opens with the default workspace consisting of four size-adjustable windows:

- Text Editor. Placed at the top left window is where you will introduce and generate the scripts or programs that want to save to execute later. It is like a regular text editor but suited for introducing code in R language.
- Environment. Placed at the top right window and has several tabs. It is where you will find everything related with the current session: variables, history of commands,
- Console. Placed at the bottom left window is where tasks are executed. It is how the whole screen of R without RStudio would look like. You can introduce code that you want to execute immediately without saving it to a file for later use. It also has tabs where you can follow the execution process, obtain logs and debug errors.
- File Browser, Plots and Help. At the bottom-right window there are three useful tabs.
 - Files shows the Files in your current working directory. The working directory is something you want to control as it is where R will look for external files such as data that you want to use, and also is where it will save the code, plots or results you generate. At section 3.4 you can find how to set the desired working directory.
 - Plot stacks in sequential order the plots that you generate during a session. From there you can choose the plots you want to save, delete or export with the desired format such as .png, .pdf or .jpg.
 - The Help tab is very useful, specially while learning new functions or areas of R. There are two ways to display the contents on the help files in R: (1) typing in the search box of the tab Help or (2) by introducing in the console a command made up of the ? operator and the name of the R function or characteristic we want to know more as shown in the following two lines of code:

```
> ?plot #will open the help for plot() function  
> ?iris #will open the help for iris data object
```

The greater than symbol, `>`, is the default prompt after which commands can be introduced in the R console. The number sign, `#`, introduced in a code line is used for introducing comments which are not executed.

R is an open source software made with the contribution of a large community of researchers and expert users. Apart from the information and software you can find at the official web site repositories <https://cran.r-project.org/web/packages/>, there are also many useful forums, tutorials and courses about R. You just have to do a search in Google and probably will find ideas or support. If you are looking for something very specific it is a good idea to directly copy - paste the message errors or outputs from R.

3 Working with Data

3.1 Reading Raw Data

R admits all types of data such as numbers, text, dates, ... which are stored in different classes of objects such as vectors, data frames and matrices. R is an Objected Oriented Program, OOP such as Java, Python and other widely used programming languages.

3.1.1 Manual Input of Data

The simplest way to introduce data in R is to use the console. For example, we can introduce the number *1* in a vector we call *first*. Vectors are the most elemental class of R objects.

```
> first <-1 # nothing printed
> first      # printed

[1] 1
```

The `<-` symbol is the assignment operator . The first line creates a new object we have named "first" which contains just one item, the number 1. The second line of code prints the object on the screen.

It is also possible to manually introduce data in a faster way by using R operators such as :

```
> vector <-10:20
> vector

[1] 10 11 12 13 14 15 16 17 18 19 20
```

or functions such as `rep()`:

```
> fiveTwos<-rep(2,5)
> fiveTwos

[1] 2 2 2 2 2
```

...or `seq()`:

```
> by2<-seq(5.5,19.2,2)
> by2

[1] 5.5 7.5 9.5 11.5 13.5 15.5 17.5
```

R functions take the form *nameOfFunction(x)*, where x is the argument or arguments that the function can take.

The objects we have created so far only contain numeric data. We can have also objects which contain more than one type of data with the function `c()`:


```
> car <-c("Ford",4,"red",FALSE,"as new",NA)
> car

[1] "Ford"    "4"       "red"     "FALSE"   "as new" NA
```

The vector with name `car` contains 6 elements of the following types: numerical(4), text (Ford, red, as new), logical (FALSE), and a peculiar type of data, missing values (NA). Notice that when printing the vector `car` on the screen, the elements appear around of quotation marks `" "`. This is due to the fact that the class of object vector can contain only one type of data, and when more than one type of data is used, it converts (coerces) everything to text which appears between "quotation marks".

If we want to store different types of data in the same object, we can use other types of object such as data frames. Data frames are excellent for storing data organized in columns and rows or tabular data. Let's create a data frame called `myCars` with 2 columns and 6 rows. We will assign the names attribute and car to each column:

```
> myCars <-data.frame(
+   attributte = c("Name","cilynders","colour",
+                 "sport car", "observation",
+                 "price"),
+   car=c("Ford",4,"red",FALSE,"as new", NA)
+ )
> myCars
```

Notice that as the data entered in the column `car` is the same as the former vector `car`, we could have saved work by doing:

```
> myCars <-data.frame(
+   attributte =c("Name","cilynders","colour",
+                 "sport car", "observation","price"),car)
> myCars
```

	attributte	car
1	Name	Ford
2	cilynders	4
3	colour	red
4	sport car	FALSE
5	observation	as new
6	price	<NA>

Notice that object `"car"` that we call as an argument of the function `c()` does not take quotation marks in contrast to `"Name"`, `"cilynders"` The objects which are resident (that means that have been created or loaded) in the current session appear in the top right window of RStudio (Environment) and are called without using quotation marks.

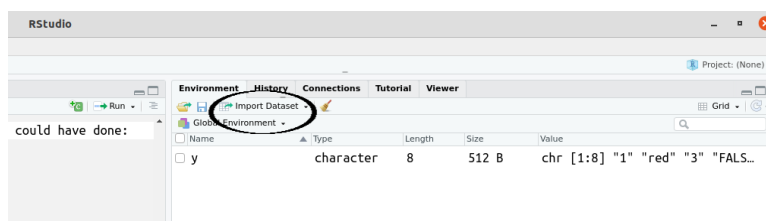
3.1.2 Import Data Files

Fortunately, in many cases data is not introduced to objects by hand. Often data is imported from other files such as spreadsheets, comma separated files, SQL databases, or configuring connections to other sources such as web scrapping services.

To import data from a file we can use the function `read.csv()`, `read.xls()` or `read.table()` depending on the file type.

```
> myxlsdata<-read.xls("myspreadsheet.xls")
> mytxtdata<-read.table("mytextfile.txt")
```

Another possibility is to take advantage of the RStudio graphical interface and click on the Import data set Tab in the top right environment window and follow the required steps.



3.2 Handling and Transformation of Data Sets

To perform statistical analysis, data should be distributed in rows and columns in what is known as tidy data, and has the following characteristics :

1. Each column contains only data from a single variable.

untidy				tidy			
name	sex-age	country		name	sex	age	country
Paul	m-24	USA		Paul	m	24	USA
John	m-22	UK		John	m	22	UK
Melinda	f-27	UK		Melinda	f	27	UK

2. Each variable contains only one type of data.

untidy				tidy			
name	sex	age	country	name	sex	age	country
Paul	m	24	USA	Paul	m	24	USA
John	m	less than 25	UK	Melinda	f	27	UK
Melinda	f	27	UK				

3. Each row contains only data from the same instance, a single observation.

untidy		tidy	
male	female	name	sex
Paul - John	Melinda	Paul	m
		John	m
		Melinda	f

4. Each data set (table) pertains to the same observational unit.

untidy				
name	sex	age	country	unemployment rate
Paul	m	24	USA	4%
John	m	22	UK	8%
Melinda	f	27	UK	8%

tidy			
country	unemployment rate	name	sex
USA	4%	Paul	m
UK	8%	John	m
		Melinda	f

In some cases we will clean and prepare the data in the original file using other programs such as spreadsheets which might be very handy to prepare not very large data sets using pivot and dynamic tables. Once we import tidy data to R it is ready to use. In other cases however we will prefer or will have to do all these transforming operations in R.

3.2.1 First Exploration of the data set

Once we have loaded either a tidy or an untidy data set in R we will be interested on doing a first and fast exploration.

As an example we will use the iris data set. which is one of the data sets included in the default installation of R, ready to use without having to worry to load data from anywhere.

```
> class(iris) #see the object class
```

```
[1] "data.frame"
```

```
> dim(iris) #see dimensions: rows and cols
```

```
[1] 150 5
```

```
> head(iris) #print just the top lines
```

	Sepal.Length	Sepal.Width	Petal.Length	Petal.Width	Species
1	5.1	3.5	1.4	0.2	setosa
2	4.9	3.0	1.4	0.2	setosa
3	4.7	3.2	1.3	0.2	setosa
4	4.6	3.1	1.5	0.2	setosa

5	5.0	3.6	1.4	0.2	setosa
6	5.4	3.9	1.7	0.4	setosa

Instead of scrolling through the set in R we will explore the set using informative functions of our choice such as `class()`, `dim()` or `head()`. We see that `iris` is an object of the class data frame, which has 150 rows and 5 columns, and with the `head()` function we print on the screen a few initial rows of the set. These commands are very useful to get a glance of the data at hand. If the number of columns and rows is reasonably small, R also has the function `View()` to scroll through the data and which is covered at section 4.1.

3.2.2 Selecting elements, rows and/or columns

Often we will like to select just some elements, columns or rows from a data set. We can do it using the brackets operator `[,]` with indexes for the rows and columns as shown in the following examples:

```
> v<-c("blue","grey","orange") #creates vector v
> v[2] #selects the second element of v

[1] "grey"

> iris[2,1] # the element of the second row and first column

[1] 4.9

> iris[1:3,] # the first three rows (and all the columns)

  Sepal.Length Sepal.Width Petal.Length Petal.Width Species
1          5.1          3.5          1.4          0.2  setosa
2          4.9          3.0          1.4          0.2  setosa
3          4.7          3.2          1.3          0.2  setosa

> iris[1:3,4:5] # first three rows and last two columns

  Petal.Width Species
1          0.2  setosa
2          0.2  setosa
3          0.2  setosa

> tail(iris[,4:5]) # last two columns and print just ending rows

  Petal.Width Species
145          2.5 virginica
146          2.3 virginica
147          1.9 virginica
148          2.0 virginica
149          2.3 virginica
150          1.8 virginica
```

With the last line of code we only print the last few of the 150 rows of the last subset using the function *tail()* on columns 4 and 5.

We can also use the minus sign to exclude elements from the data set. Look at the example:

```
> s1<-iris[2:4,-c(1,5)] #excludes column 1 and 5
> s1
```

	Sepal.Width	Petal.Length	Petal.Width
2	3.0	1.4	0.2
3	3.2	1.3	0.2
4	3.1	1.5	0.2

We could obtain the same with:

```
> iris[-c(1,5:150),c(2:4)] # or
```

	Sepal.Width	Petal.Length	Petal.Width
2	3.0	1.4	0.2
3	3.2	1.3	0.2
4	3.1	1.5	0.2

```
> s2<-iris[-c(1,5:nrow(iris)),c(2:4)] #nrow() gives the number of rows of the set
> s2
```

	Sepal.Width	Petal.Length	Petal.Width
2	3.0	1.4	0.2
3	3.2	1.3	0.2
4	3.1	1.5	0.2

We are using functions as arguments of other functions, which results in a compact code syntax.

To check that the objects *s1* and *s2* are equal we can use the *identical()* function:

```
> identical(s1,s2)
```

```
[1] TRUE
```

To select columns we can use the dollar operator, *\$*, which takes column names instead of indexes as in the following example:

```
> iris$Sepal.Length #just the column of lengths of Sepals.
> iris$Species #just the column of Species.
```

We can select rows which meet certain criteria by using the function *which()* as in the following example:

```
> s3<-iris[which(iris$Species=="setosa"),c(2)] # col 2 of plants"setosa"
> head(s3) #just print the first lines
```

```
[1] 3.5 3.0 3.2 3.1 3.6 3.9
```

Notice that "=" and "==" operators have very different meanings: "=" is used to assign while "==" checks whether the two elements are equal.

```
> a=3 #creates object a
> b=4 #creates object b
> c=3 #creates object c
> a==b #checks if a is equal to b
```

```
[1] FALSE
```

```
> a==c #checks if a is equal to c
```

```
[1] TRUE
```

Apart than using the minus sign we can also do selections by exclusion with the exclamation mark "!". And we can also concatenate conditions with the ampersand "&".

```
> iris[which(!iris$Species=="setosa"&iris$Petal.Length>6.5),]

   Sepal.Length Sepal.Width Petal.Length Petal.Width Species
106          7.6         3.0         6.6         2.1 virginica
118          7.7         3.8         6.7         2.2 virginica
119          7.7         2.6         6.9         2.3 virginica
123          7.7         2.8         6.7         2.0 virginica
```

We have selected the instances (rows) of species different from setosa which have petals longer than 6.5.

If we want to save any of the subsets or selections for later use, we have to assign them to a new object just by giving it a name:

```
> firstThreeRows<-iris[1:3,]
> firstThreeRows # print
```

```
   Sepal.Length Sepal.Width Petal.Length Petal.Width Species
1          5.1         3.5         1.4         0.2  setosa
2          4.9         3.0         1.4         0.2  setosa
3          4.7         3.2         1.3         0.2  setosa
```

```
> firstThreeRows[3,3] # the element length of petal of the third observation
```

```
[1] 1.3
```

If the object with the chosen name already exists, it will be replaced with the new assigned content without displaying any warning.

3.2.3 Merging data sets

In some cases we will want to create data sets by merging other data sets which have the same length and order. Equal length refers to equal number of rows, and equal order refers to the order of rows.

With the following code we create two simple vectors, `names` and `ages`, which have the same length (3) and order (Paul, John and Melinda):

```
> names <- c("Paul", "John", "Melinda")
> ages <- c(24, 22, 27)
```

We can create a 2 rows and 3 columns matrix with the function `rbind()`, or a 3 rows and 2 columns matrix with the function `cbind()`:

```
> threecol<-rbind(names,ages)
> threecol

      [,1] [,2] [,3]
names "Paul" "John" "Melinda"
ages  "24"  "22"  "27"
```

```
> twocol<-cbind(names,ages)
> twocol

      names  ages
[1,] "Paul"  "24"
[2,] "John"  "22"
[3,] "Melinda" "27"
```

The matrix object class as well as the vector class do only allow elements of the same type, and coerces numeric elements such as age, to text. Notice that text is printed between quoted marks. To mix different types of elements in the same object we can use objects of the class data frame as seen in section 3.1.1. We create the data frame with the function `data.frame()` as in the following example:

```
> namesAges<-data.frame(names,ages)
> namesAges

      names  ages
1    Paul   24
2   John   22
3 Melinda   27
```

We can enlarge the last data set by adding a data frame which contains the height (cm) and weight (kg) (in the same order Paul, John and Melinda):

```
> heightsWeights<-data.frame(heights=c(170,167,177),
+                               weights=c(70,67,77))
```

```
> cbind(namesAges,heightsWeights)
```

	names	ages	heights	weights
1	Paul	24	170	70
2	John	22	167	67
3	Melinda	27	177	77

Many times however, we will want to merge data from sets with different lengths and/or order of observations. In that case we will do the merge using an index variable, a variable which appears in both sets and unequivocally allows to connect matching instances of both sets. R does this with the *merge()* function. In the following example the data frame *heightsWeights2* contains the observation Tim, which is not included in the data frame *namesAges*, but we will overcome this difference by assigning the index variable *names* which appears in both sets.

```
> namesAges
```

	names	ages
1	Paul	24
2	John	22
3	Melinda	27

```
> heightsWeights2<-data.frame(names=c("Paul","John","Melinda","Tim"),
+                               heights=c(160,167,177,168),
+                               weights=c(70,67,102,68))
```

```
> heightsWeights2
```

	names	heights	weights
1	Paul	160	70
2	John	167	67
3	Melinda	177	102
4	Tim	168	68

```
> merge(namesAges,heightsWeights2,by="names")
```

	names	ages	heights	weights
1	John	22	167	67
2	Melinda	27	177	102
3	Paul	24	160	70

The *merge()* function with optional parameters *by*, and *all*, *all.x* and *all.y* allows to perform the inner, outer, left, right and cross join clauses used with SQL language.

```
> merge(namesAges,heightsWeights2,by="names",all=FALSE) #inner join
```

	names	ages	heights	weights
1	John	22	167	67


```

2 Melinda  27    177    102
3   Paul   24    160     70

> merge(namesAges,heightsWeights2,by="names",all=TRUE) #outer join

  names ages heights weights
1   John  22    167     67
2 Melinda 27    177    102
3   Paul  24    160     70
4    Tim  NA    168     68

> merge(namesAges,heightsWeights2,by="names",all.x=TRUE) #left join

  names ages heights weights
1   John  22    167     67
2 Melinda 27    177    102
3   Paul  24    160     70

> merge(namesAges,heightsWeights2,by="names",all.y=TRUE) #right join

  names ages heights weights
1   John  22    167     67
2 Melinda 27    177    102
3   Paul  24    160     70
4    Tim  NA    168     68

> merge(namesAges,heightsWeights2,by=NULL) #cross join

  names.x ages names.y heights weights
1    Paul  24    Paul    160     70
2    John  22    Paul    160     70
3 Melinda 27    Paul    160     70
4    Paul  24    John    167     67
5    John  22    John    167     67
6 Melinda 27    John    167     67
7    Paul  24 Melinda    177    102
8    John  22 Melinda    177    102
9 Melinda 27 Melinda    177    102
10   Paul  24     Tim    168     68
11   John  22     Tim    168     68
12 Melinda 27     Tim    168     68

```

3.2.4 Add calculated data to a data frame

In the last two subsections we have covered how to add existing data to a data frame. In some cases we might want to add data which is obtained in the original data set.

Let's add a calculated column (variable) Body Mass Index (BMI) which is the weight in kg divided by the height in meters squared. We round it to 0 decimals with *round()*.

```
> mydf<-merge(namesAges,heightsWeights2,by="names",all=FALSE)
> bmi<-round(mydf$weights/(mydf$heights/100)^2,0)
> mydf<-cbind(mydf,bmi)
> mydf
```

	names	ages	heights	weights	bmi
1	John	22	167	67	24
2	Melinda	27	177	102	33
3	Paul	24	160	70	27

We can also add a calculated row with the average of the numeric variables. We use the function *colMeans()* selecting only numeric columns:

```
> avg<-colMeans(mydf[,2:5])
> round(avg,2) #round at 2 decimals
```

	ages	heights	weights	bmi
24.33	168.00	79.67	28.00	

We can not add the row avg to mydf using *rbind()* as does not contains the column name. We have to add the missing column:

```
> avg<-colMeans(mydf[,2:5])
> newAvg<-mydf[1,]
> newAvg[1]<-"AVERAGE"
> newAvg[2:5]<-round(avg,2)
> newAvg
```

	names	ages	heights	weights	bmi
1	AVERAGE	24.33	168	79.67	28

```
> rbind(mydf,newAvg)
```

	names	ages	heights	weights	bmi
1	John	22.00	167	67.00	24
2	Melinda	27.00	177	102.00	33
3	Paul	24.00	160	70.00	27
4	AVERAGE	24.33	168	79.67	28

We can also add new variables using conditions . For example we can add the label overweight which takes the logical label TRUE if BMI is greater or equal to 30 and FALSE otherwise.

```
> mydf
```

```

      names ages heights weights bmi
1     John   22     167      67  24
2 Melinda   27     177     102  33
3     Paul   24     160      70  27

> overweight <- mydf$bmi>=30
> cbind(mydf,overweight)

      names ages heights weights bmi overweight
1     John   22     167      67  24      FALSE
2 Melinda   27     177     102  33      TRUE
3     Paul   24     160      70  27      FALSE

```

or instead of using the BMI we can invent the condition: TRUE if weight is higher or equal than the mean AND height is lower than the mean:

```

> overweight <- mydf$weights>=mean(mydf$weights)&
+   mydf$heights<mean(mydf$heights)
> cbind(mydf,overweight)

      names ages heights weights bmi overweight
1     John   22     167      67  24      FALSE
2 Melinda   27     177     102  33      FALSE
3     Paul   24     160      70  27      FALSE

```

3.2.5 Dealing with missing values

Raw data might contain missing values for some or all of the variables (columns) of one or more instances (rows). The treatment of missing values is something that the analyst has to decide based on the available data and the type of analysis that has to perform. However, when possible a simple solution might be just to delete the instance that contains missing data. This can be done with the function *na.omit()* as shown in the following example:

```

> myCars

      attributte    car
1      Name      Ford
2 cilynders      4
3    colour    red
4 sport car  FALSE
5 observation as new
6      price    <NA>

> myCars<-na.omit(myCars)
> myCars

```

```

      attributte   car
1      Name      Ford
2  cilynders      4
3      colour    red
4  sport car  FALSE
5 observation as new

```

After applying the `na.omit()` function and assigning the output to the same `myCars` object, the missing value is removed.

3.3 Save and export Data, Results and Graphics

3.3.1 Save all the elements of a session

When ending a session R saves by default all the elements (environment) to a file with extension `.Rdata` in the working directory. When starting a session it reopens the last session loading the `.Rdata` file.

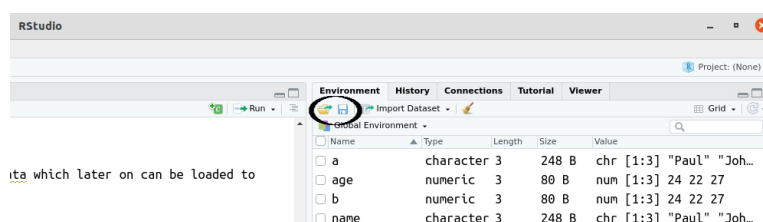
It is possible to choose a specific name for a session, and save only some of the objects. The following example saves objects `x`, `y` and `z` in a file named `mydata.Rdata`.

```
> save(x, y, z, file = "mydata.Rdata")
```

Later on, it is possible to load an specific `.Rdata` file and re-start the session at the point we ended.

```
> load("mydata.Rdata")
```

We could use also use the tabs save and load workspace in the top-right environment window of RStudio, to perform the above tasks using the graphical interface.



3.3.2 Saving Data Objects to csv files

Results, intermediate generated data or data sets can be exported to comma separated values, `csv`, files that be opened with other software such as spreadsheets. To export to `csv` files we can use the console with the function `write.csv`.

For instance, we can export a summary of descriptive statistics or a data set as in the following example:

```
> write.csv(pt_data, file = "pt_data.csv")
```

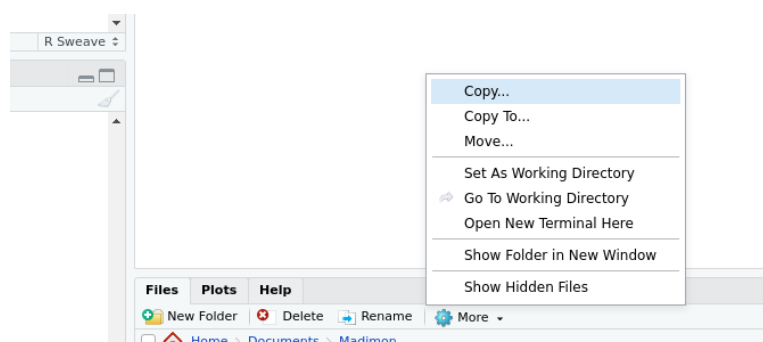
The package *writexl* has the function *write.xls()* which can be used to save content directly with spreadsheet format. If the function is not available probably have to install the package as described in 2.2.

3.4 Setting the Working Directory

When starting a project we will probably want to have raw data files, scripts (code), results, images, and files that R will automatically generate in a separate folder, or folder tree. This folder is the working directory. To know the path of the current working directory we can use the console:

```
> getwd()
> "/user"
```

If we want to change the working directory we can use either the function *setwd()* in the console or the graphical interface in the bottom right window of RStudio to set a desired working directory:



```
> setwd("/user/myproject")
```

The working directory for the session, after running the above line of code, will be `/user/myproject`, and it is that place where RStudio will point for loading and saving data files, export files and images, create intermediate files ..., unless we provide a different path.

4 Descriptive Statistics

4.1 Viewing Data Objects

Apart from the exploratory functions presented in section 3.2.1, R provides the `View()` function which opens a window with the data set and allows scrolling through rows and columns as in a spreadsheet.

The code of the following example opens a new tab in the top-left window of RStudio, where we can explore the iris data set, sort by columns, filter,

...

```
> View(iris)
```

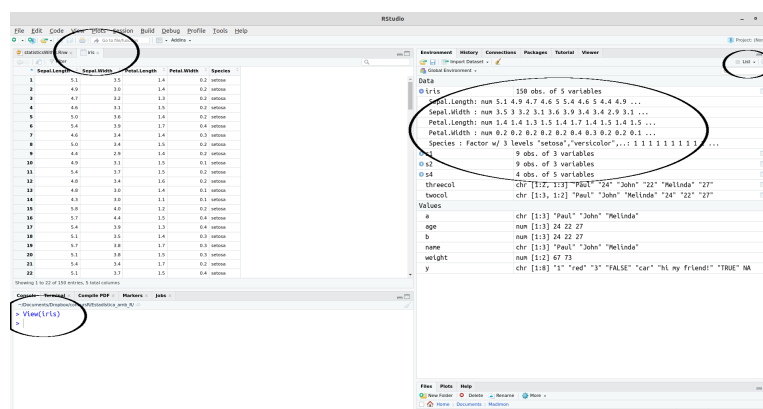


Figure 1: Viewing a data set

As shown in figure 1, we can display the view of the data set with the `view()` function as well as by clicking on its name in the environment objects list.

4.2 Frequency Tables

Calculating frequencies is about counting occurrences.

Following there are several examples with the data set `absenteeism` of the R package `openintro` which contains school absenteeism data from New South Wales. We call the data set `absenteeism` without installing the full `openintro` package by using the operator `::`.

```
> absenteeism<-openintro::absenteeism #creates data object
```

```
> head(absenteeism) #head() prints a few lines
```

```
# A tibble: 6 x 5
  eth  sex  age  lrn  days
<fct> <fct> <fct> <fct> <int>
1 A    M    F0   SL    2
```

```

2 A      M      FO      SL      11
3 A      M      FO      SL      14
4 A      M      FO      AL      5
5 A      M      FO      AL      5
6 A      M      FO      AL      13

```

```

> names(absenteeism) #variables/cols names
[1] "eth" "sex" "age" "lrn" "days"
> dim(absenteeism) #dimensions rows and cols
[1] 146 5

```

We see that we have an object of the class tibble, similar to a data frame, which has 146 rows (observations) with 5 variables each: ethnicity (Aboriginal or Not), gender (M, F), age bucket, learner status (Average Learner or Slow Learner) and days (number of days absent). The four first variables are categorical and the last one (days) numerical.

To count the elements of each type in a vector of the data frame we will use the function `table()`.

```

> names(absenteeism) #to see col names
[1] "eth" "sex" "age" "lrn" "days"
> table(absenteeism$sex) #to count occurrences of each sex
 F  M
80 66
> table(absenteeism$lrn) #to count occurrences of each learner status
AL SL
83 63

```

If we count for more than a variable at the same time we will create a contingency table:

```

> table(absenteeism$sex, absenteeism$lrn)
      AL SL
F  40  40
M  43  23

```

We can turn frequency tables into proportions tables with `prop.table()`

```

> mytable<-table(absenteeism$sex, absenteeism$lrn)
> mytable<-prop.table(mytable)
> round(mytable*100,1) #in % rounded to 1 dec

```

	AL	SL
F	27.4	27.4
M	29.5	15.8

We can add marginals sums to frequency tables with *addmargins()*

```
> mytable<-table(absenteeism$sex, absenteeism$lrn)
> addmargins(mytable)
```

	AL	SL	Sum
F	40	40	80
M	43	23	66
Sum	83	63	146

At section 5.3 we cover how to use contingency tables as an statistical test for categorical variables.

4.3 Descriptive Measures

Numerical descriptive measures together with graphics form the starting point of most statistical analysis.

4.3.1 The *summary* function

The function *summary()* provides some of the common descriptive measures of a data set.

```
> summary(absenteeism)
```

eth	sex	age	lrn	days
A:69	F:80	F0:27	AL:83	Min. : 0.00
N:77	M:66	F1:46	SL:63	1st Qu.: 5.00
		F2:40		Median :11.00
		F3:33		Mean :16.46
				3rd Qu.:22.75
				Max. :81.00

For the categorical variable *setosa* it calculates frequencies, and for continuous variables calculates Min, Max, quartiles and median; simple measurements but a good starting point’.

4.3.2 Selecting Descriptive Statistics

We can use functions to calculate specific descriptive statistical measures not included in the *summary* function. We can also focus our analysis to only some of the variables of the data set. Some of the useful descriptive statistic measures loaded in the core R libraries are *mean()*, *median()*, *sd()*,

max(), *min()*, *range()*, or *mad()* ³. These functions are often applied to some of the variables and/or specific subsets of data.

When we want to analyze separate columns of the data frame it is useful to use the function *attach()*, which makes available for use the variables of a data frame without indicating the data frame to which pertain:

```
> attach(absenteeism)
```

After attaching the set we can call the variable without its data set:

```
> mean(days) #instead of mean(absenteeism$days)
```

```
[1] 16.4589
```

```
> sum(days)
```

```
[1] 2403
```

When finished working with the attached variables is a good practice to detach them back:

```
> detach(absenteeism)
```

If we want to compute a descriptive statistic of a variable but we want to split it in categories we can use the function *tapply()*. The *tapply()* function takes at least three arguments: the first one is the column where we want to apply the function, the second the categories we want to use, and the third the function we want to apply. We can calculate again the mean of absenteeism days that for the whole set we have calculated is 16.5, but we will split it by sex, that is we will have the mean of absent days for men and mean of absent days for women.

```
> tapply(days,list(sex),mean)
```

```
      F      M
15.22500 17.95455
```

If instead of the mean we want to sum the days and instead of split by sex we want to split by sex and ethnicity we can use:

```
> tapply(days,list(sex,eth),sum)
```

```
      A      N
F 795 423
M 670 515
```

We can save any of the tables we have created to a data frame which we can use later or export to a file:

³Many other statistics can be found in specific packages such as (*pastecs*).

```
> meanDaysbySex<-tapply(days,list(sex),mean)
> write.csv(meanDaysbySex,file="daysbySex.csv")
```

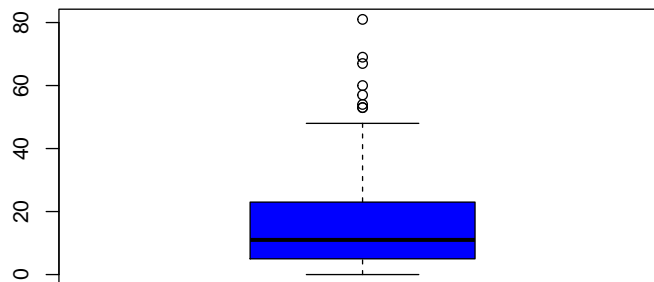
4.4 Descriptive Graphics and Charts

The core installation of R provides many plotting functions and there are also dedicated libraries for advanced approaches to plotting⁴. We will use here the core plotting functions without installing any of the additional packages.

4.4.1 Plots of one Variable: Boxplots and Histograms.

Boxplots are a visual representation of the descriptive measures obtained with the *summary* function, quartiles, min, median,, max ..., plus a bit more information. This is implemented via the *boxplot()* function.

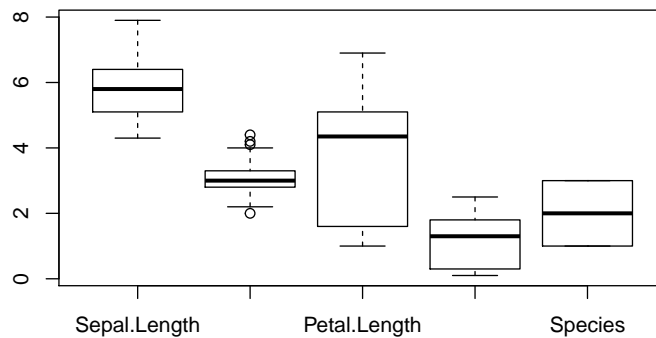
```
> boxplot(absenteeism$days,col="blue")
```



The blue region shows the interquartile range with the median (dark line). The circle dots outside the min and max lines identify outliers. We can also plot several variables side-to-side which can be useful to compare them. With the variables of the data set *iris*:

```
> boxplot(iris)
```

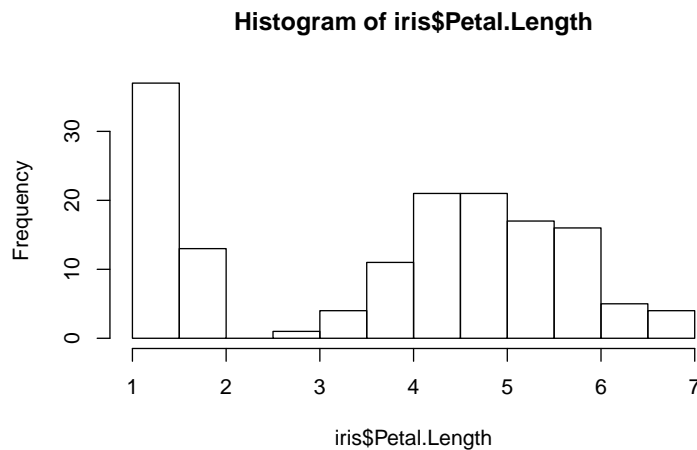
⁴ggplot2 is a very attractive plot library



As you see it is a very good way to compare visually continuous variables. The boxplot of the categorical variable Species does not make much sense; boxplots are not used for categorical variables.

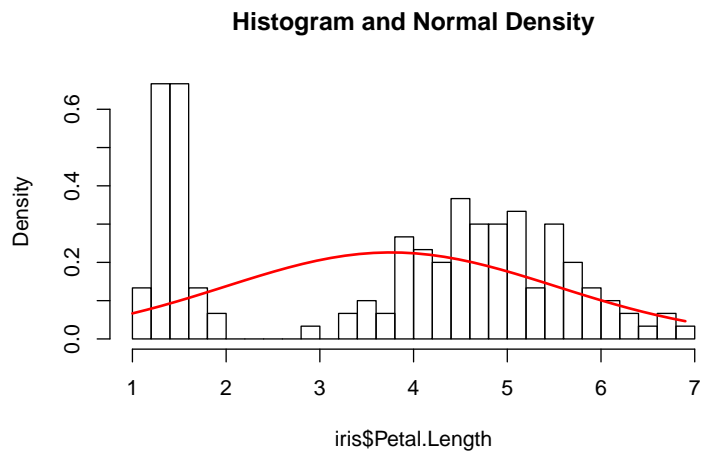
Histograms visualize the distribution of frequencies of quantitative variables, and are very useful to learn about the underlying probability distribution function of the variable. In R histograms are created with the `hist()` function:

```
> hist(iris$Petal.Length)
```



The `hist()` function can take options for customizing different aspects such as the size of bins with `breaks=`, use the density probability instead of the absolute frequency in the vertical axis with `prob=TRUE` and add on top the normal density calculated with the mean and standard deviation of the variable, as shown in the following plot:

```
> hist(iris$Petal.Length,breaks=40,prob=TRUE, main="Histogram and Normal Density")
> x <- seq(min(iris$Petal.Length), max(iris$Petal.Length), length = 40)
> f <- dnorm(x, mean = mean(iris$Petal.Length), sd = sd(iris$Petal.Length))
> lines(x, f, col = "red", lwd = 2)
```

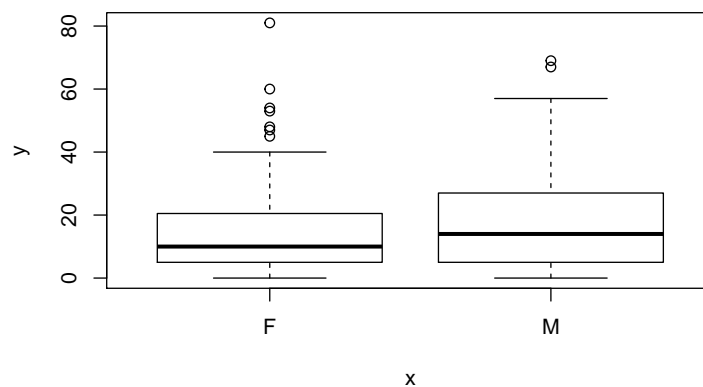


The variable Petal.Length has excessive load at the left to adjust to the normal distribution, we would say that Petal.Length is skewed to left.

4.4.2 Plots of two Variables

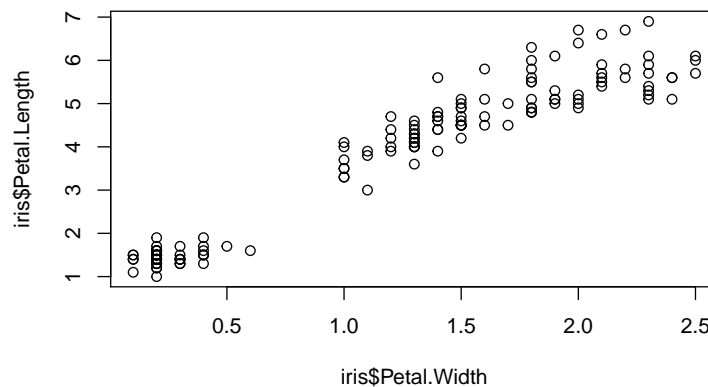
- One variable and factor levels. Factor levels are categorical variables. We can plot for example the variable number of absent days by sex from the absenteeism data set.

```
> plot(absenteeism$sex,absenteeism$days)
```



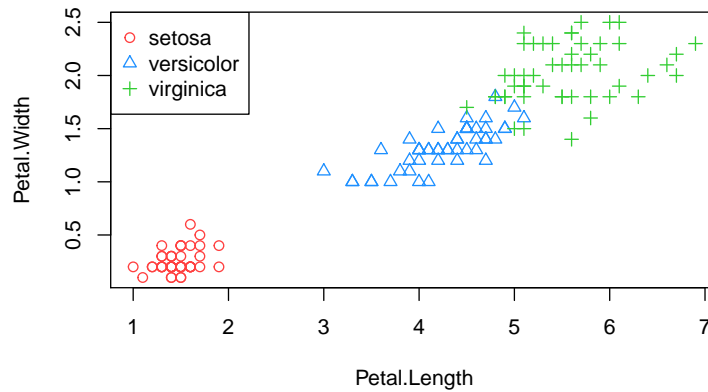
- Explanatory and Response variable. When we are interested on identifying potential causal effects between variables, the scatterplot can be a good choice. The `plot()` function draws axes and adds a scatterplot of points.

```
> plot(iris$Petal.Width,iris$Petal.Length)
```



It seems that the wider the petal the longer it is However the exploratory analysis as the name says is intended to explore not yet to make assessments with confidence. May be adding some colours to distinguish the three species of the iris set might be useful:

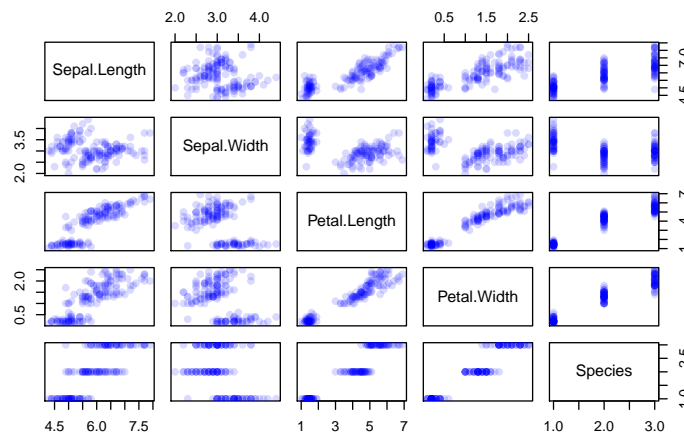
```
> attach(iris)
> plot(Petal.Length,Petal.Width,data=iris,
+      col=c("brown1","dodgerblue1","limegreen")[as.integer(Species)],
+      pch=c(1,2,3)[as.integer(Species)])
> legend(x="topleft",
+      legend=c("setosa","versicolor","virginica"),
+      col=c("brown1","dodgerblue1","limegreen"),
+      pch=c(1,2,3))
```



Yes, it helps. Within the same specie, at least in the case of setosa specie, it seems the relationship between width and length of petals is weaker than it seemed.

The multiple scatterplot can be useful to take a first glance to the relationship among several pairs of variables in a single plot:

```
> plot(iris,
+      col=rgb(0,0,1,.15),
+      pch=19)
```



5 Statistical Analysis

The characteristics of the data set and the research question we want to answer, will determine the choice of the statistical test.

5.1 Data sets and Variables

Variables are characteristics which varies among independent instances. Each variable is distributed across one column and each instance (observation) across one row of the data set, thus, the number of columns is the number of variables and the number of rows the number of observations. We can get both with functions *ncol()* and *nrow()*.

Variables can be categorical such as gender or nationality, with limited values, or continuous when are measured on a proper scale such as age or weight. Furthermore, categorical variables can be ordinal, when indicate order or preference (e.g. first, second, third ...) in contrast to nominal when one category is no better than another (e.g. ethnicity).

Some data sets contain variables which are independent among them in contrast to sets where one or more dependent variables are influenced by one or more independent variables. Sets which contain dependent (response) and independent (predictors) variables can be used to elaborate predictions using supervised learning statistical tools. In contrast, sets without any dependent variable are used to classify observations in groups using unsupervised learning statistical tools. Supervised and unsupervised learning are domains of machine learning and artificial intelligence.

Finally, we will also distinguish variables which values are determined by some common probability distribution function such as the normal, t-student or poisson and we will use parametric statistical tests in contrast to other variables which do not adjust to any of these probability distributions and will have to use non-parametric tests.

The length of the data set, the number of occurrences (rows), can also be decisive to choose the statistical test as only some tests are valid for small data sets (such as less than 30 occurrences).

5.2 Research Question

Sometimes we will elaborate the research question based on the available data; in other cases we will design a process (design of experiments) to collect data that we expect will help to answer our question.

Our research questions, from an statistical perspective, will fail in one of the following two groups: questions to be answered by learning about the

influence among variables or, questions to be answered by learning about how observations can be grouped or classified.

The first group is addressed using statistical tests which fail in the realm of Supervised Learning. The objective of these tools is to learn (discover) about the function f that relates response (dependent, output) variable and predictors (independent or input) variables. Knowing existing relationships is very useful as allows to make predictions of outcomes based on inputs and allows to make inference or judgments about a population based on a sample of data. Some examples of prediction and inference questions cited by Hastie et al. (2021):

- Which predictors are associated with the response? It is often the case that only a small fraction of the available predictors are substantially associated with Y . Identifying the few important predictors among a large set of possible variables can be extremely useful, depending on the application.
- What is the relationship between the response and each predictor? Some predictors may have a positive relationship with Y , in the sense that larger values of the predictor are associated with larger values of Y . Other predictors may have the opposite relationship. Depending on the complexity of f , the relationship between the response and a given predictor may also depend on the values of the other predictors.
- Can the relationship between Y and each predictor be adequately summarized using a linear equation, or is the relationship more complicated? Historically, most methods for estimating f have taken a linear form. In some situations, such an assumption is reasonable or even desirable. But often the true relationship is more complicated, in which case a linear model may not provide an accurate representation of the relationship between the input and output variables,

Classical supervised learning test covered in this manual are contingency tables, linear regression and ANOVA.

The second group of research questions is addressed with unsupervised learning tools which have the objective of classifying observations in meaningful groups. Hastie et al. (2021) cites examples of classification problems:

- A person arrives at the emergency room with a set of symptoms that could possibly be attributed to one of three medical conditions. Which of the three conditions does the individual have?
- An online banking service must be able to determine whether or not a transaction being performed on the site is fraudulent, on the basis of the user's IP address, past transaction history, and so forth.

- On the basis of DNA sequence data for a number of patients with and without a given disease, a biologist would like to figure out which DNA mutations are deleterious (disease-causing) and which are not.

Some of the classification methods included in this course and others are logistic regression, log-linear analysis and cluster analysis.

5.3 Contingency Tables

In section 4.2 we covered frequency tables to count classes of occurrences within a variable, and contingency tables to count occurrences cross-combining categories and/or variables. Apart from exploratory tools tables can also be used as statistical tools when used together with statistical tests.

5.3.1 Pearson's Chi-Square Test

The simplest test to apply to a contingency table is the Chi-Square Test, χ^2 , which its statistic is calculated using the following function:

$$\chi^2 = \frac{(\text{observed} - \text{expected})^2}{\text{expected}}$$

The χ^2 distributions tells the size of the test statistic that could be expected by chance alone (i.e. when the null hypothesis was true). A big value of the test statistic tells that something is happening, and hence that the null hypothesis is false. The χ^2 distribution defines what constitutes a big value of the test statistic (its critical value).

The simplest case for applying the χ^2 test can be to check if two frequencies of two classes of the same variable are different. An example can be the following: an organizer of a music festival takes a random sample of 250 assistants and counts 109 women (43,6%) and 141 men (56,4%). Can the organizer conclude that the sex ratio of assistants is significantly different from 50:50? Using a sample of 250 individuals he wants to infer if the frequency of men and woman are different in the whole group of assistants (population) . The test statistic is computed as:

$$\chi^2 = \frac{(109 - 125)^2}{125} + \frac{(141 - 125)^2}{125} = 4.096$$

The p-value with 1 degree of freedom for 4.096 is 0.04298. The null hypothesis can be discarded with only 4,29% probability of committing a type I error. , that is that is to say, we allow less than 1 in 20 chance of rejecting the null hypothesis when it is actually true. We can infer that the ratio of men is higher than the ratio of woman among the festival assistants. We obtain the p-value for a statistic of 4.096 with 1 degree of freedom which follows the χ^2 probability distribution with the function `pchisq()`:

```
> 1-pchisq(4.096,1)
```

```
[1] 0.0429848
```

To implement the chi-squared test in R we will just use the function `chisq.test()`, which takes as arguments the observed frequencies and in a second vector the expected probabilities:

```
> chisq.test(c(109,141),p=c(0.5,0.5))
```

```
Chi-squared test for given probabilities
```

```
data: c(109, 141)
```

```
X-squared = 4.096, df = 1, p-value = 0.04298
```

We obtain the same result as before, we can reject the null hypothesis and accept that there are more men than women among the assistants.

If the sample had only 100 assistants, and obtained the same percentages of 43,6% women and 56,4% men, would the conclusion be the same?

```
> chisq.test(c(44,56),p=c(0.5,0.5))
```

```
Chi-squared test for given probabilities
```

```
data: c(44, 56)
```

```
X-squared = 1.44, df = 1, p-value = 0.2301
```

No in that case can not discard the null hypothesis, as these frequencies can happen by chance 23.01% of times. Sample size affects the outcome of the χ^2 test.

Chi-Square Test can also be used to test the independence among categorical variables. The null hypothesis of the test is that variables are independent. We use the function `chisq.test()` passing as argument a contingency table created with `table(x,y)`. In the following example we run the chi-square test to test for independence of the variable `lrn` (learning status slow or advanced) respect to ethnicity of the data set `absenteeism`:

```
> absenteeism<-openintro::absenteeism #create data frame
```

```
> attach(absenteeism)
```

```
> mytablrneth<-table(lrn,eth)
```

```
> addmargins(mytablrneth)
```

```
      eth
lrn    A    N Sum
AL    40   43  83
SL    29   34  63
Sum   69   77 146
```

```
> chisq.test(mytablrneth)
```

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

```
data: mytablrneth
```

```
X-squared = 0.0084084, df = 1, p-value = 0.9269
```

The p-value of the test is higher than 0.05 and thus we can not discard the null hypothesis and conclude that these two variables are independent of each other. If we run the test with the variable age we will see that there is dependency between lrn and age as the p-value is lower than 0.05.

```
> mytablrnage<-table(lrn,age)
```

```
> addmargins(mytablrnage)
```

	age				
lrn	F0	F1	F2	F3	Sum
AL	19	15	16	33	83
SL	8	31	24	0	63
Sum	27	46	40	33	146

```
> chisq.test(mytablrnage)
```

```
Pearson's Chi-squared test
```

```
data: mytablrnage
```

```
X-squared = 42.708, df = 3, p-value = 2.838e-09
```

5.3.2 Log-Linear Analysis

Log-linear analysis can also be applied to contingency tables to test the independence or interaction of categorical variables.

As an example, we will build a log-linear model to test the independence among the categorical variables of the absenteeism data set. The objective is to assess if ethnicity, sex, and learning status variables interact among them.

```
> #convert tibble to data frame
```

```
> absenteeism.df<-as.data.frame(absenteeism)
```

```
> # discard vars days and age
```

```
> absenteeism.df <-absenteeism.df[,-c(3,5)]
```

```
> # create frequency table
```

```
> mytab<-ftable(table(absenteeism.df),col.vars = c("lrn"))
```

```
> print(mytab)
```

```
      lrn AL SL
eth sex
```

A	F	19	19
	M	21	10
N	F	21	21
	M	22	13

```
> # optional: check if all the observations have been counted
> nrow(absenteeism)==sum(mytab)
```

```
[1] TRUE
```

With the log-linear test we want to answer questions such: is it significant that the number of slow learners in the group of aboriginal boys is the lowest of all the frequencies?

With the log-linear analysis of contingency tables we will estimate the theoretical frequency, $y_{i,j}$, for each cell of the table. We will do it calculating the cross probability considering that the variables are independent variables and follow the poisson distribution. The poisson probability distribution adjusts well to variables which count occurrences (such as contingency tables).

$$y_{ij} = n \times pr(i) \times pr(j)$$

y_{ij} is the expected frequency in the $cell_{i,j}$ of the contingency table; n is the number of observations and $pr(i)$ and $pr(j)$ are the probabilities of categories i and j .

If we take logarithms on the above equation we obtain the linear model we will use to fit our data points⁵. This is the reason why this model is called log-linear.

$$\log(y_{ij}) = \log(n) + \log(pr(i)) + \log(pr(j))$$

We will test the "mytab" contingency table we have created using the *glm()*, Generalized Linear Models, which can be used to fit log-linear and many other types of linear models. To run the test we will convert the table into a data frame and will set the option *family=poisson*.

```
> mytab.df<-as.data.frame(mytab) #convert the contingency table to a d.f.
> mytab.df
```

	eth	sex	lrn	Freq
1	A	F	AL	19
2	N	F	AL	21
3	A	M	AL	21
4	N	M	AL	22

⁵ $\log(a \times b) = \log(a) + \log(b)$

```

5  A  F  SL  19
6  N  F  SL  21
7  A  M  SL  10
8  N  M  SL  13

```

```

> model<-glm(Freq~eth+sex+lrn, data = mytab.df, family=poisson)
> summary(model)

```

Call:

```
glm(formula = Freq ~ eth + sex + lrn, family = poisson, data = mytab.df)
```

Deviance Residuals:

	1	2	3	4	5	6	7	8
	-0.5488	-0.6230	0.7538	0.4884	0.6478	0.6391	-0.9884	-0.5336

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	3.0678	0.1592	19.271	<2e-16 ***
ethN	0.1097	0.1658	0.662	0.5081
sexM	-0.1924	0.1663	-1.157	0.2473
lrnSL	-0.2757	0.1671	-1.650	0.0989 .

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for poisson family taken to be 1)

Null deviance: 8.1172 on 7 degrees of freedom
 Residual deviance: 3.5858 on 4 degrees of freedom
 AIC: 49.348

Number of Fisher Scoring iterations: 4

From the *summary()* function we observe that none of the variables are significant, we can not conclude that eth, sex and lrn interact among them.

We can also check which is the difference between the observed and expected frequencies:

```

> cbind(model$data,
+       fitted=round(fitted(model),0),
+       dif=round(model$data$Freq-fitted(model),0))

```

	eth	sex	lrn	Freq	fitted	dif
1	A	F	AL	19	21	-2
2	N	F	AL	21	24	-3
3	A	M	AL	21	18	3
4	N	M	AL	22	20	2

5	A	F	SL	19	16	3
6	N	F	SL	21	18	3
7	A	M	SL	10	13	-3
8	N	M	SL	13	15	-2

The differences between the observed frequencies (Freq) and the expected frequencies (fitted) are small. Another check about the model fitness (which can be done using the output of the summary function) is that the Residual deviance 3.59 should not be higher than the degrees of freedom 4, which indicates a good fit of the model.

5.4 Confidence Intervals, CI

CI measures the unreliability of the estimate of the mean, \bar{X} , based on the variance, s^2 and size, n , of the sample from which is obtained. The larger the variance and smaller the size the more unreliable is the estimate, and larger will have to be the CI.

The measure of unreliability used to calculate CI is the standard error, $s.e. = \sqrt{\left(\frac{s^2}{n}\right)}$, where s^2 is the variance and n the sample size.

To obtain the CI for small samples or when the variance of the population is not known the T-Student distribution is used:

$$CI = \bar{X} \pm t_{\alpha/2, d.f.} \times s.e.$$

$t_{\alpha/2}$ is the T-student distribution value for α probability and the number of degrees of freedom is $n - 1$.

When the experimental design/sample sizes are larger or when the standard deviation of the population is known the normal distribution is used and the CI formula is:

$$CI = \bar{X} \pm z_{\alpha/2} \times s.e.$$

We will calculate the confidence interval for the variable days of the data set absenteeism.

```
> sample.mean<-mean(absenteeism$days) # mean
> sample.mean

[1] 16.4589

> n <- length(absenteeism$days)
> n

[1] 146
```

```

> sample.se<-sd(absenteeism$days)/(sqrt(n)) #se
> #will use t distribution with 5% prob with
> # the quantile function qt()
> # with lower tail=F,  $P[X>x]=0.05$ , we get
> #the value x.
> # for normal distribution use qnorm()
> alpha <-0.05
> t.score <-qt(p=alpha/2,df=n-1,lower.tail = F)
> t.score

[1] 1.97646

> margin.error <- t.score*sample.se
> lower.bound <- sample.mean - margin.error
> upper.bound <- sample.mean + margin.error
> print(c(lower.bound,upper.bound))

[1] 13.80032 19.11749

```

With a sample mean of 16.46 days, sample size of 146 observations and standard deviation 16.25, we can assume with 95% probability that the population mean for the variable number of days is within the interval from 13.8 to 19.12.

When implementing CI with R we can just use the function *t.test()* if we want to use the T-Student distribution:

```

> t.test(absenteeism$days)

      One Sample t-test

data:  absenteeism$days
t = 12.236, df = 145, p-value < 2.2e-16
alternative hypothesis: true mean is not equal to 0
95 percent confidence interval:
 13.80032 19.11749
sample estimates:
mean of x
 16.4589

```

To compute the CI using the normal distribution is done with the function *confint* which take as an argument a fitted model such as a linear regression. However rarely we will assume that the variance of the population is known, and it is more frequent to use the t-distribution probability function.

5.5 ANOVA analysis

ANOVA is a powerful statistical model to study the difference of means and standard errors of different groups of data. ANOVA consists in decomposing the variance of the data set in the between-group-variance and the within-group-variance. Both are measured with sum of squares, SS. The ratio of the between SS divided by within-SS results in an F-statistic which is the test statistic for ANOVA. We will use the F-Probability Distribution to assess the critical value from which the null hypothesis will be discarded.

We will use the absenteeism data set as an example. Can we assume that in the schools of New South Wales absenteeism is higher in the aboriginal ethnicity?

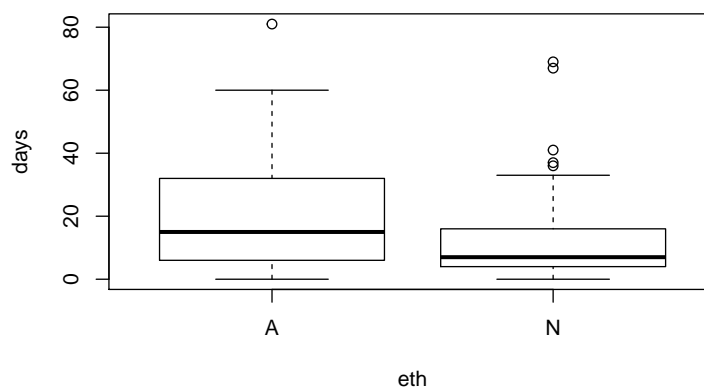
```
> attach(absenteeism)
> means<-tapply(days,list(eth),mean)
> round(means,1)
```

```
  A    N
21.2 12.2
```

The means of the variable absent days, 21.2 and 12.2, might induce to think so.

But if we compare the box-plots of both categories of data, the difference seems less straightforward. Almost all the data of non-aboriginals is comprehended within the data range of aboriginals.

```
> plot(days~eth)
```



To know if we can infer the mean of the population by using the sample data, we will use the ANOVA test, implemented in R with the function `aov()`.


```
> s<-(aov(days~eth))
> summary(s)
```

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
eth	1	2981	2980.5	12.15	0.000651 ***
Residuals	144	35324	245.3		

```
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Now, with the ANOVA result we can assume that aboriginals take more absent days. The between-group mean of the sum of squares is 2980.5 and the within-group mean of the sum of squares is 245.3 which results in an F value of $\frac{2980.5}{245.3} = 12.15$. The p-value for this F value in an F distribution with 1 and 144 degrees of freedom is 0.000651. We discard the null hypothesis.

And, can we assume that in the schools of New South Wales men are absent more days than women?

```
> attach(absenteeism)
> means<-tapply(days,list(sex),mean)
> round(means,1)
```

```
      F      M
15.2 18.0
```

It is large enough the difference between sample means?

```
> s<-(aov(days~sex))
> summary(s)
```

	Df	Sum Sq	Mean Sq	F value	Pr(>F)
sex	1	269	269.4	1.02	0.314
Residuals	144	38035	264.1		

The answer is No. The p-value for the F-statistic is 0.31. Too high to be statistically significant.

And what about adding both factors at once, that is sex and ethnicity?. That is, we have one dependent variable, number of absent days, and two independent variables, ethnicity and sex; that is a multivariate analysis of variance.

```
> attach(absenteeism)
> means<-tapply(days,list(sex,eth),mean)
> round(means,1)
```

```
      A      N
F 20.9 10.1
M 21.6 14.7
```

```

> s<-(aov(days~sex+eth))
> summary.lm(s)

Call:
aov(formula = days ~ sex + eth)

Residuals:
    Min       1Q   Median       3Q      Max
-20.762  -9.919  -5.729   5.183  61.016

Coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept)   19.984      2.218   9.010 1.2e-15 ***
sexM           2.778      2.603   1.067 0.287771
ethN          -9.065      2.595  -3.493 0.000636 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 15.65 on 143 degrees of freedom
Multiple R-squared:  0.0851,    Adjusted R-squared:  0.0723
F-statistic:  6.65 on 2 and 143 DF,  p-value: 0.001731

```

From the *aov* output:

- the t-statistic for the intercept is significant
- the t-statistic for the variable ethnicity is significant
- the t-statistic for the variable sex is not significant.
- the F-statistic for the overall model is significant
- the Multiple R-squared of the model is low, only 8.51 percent of the variance is explained by the sex and ethnicity variables.

When there is more than one independent variable in the model, sometimes the ANOVA analysis is conducted in what is called the two-way analysis, that is considering new factors consisting on the interaction of independent variables.

The two-way analysis using as independent variables sex and ethnicity would be:

```

> s<-(aov(days~sex*eth))
> summary.lm(s)

Call:
aov(formula = days ~ sex * eth)

```

Residuals:

Min	1Q	Median	3Q	Max
-20.921	-9.921	-4.893	6.022	60.079

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	20.9211	2.5433	8.226	1.12e-13 ***
sexM	0.6919	3.7944	0.182	0.8556
ethN	-10.8496	3.5101	-3.091	0.0024 **
sexM:ethN	3.9510	5.2224	0.757	0.4506

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 15.68 on 142 degrees of freedom

Multiple R-squared: 0.08877, Adjusted R-squared: 0.06952

F-statistic: 4.611 on 3 and 142 DF, p-value: 0.004134

Although we have added a variable (interaction sex and ethnicity) the performance of the model has decreased as it has a higher p-value for the F-Statistic and lower adjusted R-squared.

5.6 Tests for applying hypothesis contrast and confidence interval.

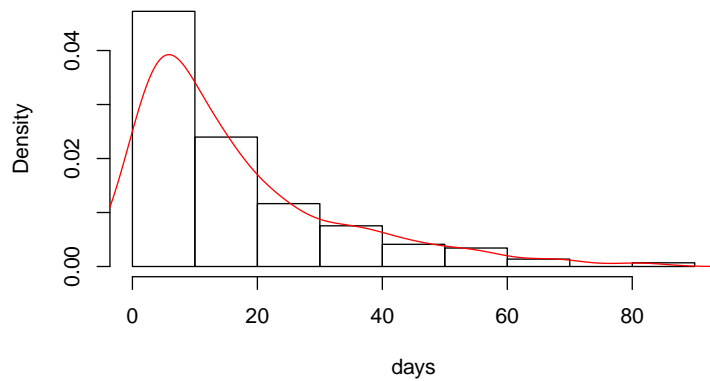
Statistical analysis models become simpler if some assumptions about the probability distribution of the data can be made. Some of the assumptions that will like to test specially if want to build predictive models are:

- Does the data follow a normal distribution?
- Is variance constant?

5.6.1 Normality test.

Plotting a histogram of the variable of interest will show the shape of the probability density function which can be a good indication to know if the data follows a normal distribution.

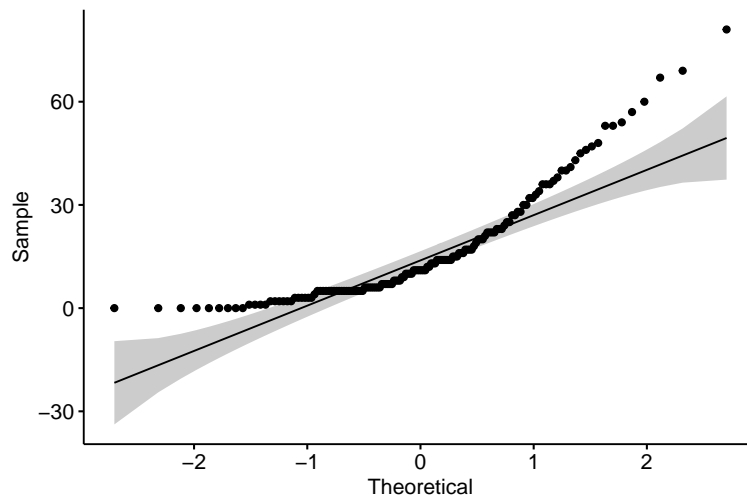
```
> hist(absenteeism$days,probability=T, main="",xlab="days")
> lines(density(absenteeism$days),col=2)
```



The histogram shows that the variable departs from a normal distribution.

Another visual judgment about whether the distribution is bell shaped, is the Q-Q plot or quantile-quantile plot which draws the correlation between a given sample and the normal distribution.

```
> library(ggpubr)
> ggqqplot(absenteeism$days)
```



If the data is normally distributed, all the dots will be close to the line. Again, absent days is suspiciously non-normal.

Kolmogorov-Smirnov and Shapiro-Wilk's methods are two widely used normality tests in addition to the mentioned visual methods. Both tests can be called in R with their respective functions, *ks.test* and *shapiro.test*.

```
> ks<-ks.test(absenteeism$days,pnorm, mean(absenteeism$days),sd(absenteeism$days))
> ks
```

One-sample Kolmogorov-Smirnov test

```
data: absenteeism$days
D = 0.16971, p-value = 0.000445
alternative hypothesis: two-sided
```

The p-value for the Kolmogorov-Smirnov normality test is 0.00045, < 0.05 , so we reject the null hypothesis that the data follows a normal distribution with more than 95% confidence.

```
> shapiro<-shapiro.test(absenteeism$days)
> shapiro
```

Shapiro-Wilk normality test

```
data: absenteeism$days
W = 0.83631, p-value = 1.81e-11
```

The p-value for the Shapiro-Wilk normality test is very low, $1.81e-11$, so we can confidently reject the null hypothesis that the data follows a normal distribution.

5.6.2 Constant Variance Assumption. Homoscedasticity test.

Variance analysis is conducted after specifying a lineal model using continuous variables and analyzing the model residuals, the variability not explained by the model.

We will use as an example the cars data set which has 50 observations of two continuous variables: speed (mph) and dist for stopping distance (ft).

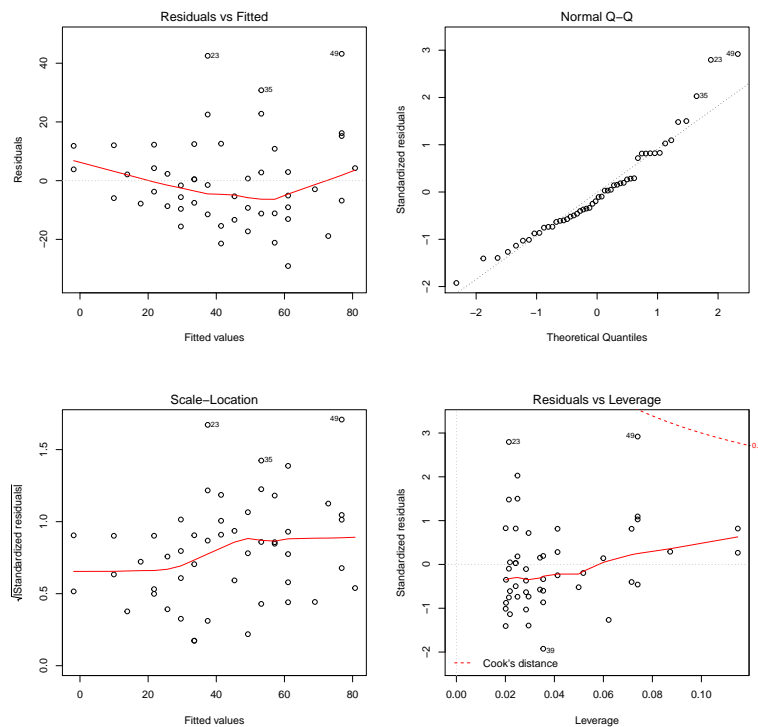
```
> head(cars)
```

	speed	dist
1	4	2
2	4	10
3	7	4
4	7	22
5	8	16
6	9	10

We will build a model using as dependent variable, the stopping distance, and as independent variable the speed, and will analyze the residuals to assess if the variance of the variable dist depends on the variable speed,

and thus, there is heteroscedasticity which would invalidate some of the conclusions we can obtain with the regression model.

```
> #create linear model
> lmMod<-lm(dist~speed,data=cars)
> # config 2x2 = 4 charts in 1 panel
> par(mfrow=c(2,2))
> plot(lmMod)
```



The the two plots of the left side, and their red lines indicates that the dispersion of residuals are related with the fitted values. If variance is constant across observations (homoscedasticity) the red lines are flat, but if as in this example the variance is not constant (heteroscedasticity), the lines take some different shapes. In the example, the higher the speed the higher the variance, variance is not constant.

There are also statistical tests for homoscedasticity. A usual one is the Breusch-Pagan test which can be called with the `bptest()` function of the `lmtest` library.

```
> bp<-lmtest::bptest(lmMod)
> bp
```

studentized Breusch-Pagan test

```
data:  lmMod
BP = 3.2149, df = 1, p-value = 0.07297
```

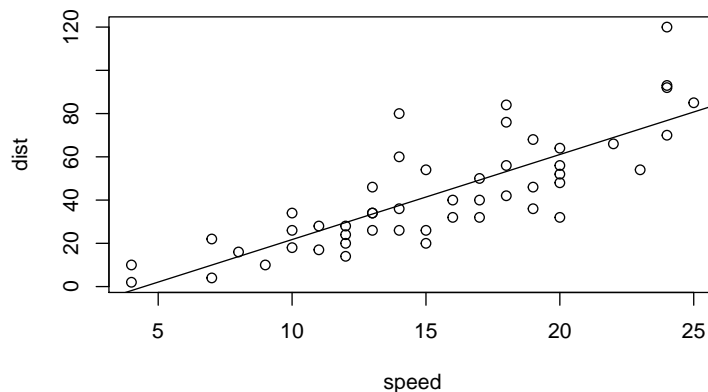
The p-value 0.07297 is higher than 0.05 and we can not reject with 95% confidence the null hypothesis that variance is constant. We confirm the heteroscedasticity observed with the Q-Q plots.

5.7 Correlation analysis

The correlation coefficient ρ is a measure of the closeness of association of the points of a scatterplot to the regression line based on those points. It is a measure of the strength of the linear relationship between two continuous variables. A correlation of -1 indicates perfectly negative correlation, 0 absence of correlation, and +1 perfectly positive correlation.

The following chart contains the regression line between the dependent variable dist and the independent variable speed of the data set cars, together with the observation points.

```
> attach(cars)
> lModel<-lm(dist~speed, data=cars)
> plot(speed,dist)
> abline(lModel)
```



Visually, it seems that the distance and speed are positively correlated, the higher the speed the longer the distance needed to stop the car.

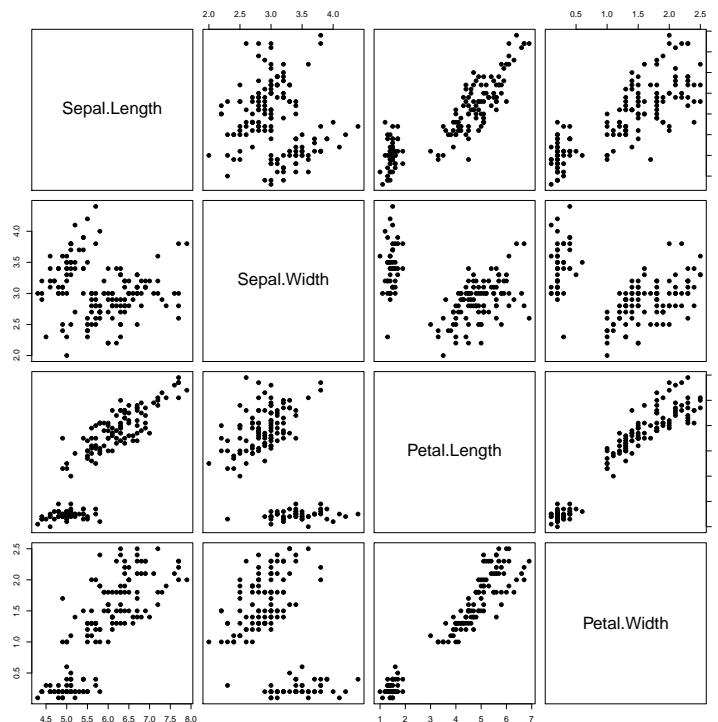
```
> cor(dist,speed)
[1] 0.8068949
```

The high correlation coefficient of 0.81, confirms the visual intuition.

A very useful R function to visually scan potential correlations among pairs of variables is the function *pairs* which plots all the possible pairs of scatter plots.

We can plot correlations among the continuous variables of the iris data set. We exclude column 5, species, which is categorical.

```
> pairs(iris[,1:4], pch = 19)
```



Visually, it seems that all the variables have some positive correlation among them with the exception of Sepal.Width. The the following code from STHDA (2021) creates a more informative and attractive plot.

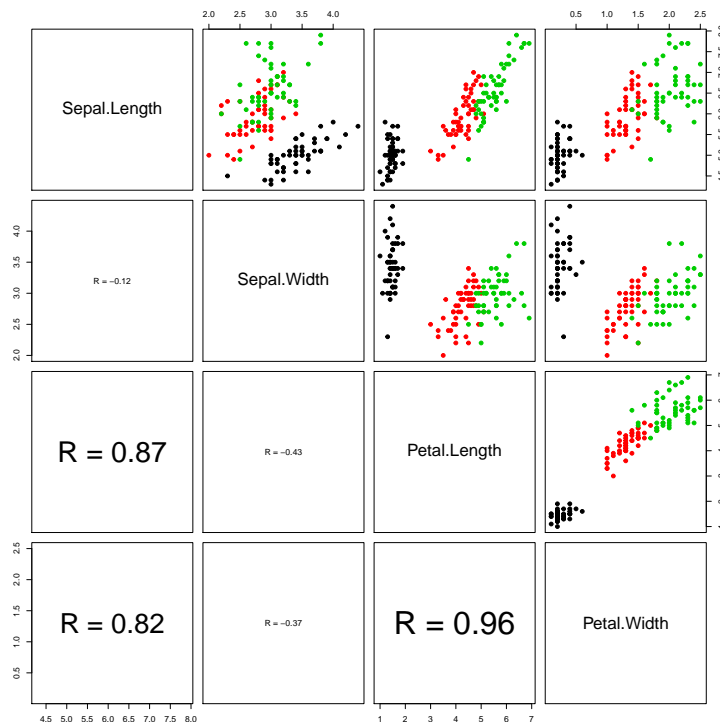
```
> # Correlation panel
> panel.cor <- function(x, y){
+   usr <- par("usr"); on.exit(par(usr))
+   par(usr = c(0, 1, 0, 1))
+   r <- round(cor(x, y), digits=2)
+   txt <- paste0("R = ", r)
+   cex.cor <- 0.8/strwidth(txt)
+   text(0.5, 0.5, txt, cex = cex.cor * r)
+ }
> # Customize upper panel
> upper.panel<-function(x, y){
```



```

+ points(x,y, pch = 19, col = iris$Species)
+ }
> # Create the plots
> pairs(iris[,1:4],
+       lower.panel = panel.cor,
+       upper.panel = upper.panel)

```



5.8 Regression analysis

Regression is a useful tool for predicting a quantitative response (dependent variable) based on input data or predictors (independent variables). Regression analysis is the basis of advanced statistical models but also is widely used in its simplest form.

With regression we can address questions such as there is a relationship between CO_2 emissions and atmosphere temperature?, How strong is the relationship between advertising campaign budget and number of seats in the Spanish Parliament?, How accurately can we predict the unemployment rate? There is a linear relationship between a country democracy index and the percentage of university students that obtain a grade in political sciences?

We will cover three types of regression analysis:

- Simple Linear Regression
- Multiple Linear Regression
- Logistic Regression

5.8.1 Simple Linear Regression

It is a model where we have one dependent variable (Y) and one independent variable (X), and we assume that they are linearly correlated:

$$Y \approx \beta_0 + \beta_1 X$$

β_0 and β_1 are the unknown coefficients and we will estimate them by using the pairs of data $(x_1, y_1), (x_2, y_2), \dots, (x_n, y_n)$

We compute the linear regression in R with the function `lm()`. We will use as example the data set `cars` with dependent variable (Y) distance to stop, and independent variable (X), speed. Notice the use of the symbol⁶ to separate the dependent (left) from the independent variables (right).

```
> fit = lm(dist ~ speed, data = cars)
> fit
```

Call:

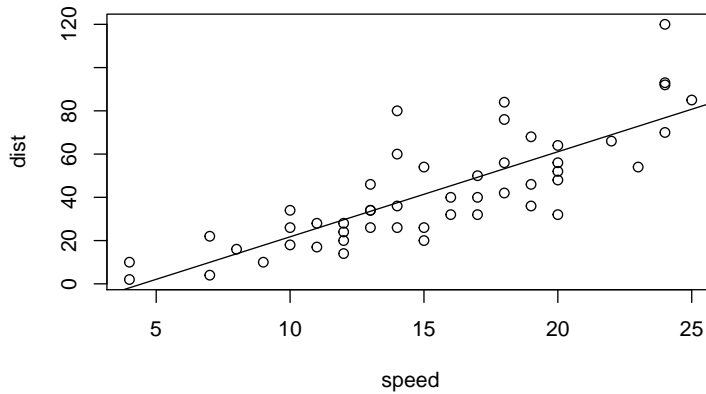
```
lm(formula = dist ~ speed, data = cars)
```

Coefficients:

(Intercept)	speed
-17.579	3.932

```
> b = coef(fit)
> plot(cars) #scatterplot
> abline(fit) #plot regression line
```

⁶in most keyboards AltGr+F4



The equation line is:

$$dist_i = -17.58 + 3.93speed_i + \epsilon_i$$

The minimum sum of squares method used to obtain the regression analysis assumes that the mean of ϵ_i is zero. The *summary* function provides most of the needed output of the regression analysis.

```
> summary(fit)
```

Call:

```
lm(formula = dist ~ speed, data = cars)
```

Residuals:

	Min	1Q	Median	3Q	Max
	-29.069	-9.525	-2.272	9.215	43.201

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	-17.5791	6.7584	-2.601	0.0123 *
speed	3.9324	0.4155	9.464	1.49e-12 ***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 15.38 on 48 degrees of freedom

Multiple R-squared: 0.6511, Adjusted R-squared: 0.6438

F-statistic: 89.57 on 1 and 48 DF, p-value: 1.49e-12

The model has very low (significant) p-values.

5.8.2 Multiple Linear Regression

Here we have one dependent variable and several independent variables. In order to fit a multiple linear regression model using least squares, we again use the `lm()` function. The syntax `lm(y ~ x1 + x2 + x3)` is used to fit a model with three predictors, `x1`, `x2`, and `x3`. The `summary()` function now outputs the regression coefficients for all the predictors, in this case we will have as many slope coefficients as dependent variables. Each coefficient will also have its own p-value.

```
> model<-lm(iris$Petal.Length~.,data=iris[,c(1:2,4)])
> summary(model)
```

Call:

```
lm(formula = iris$Petal.Length ~ ., data = iris[, c(1:2, 4)])
```

Residuals:

	Min	1Q	Median	3Q	Max
	-0.99333	-0.17656	-0.01004	0.18558	1.06909

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)
(Intercept)	-0.26271	0.29741	-0.883	0.379
Sepal.Length	0.72914	0.05832	12.502	<2e-16 ***
Sepal.Width	-0.64601	0.06850	-9.431	<2e-16 ***
Petal.Width	1.44679	0.06761	21.399	<2e-16 ***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

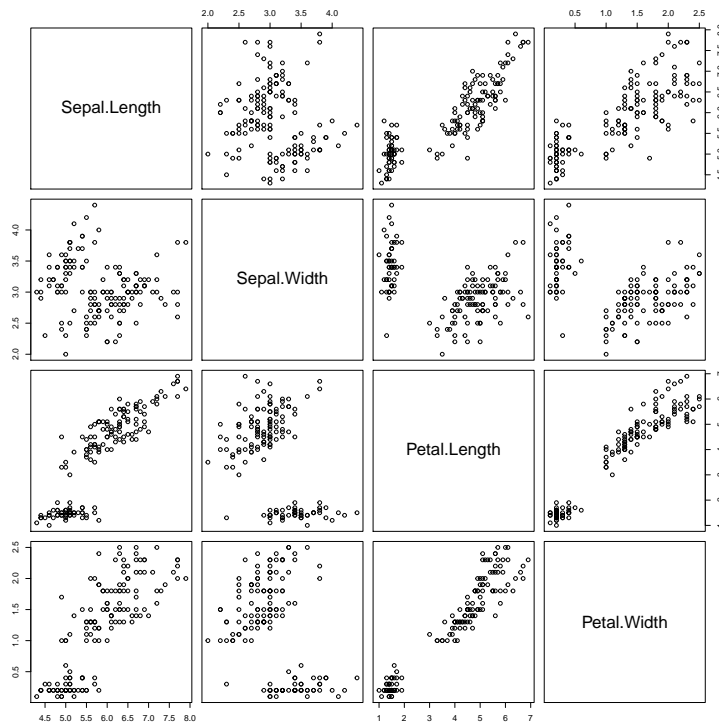
Residual standard error: 0.319 on 146 degrees of freedom

Multiple R-squared: 0.968, Adjusted R-squared: 0.9674

F-statistic: 1473 on 3 and 146 DF, p-value: < 2.2e-16

The three predictors have coefficients with very low p-values, that is they are statistically significant. Nevertheless, it is interesting to take a look at the scatterplot:

```
> pairs(iris[,-5])
```



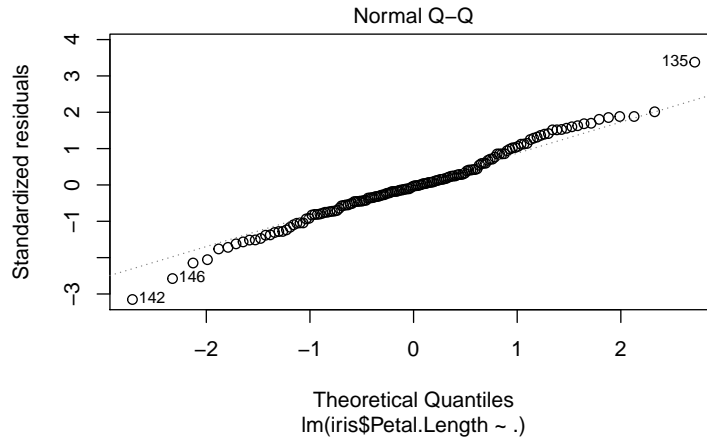
- Petal.Length and Sepal.Length do not seem to have a linear relationship, specially the small ones. Points do not seem to spread around a regression line. This would invalidate the use of Sepal.Length as a predictor.
- It seems there is linear correlation between Sepal.Length and Petal.Width. Robust linear models have independent variables uncorrelated between. When this happens the model has collinearity which can be an issue to fit models as well as to interpret results.

A further step needed to assess the robustness of the model is the analysis of residuals. We will use the Durbin Watson test to investigate serial correlation, and the Q-Q plot to assess if residuals follow a normal distribution. Serial correlation appears when a variable is correlated with its self, as if it had memory of the last values it has taken.

```
> library(car)
> durbinWatsonTest(model)

lag Autocorrelation D-W Statistic p-value
  1      0.1030362      1.782967   0.124
Alternative hypothesis: rho != 0

> plot(model, which=2)
```



The p-value of the Durbin Watson test is higher than 0.05. We can not discard the null hypothesis that residuals are not correlated. There is no serial correlation. The distribution of residuals in the Q-Q plot are around the diagonal which indicates that follow a normal distribution.

5.8.3 Logistic Regression

The Y variable of logistic regression models is a probability estimate. The objective of the model is to assess if, and how much, the X independent variables influence the value of the Y dependent variable. The probability estimated with the logistic model is often used as a probability related with a binary variable such as a yes / no, success/ failure, The response variable expressed in this way is categorical in contrast to the one of the linear regression model which always is continuous. The logistic regression model, used in this way, is a classification statistical tool which assesses if an observation belongs to a certain class, i.e. yes or no. It is possible to extend the logistic model to multinomial logistic regression models, not covered in this course, with categorical response variables with more than two classes (not binary).

The response variable of the logistic regression is constrained to the range $[0,1]$ because it is a probability. The logistic model instead of a linear function (which has no limited range) uses the logit function which always has the outcome between $[0,1]$. Another related model, is the probit model, not covered in this course, which uses the normal distribution probability function, which also accomplishes the $[0,1]$ requirement but might be harder to compute.

The probability in the logistic model is computed with the following function:

$$p = \frac{e^{\beta_0 + x_1\beta_1 + \dots + x_q\beta_q}}{1 + e^{\beta_0 + x_1\beta_1 + \dots + x_q\beta_q}}$$

and can be transformed to:

$$\frac{p}{1-p} = e^{\beta_0 + x_1\beta_1 + \dots + x_q\beta_q}$$

and taking logarithms to both sides we have:

$$\log\left(\frac{p}{1-p}\right) = \beta_0 + x_1\beta_1 + \dots + x_q\beta_q$$

The $\frac{p}{1-p}$ ratio is called the odds ratio which is a different way of presenting information on probabilities, and the log of the odds ratio is called the logit. Odds is an alternative way to present probabilities. For example, in horse race betting, odds for 2 to 1 for a horse, means that expects winning two of every three races (will win 2 and loose 1) and the probability to win is $p = \frac{2}{3} = 0.66$. Odds for can be obtained from probabilities with $\frac{p}{1-p}$ and odds against with the inverse, $\frac{1-p}{p}$.

As in the case of linear regression, we can have simple and multiple regression depending on the number of predictors or independent X variables included in the model.

Logistic regression is implemented in R with the *glm()* function which instead of the minimum squares method used in linear regression, uses maximum likelihood to fit the model. Maximum likelihood (L) is a very general approach that can fit both categorical and continuous response variables no matter if they are modeled thorough linear or a non-linear relationship with the predictor variables.

The following example adapted from ? uses a logistic regression model with the iris data set to classify plants of the specie virginica. To apply the logistic regression will follow the following steps:

1. Prepare the data: Iris is already a tidy data set. We will only have to create a binary variable `iris$virginica` which will take the value True if the observation belongs to the virginica specie or False if otherwise. We also add a useless variable to later test if the the logistic regression takes or drops it from the model. Can check 3.2.4 on how to create and add calculated variables to a data set.
2. Create a Logistic Regression Model: We will use the *glm()* function of the *stats* package of R. The model will link the binary variable we have created with the other variables of the set using the logit function.

3. Check which variables are significant. We use the *summary()* function and check the p-values for the coefficients of the model.
4. Improve the model by adding - dropping variables . Initially all the variables of the set are offered to the model, however we will use the *step* function which iterates on the model by adding and subtracting variables to improve its fitness measured with the AIC (Akaike's An Information Criterion). For models fitted by maximum likelihood, the smaller the AIC the better the fit. We follow the parsimonious modeling principle.
5. Use the model to classify observations as virginica or not. Here we use in-sample testing on the data we learned the data from. To get a generalization error estimate you should use a test set or cross-validation!
6. Check classification Performance

```
> #step 1
> data(iris)
> x <- iris[sample(1:nrow(iris)),]
> x <- cbind(x, useless = rnorm(nrow(x)))
> x$virginica <- x$Species == "virginica"
> x$Species <- NULL

> #step 2
> model <- glm(virginica ~ .,
+   family = binomial(logit), data=x)
> model
```

```
Call:  glm(formula = virginica ~ ., family = binomial(logit), data = x)
```

Coefficients:

(Intercept)	Sepal.Length	Sepal.Width	Petal.Length	Petal.Width
-97.305	-3.894	-12.837	17.533	43.325
useless				
3.592				

```
Degrees of Freedom: 149 Total (i.e. Null); 144 Residual
```

```
Null Deviance: 191
```

```
Residual Deviance: 9.202 AIC: 21.2
```

About the warning: glm.fit: fitted probabilities numerically 0 or 1 occurred means that the data is possibly linearly separable.

```
> #step 3
> summary(model)
```

```
Call:
```



```
glm(formula = virginica ~ ., family = binomial(logit), data = x)
```

Deviance Residuals:

Min	1Q	Median	3Q	Max
-1.773	0.000	0.000	0.000	1.591

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	-97.305	68.878	-1.413	0.158
Sepal.Length	-3.894	3.660	-1.064	0.287
Sepal.Width	-12.837	8.290	-1.549	0.122
Petal.Length	17.533	11.834	1.482	0.138
Petal.Width	43.325	28.735	1.508	0.132
useless	3.592	3.098	1.160	0.246

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 190.9543 on 149 degrees of freedom
 Residual deviance: 9.2025 on 144 degrees of freedom
 AIC: 21.202

Number of Fisher Scoring iterations: 14

```
> #step 4
```

```
> model2<-step(model,data=x)
```

Start: AIC=21.2

```
virginica ~ Sepal.Length + Sepal.Width + Petal.Length + Petal.Width +  
            useless
```

	Df	Deviance	AIC
- Sepal.Length	1	10.7755	20.776
<none>		9.2025	21.202
- useless	1	11.8985	21.899
- Sepal.Width	1	15.1706	25.171
- Petal.Length	1	23.0471	33.047
- Petal.Width	1	23.7618	33.762

Step: AIC=20.78

```
virginica ~ Sepal.Width + Petal.Length + Petal.Width + useless
```

	Df	Deviance	AIC
<none>		10.775	20.776
- useless	1	13.266	21.266
- Sepal.Width	1	20.453	28.453

```
- Petal.Length  1  25.220 33.220
- Petal.Width   1  31.451 39.451
```

The best model includes only Sepal.Width, Petal.Length and Petal.Width and has an AIC of 20.78 from an initial model which had all the variables with an AIC of 21.2. The significance of the model is obtained with the *summary* function.

```
> summary(model2)
```

Call:

```
glm(formula = virginica ~ Sepal.Width + Petal.Length + Petal.Width +
     useless, family = binomial(logit), data = x)
```

Deviance Residuals:

	Min	1Q	Median	3Q	Max
	-1.47653	-0.00001	0.00000	0.00000	1.84779

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	-74.561	46.421	-1.606	0.1082
Sepal.Width	-12.638	7.089	-1.783	0.0746 .
Petal.Length	10.090	5.913	1.706	0.0879 .
Petal.Width	36.889	22.376	1.649	0.0992 .
useless	2.424	2.187	1.108	0.2677

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for binomial family taken to be 1)

```
Null deviance: 190.954 on 149 degrees of freedom
Residual deviance: 10.776 on 145 degrees of freedom
AIC: 20.776
```

Number of Fisher Scoring iterations: 13

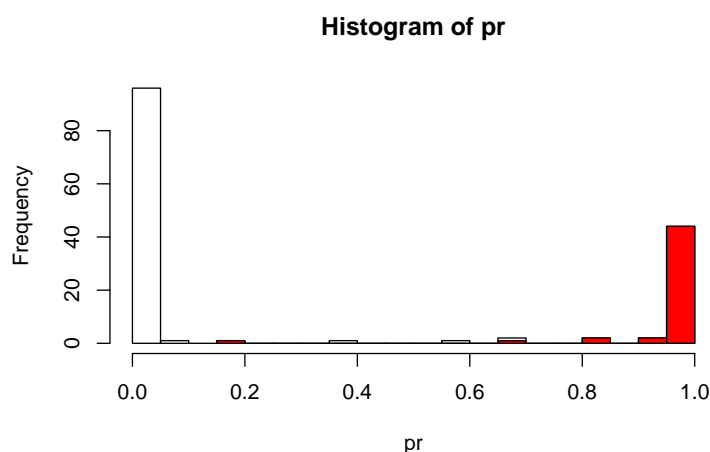
The estimates β_0, β_1, \dots are log-odds and can be converted into odds using e^β . A negative log-odds ratio means that the odds go down with an increase in the value of the predictor. A predictor with a positive log-odds ratio increases the odds. In this case, the odds of looking at a Virginica iris goes down with Sepal.Width and increases with the other two predictors.

```
> #step 5
> pr <- predict(model2, x, type="response")
> head(round(pr, 2))
```

```
94 115 134 44 27 74
```

```
0.00 1.00 0.18 0.00 0.00 0.00
```

```
> hist(pr, breaks=20)
> hist(pr[x$virginica==TRUE], col="red", breaks=20, add=TRUE)
```



```
> table(actual=x$virginica, predicted=pr>.5)
```

	predicted	
actual	FALSE	TRUE
FALSE	98	2
TRUE	1	49

5.9 Principal Component Analysis

Principal components allow to summarize a data set with a small number of representative variables that collectively explain most of the variability in the original set. Principal components analysis (PCA) refers to the process by which principal components are computed. PCA is an unsupervised approach, since it involves only a set of features X_1, X_2, \dots, X_p and no associated response Y . Principal Component Analysis finds a set of orthogonal standardized linear combinations which together explain all of the variation in the original data. Evaluating the meaning of the obtained principal components, the variables, can be difficult although it can also provide potentially interesting insights to the analyst.

PCA is specially useful to reduce the number of variables in data sets which have correlated variables, a problem in data sets which have collinearity, that is independent variables correlated among themselves.

We will use as example the iris data set to identify the PCA of the set. The iris data set contains 150 observations of plants and using PCA we will try to find linear combinations of the numerical variables (PCA can not be applied

to categorical variables) that maximize the variation contained within them, thereby displaying most of the original variation in fewer dimensions.

We will implement PCA in R using the *prcomp* function.

```
> head(iris)

  Sepal.Length Sepal.Width Petal.Length Petal.Width Species
1          5.1         3.5          1.4          0.2  setosa
2          4.9         3.0          1.4          0.2  setosa
3          4.7         3.2          1.3          0.2  setosa
4          4.6         3.1          1.5          0.2  setosa
5          5.0         3.6          1.4          0.2  setosa
6          5.4         3.9          1.7          0.4  setosa
```

```
> model<-prcomp(iris[, -5], center=TRUE, scale=TRUE)
> model
```

```
Standard deviations (1, ..., p=4):
[1] 1.7083611 0.9560494 0.3830886 0.1439265
```

```
Rotation (n x k) = (4 x 4):

          PC1          PC2          PC3          PC4
Sepal.Length  0.5210659 -0.37741762  0.7195664  0.2612863
Sepal.Width   -0.2693474 -0.92329566 -0.2443818 -0.1235096
Petal.Length  0.5804131 -0.02449161 -0.1421264 -0.8014492
Petal.Width   0.5648565 -0.06694199 -0.6342727  0.5235971
```

We have identified four principal component variables, which have the above indicated loads on the existing four variables of the original data set. We have four new variables for each of the 150 original observations:

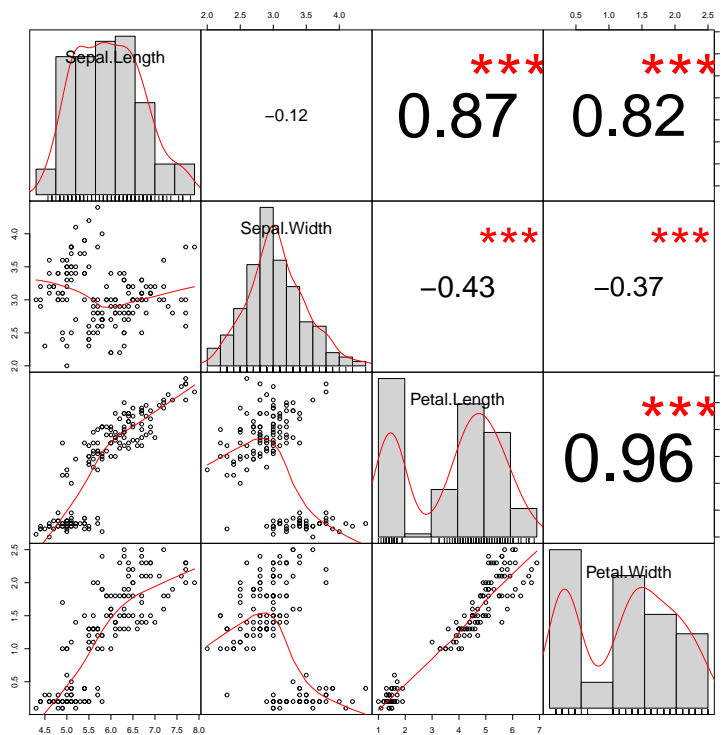
```
> head(model$x)

          PC1          PC2          PC3          PC4
[1,] -2.257141 -0.4784238  0.12727962  0.024087508
[2,] -2.074013  0.6718827  0.23382552  0.102662845
[3,] -2.356335  0.3407664 -0.04405390  0.028282305
[4,] -2.291707  0.5953999 -0.09098530 -0.065735340
[5,] -2.381863 -0.6446757 -0.01568565 -0.035802870
[6,] -2.068701 -1.4842053 -0.02687825  0.006586116
```

With the function *summary()* we can see the importance of each variable in terms of the variability that explains. `summary(model)` The first component explains 72,96% of the total variation, here is the plot of this model, showing the importance of PC1. PC3 and PC4 might be disregarded as both together do only explain 3.6% of variation.

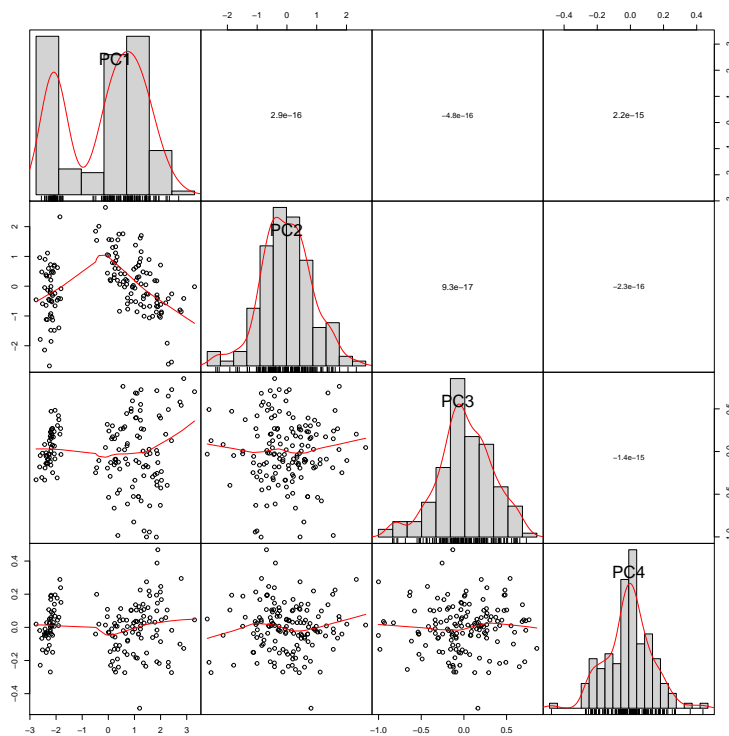
The variables contained in the original set show important correlation among them:

```
> library(PerformanceAnalytics)
> chart.Correlation(iris[, -5], method="pearson")
```



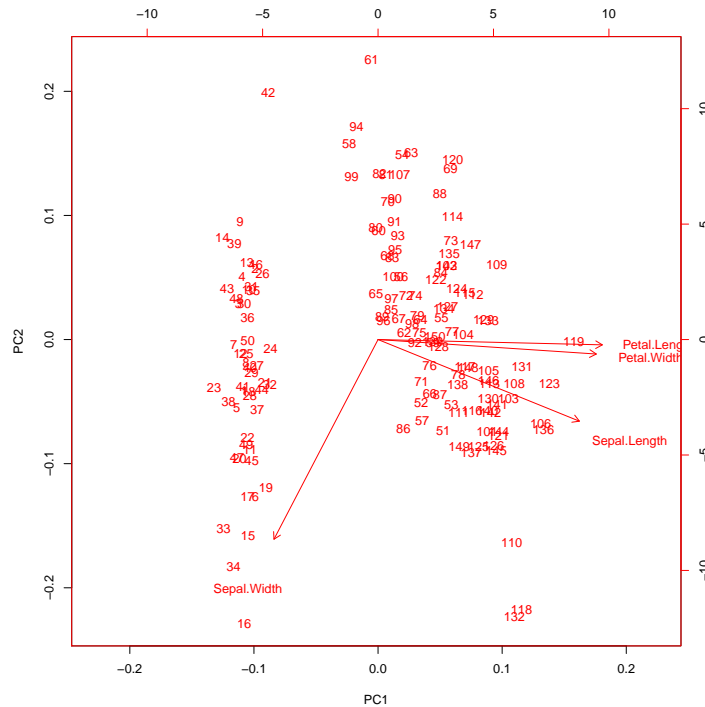
While the PCAs new variables have 0 correlation:

```
> library(PerformanceAnalytics)
> chart.Correlation(model$x, method="pearson")
```



With the *biplot* function we obtain a visual representation of the PCA:

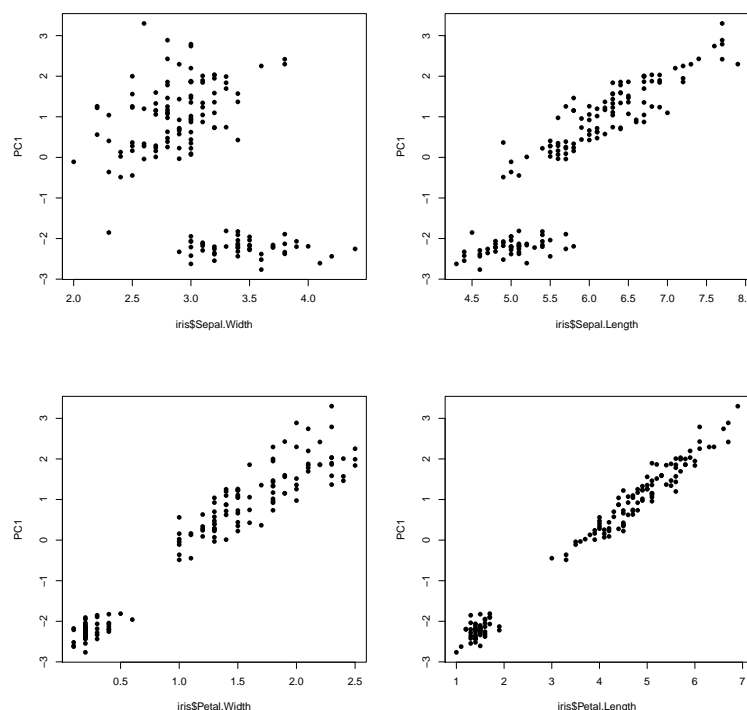
```
> biplot(model, main="", col="red", ellipse=TRUE)
```



```
> pred<-predict(model,iris[1:5,-5])
> pred
```

	PC1	PC2	PC3	PC4
1	-2.257141	-0.4784238	0.12727962	0.02408751
2	-2.074013	0.6718827	0.23382552	0.10266284
3	-2.356335	0.3407664	-0.04405390	0.02828231
4	-2.291707	0.5953999	-0.09098530	-0.06573534
5	-2.381863	-0.6446757	-0.01568565	-0.03580287

```
> PC1<-predict(model)[,1]
> PC2<-predict(model)[,2]
> par(mfrow=c(2,2))
> plot(iris$Sepal.Width,PC1,pch=16)
> plot(iris$Sepal.Length,PC1,pch=16)
> plot(iris$Petal.Width,PC1,pch=16)
> plot(iris$Petal.Length,PC1,pch=16)
```



PC1 is strongly correlated with Petal.Width and Petal.Length.

Principal Component Analysis are at the hearth of the algorithms used by many recommendation systems such as the ones used by Netflix to suggest a movie that a particular customer might like, as cited in Hastie et al. (2021).

5.10 Factor Analysis

Factor analysis (FA) is a technique similar to Principal Component Analysis (PCA) as both are dimension reduction techniques which have the objective of explaining the variability of the set with minimum number of variables. The key difference with PCA is that with FA there is the assumption that correlations among all observed variables can be explained by latent (un-observed) variables also known as factors, which are the equivalent to the principal components computed with PCA. A factor is a common element with which several other variables are correlated. For example, the socio-economic category can be a factor (not directly observable) but correlated with observable variables such as education level, income, employment status,

Computed FA factors can help to understand the data dynamics and can have meaning for the analyst, on the other hand, the principal components obtained with PCA not necessarily have a meaning.

FA is implemented in R with *factanal* function. As an example we try to reduce the four quantitative variables of the iris data set on a single factor:

```
> model<-factanal(iris[,-5],factors=1)
> model
```

Call:

```
factanal(x = iris[, -5], factors = 1)
```

Uniquenesses:

Sepal.Length	Sepal.Width	Petal.Length	Petal.Width
0.240	0.822	0.005	0.069

Loadings:

	Factor1
Sepal.Length	0.872
Sepal.Width	-0.422
Petal.Length	0.998
Petal.Width	0.965

	Factor1
SS loadings	2.864
Proportion Var	0.716

Test of the hypothesis that 1 factor is sufficient.

The chi square statistic is 85.51 on 2 degrees of freedom.

The p-value is 2.7e-19

The model explains 71,6% of variability and has a significant 0 p-value.

FA decomposes the variability of the model into: variability explained by the factor/s, and unique variability explained by the existing variable. Unique variability can be found in the model output:

```
> model$uniquenesses
```

Sepal.Length	Sepal.Width	Petal.Length	Petal.Width
0.24022872	0.82186526	0.00500000	0.06933342

The variables related with petals dimensions are very well explained by the factor, 0.5% and 6.93% uniqueness, while the width and length of the sepal are variables with high uniqueness, 82.19% and 24.02% not explained by the factor.

Cluster Analysis is a set of techniques that look for subgroups (clusters) in the data. Objects belonging to the same group resemble each other. Objects belonging to different groups are dissimilar. Cluster Analysis is an unsupervised learning method which has two main different approaches, K-means clustering and Hierarchical Clustering.

K-means clustering is implemented in R with the *kmeans()* function. Each observation is assigned to the group with the nearest centroid. The *kmeans* function minimizes the within-cluster sum of squares (variability).

K-means clustering with 2 clusters of sizes 97, 53

	Sepal.Length	Sepal.Width	Petal.Length	Petal.Width
1	6.301031	2.886598	4.958763	1.695876
2	5.005660	3.369811	1.560377	0.290566

[illegible]

```
[1] 123.79588  28.55208
(between_SS / total_SS = 77.6 %)
```

```
[1] "cluster"      "centers"      "tottss"       "withinss"     "tot.withinss"
[6] "betweenss"    "size"         "iter"         "ifault"
```

We have split the 150 observations in two clusters which explain 77.64% of set variability.

We add the cluster to the original set as a new variable called cluster:

```
> iriskm<-cbind(iris,cluster=kmModel$cluster)
> table(iriskm[,5:6])
```

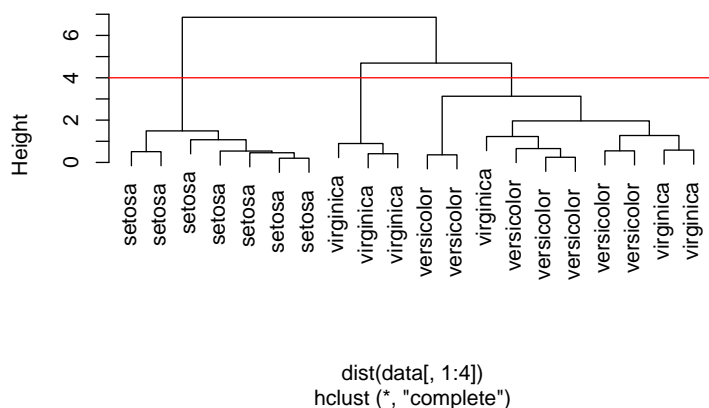
	cluster	
Species	1 2	
setosa	0 50	
versicolor	47 3	
virginica	50 0	

Cluster 1 contains all the virginica species and almost all the versicolor while cluster two contains all setosa and the left versicolor from cluster 1.

The second method, Hierarchical clustering, is implemented in R using together the `hclust()` and `dist()` functions. The process starts assigning each observation to a cluster, initially there are as many clusters as observations, and proceeds iteratively calculating distances between clusters and joining the closest ones until there is a single cluster left which contains all the observations. You can use `?hclust` in the console for details on the algorithm, distances and more.

We will also do an example with the iris data set although we will only use a sample of 20 observations to better visualize the resulting clusters in the dendrogram. In a real application we would use for sure the full data set.

```
> set.seed(3123) #to obtain the same random sample
> data<-iris[sample(nrow(iris),20),] #random sample
> irish <- hclust(dist(data[,1:4]))
> plot(irish,main= "", labels=data[,5])
> abline(h=4,col="red")
```



With the *cutree()* function we can obtain the clusters at a chosen desired height. Looking at the dendrogram at height 4, we will obtain three clusters:

```
> irish$clusters<-cutree(irish,h=4)
> table(irish$clusters,data[,5])
```

	setosa	versicolor	virginica
1	7	0	0
2	0	7	3
3	0	0	3

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